Establishment and evolution of heterochromatin

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The eukaryotic genome is packaged into transcriptionally active euchromatin and silent heterochromatin, with most studies focused on the former encompassing the majority of protein-coding genes. The recent development of various sequencing techniques has refined this classic dichromatic partition and has better illuminated the composition, establishment, and evolution of this genomic and epigenomic “dark matter” in the context of topologically associated domains and phase-separated droplets. Heterochromatin includes genomic regions that can be densely stained by chemical dyes, which have been shown to be enriched for repetitive elements and epigenetic marks, including H3K9me2/3 and H3K27me3. Heterochromatin is usually replicated late, concentrated at the nuclear periphery or around nucleoli, and usually lacks highly expressed genes; and now it is considered to be as neither genetically inert nor developmentally static. Heterochromatin guards genome integrity against transposon activities and exerts important regulatory functions by targeting beyond its contained genes. Both its nucleotide sequences and regulatory proteins exhibit rapid coevolution between species. In addition, there are dynamic transitions between euchromatin and heterochromatin during developmental and evolutionary processes. We summarize here the ever-changing characteristics of heterochromatin and propose models and principles for the evolutionary transitions of heterochromatin that have been mainly learned from studies of Drosophila and yeast. Finally, we highlight the role of sex chromosomes in studying heterochromatin evolution.

Keywords: heterochromatin; histone modifications; chromatin conformation; sex chromosomes

Heterochromatin: an evolving concept

The understanding of heterochromatin demonstrates how the connotation of a biological paradigm can evolve with the development of research technology (Fig. 1). The term “heterochromatin” was first used by Emil Heitz in 1928,1 probably with reference to heterochromosomes (i.e., sex chromosomes),2,3 to describe the chromosomal fragments that remain densely stained throughout the cell cycle, in contrast to the other fragments of euchromatin that become invisible after telophase. Heitz later extended the staining method that he developed in liverworts to over 100 plant species and several Drosophila species, including D. melanogaster. He established that heterochromatin/euchromatin comprise the fundamental architecture of eukaryotic chromosomes and hypothesized that euchromatin is genetically active and heterochromatin genetically passive. Heterochromatic chromosomes or pieces of chromosomes contain no genes or somehow passive genes.1 It was later reported that heterochromatin has a heterogeneous distribution within and between chromosomes in Drosophila cells.4–6 In fact, Heitz noticed that heterochromatin is often associated with sex chromosomes, and some chromosomal regions are only stained in certain cell types. These were later recognized as facultative heterochromatin (fHet),7 compared with constitutive heterochromatin (cHet). From the work of Thomas Morgan, Heitz soon realized that the hypothesis of heterochromatin as being...
Figure 1. The concept of heterochromatin has been evolving since its first description in 1928 by Emil Heitz. Shown is an incomplete list of events that led to changes in the concept of heterochromatin during research developments from the cytogenetic era until very recently in the 3D genomics era. The characterization of proteins (e.g., HP1) regulating constitutive heterochromatin was largely attributed to the genetic screens in *Drosophila* for mutants affecting the position-effect variegation (PEV) phenotype. The studies into facultative heterochromatin were first marked by the discovery of the Barr body, that is, the female-specific inactivated X chromosome. After the discovery of nucleosome structure as the chromatin unit for gene regulation, many PEV-related genes were found to be the reader or writer of histone modifications within the chromatin unit. With the development of chromatin immunoprecipitation (ChIP) technology targeting these histone modifications, combined with either microarray (ChIP-chip) or sequencing analyses (ChIP-seq), it became clear that heterochromatin is usually associated with highly repetitive regions and the histone methylation marks H3K9me2/3 and H3K27me3. This facilitated the definition of the heterochromatin region at the base pair resolution in the genomic and epigenomic era. Recently, it has been shown in *Drosophila* and humans that heterochromatin forms within phase-separated droplets of HP1 protein and forms distinct topologically associated domains from those of euchromatin, as detected by Hi-C technology.

Genetically inert was not entirely valid because, as Heitz noted, “genes which lie within heterochromatin do intervene in developmental process.”

More understanding of the composition and regulation of heterochromatin was initiated after Muller’s seminal finding of the variegated eye color pattern of X-ray–irradiated *D. melanogaster* in 1930. This phenotype, position-effect variegation (PEV), was shown by subsequent examinations of polytene chromosomes to be associated with chromosomal rearrangements that positioned the white gene from euchromatin near the pericentric cHet, indicating the spreading effect of silencing heterochromatin. Interestingly, PEV has also been induced for some genes (e.g., light and its nearby genes, see below) that relocated from heterochromatin to distal euchromatin, suggesting that a proper dosage of heterochromatin-enriched protein is required for the normal expression of these genes. PEV has become a very powerful tool for identifying the critical regulators of cHet (reviewed by Ref. 12) through genetic screens for secondary mutations modifying PEV phenotypes.

The discovery of cHet-associated proteins has also been facilitated by localizing the candidate genes through immunostaining of *Drosophila* polytene chromosomes, where cHet regions are under-replicated and concentrated at the chromocenter. Several identified proteins turned out to be histone lysine methyltransferases with evolutionarily conserved domains (e.g., chromodomain and SET domain) and to have an ortholog in yeast and humans that functions similarly in posttranslational modifications (PTMs) of histones. While cHet is concentrated at telomeric and centromeric regions of all chromosomes across different cell types, in 1949 Barr and Bertram reported a female-specific heterochromatic X chromosome in cat neuronal cells, later termed the Barr body. This special form of chromosome-wide fHet was hypothesized...
to reflect the random inactivation of the X chromosome (XCI) to achieve dosage compensation.\textsuperscript{16} XCI is initiated by activity of the X-linked long noncoding RNA \textit{Xist} in eutherian mammals and is precisely controlled to be completed in the blastocyst stage, which gives rise to all somatic cells.\textsuperscript{17}

Another classic paradigm of temporally regulated fHet was identified in the homeotic (Hox) genes. The repressed or activated expression state of Hox genes during embryonic development is maintained by Polycomb and Trithorax group proteins (PcG and TrxG).\textsuperscript{18} The first PcG gene, \textit{Polycomb} (Pc), was discovered in the 1940s,\textsuperscript{19} and its mutation was later characterized to be responsible for transforming the anterior segments of \textit{Drosophila} embryos to more posterior segments.\textsuperscript{18} The first identified TrxG gene, \textit{Trithorax} (Trx), was identified as a regulator of Hox genes and to counteract PcG protein expression to produce more anterior embryonic segments.\textsuperscript{20,21} Genetic screens producing a similar phenotype to that of Pc or Trx have identified many more genes that were later defined as PcG or TrxG genes. In particular, mutations in PcG genes result in the suppression of PEV associated with cHet, suggesting a distinct mechanism underlying the gene silencing effects of fHet and cHet.\textsuperscript{12,18} A shared feature between the cHet- and fHet-related genes is that many of them—for example, the cHet-related Su(var)3-9, the PcG gene \textit{E(z)}, and the TrxG genes \textit{dSet1} and \textit{Trx}—contain the SET protein domain, suggesting that histone lysine methylation can either repress or activate gene expression. By the discovery of many more forms of histone modifications, along with the landmark visualization of nucleosome structure\textsuperscript{22,23} as a chromatin unit for gene regulation, a histone code hypothesis was proposed.\textsuperscript{24} The hypothesis states that various PTMs of histone tails on specific residues, for example, acetylation or methylation, “extend” the genetic code by altering a heritable chromatin state and hence gene expression level through recruiting downstream effector proteins (so-called “readers” and “writers” of PTMs, reviewed in Ref. 25) recognizing the histone modifications. It has become clear that cHet is usually associated with HP1 protein and di- and trimethylation of either histone H3 lysine 9 (H3K9me2/3) or histone H4 lysine 20 (H4K20me2/3, particularly at pericentric cHet). fHet is associated with PcG/TrxG proteins and trimethylation of lysine 27 (H3K27me3) alone as a repressed chromatin state but can form \textit{bivalent} chromatin, together with H3K4me3, and switch to an activated chromatin state (i.e., euchromatin) in a spatiotemporal manner. For example, the Barr body is now known to be the result of H3K27me3 modification of the inactivated X chromosome in females by PcG genes directed by the \textit{Xist} RNA. The inactivated X chromosome occupies a distinct region from its active homologous chromosome in the female nuclei, where the Barr body specifically overlaps with the silenced noncoding genomic regions.\textsuperscript{26} There are also established links between cHet and fHet, where one of the PcG proteins, EED, can bind H3K9me3,\textsuperscript{27} and loss of the H3K9 methyltransferase reduces the binding of PcG proteins to chromatin.\textsuperscript{28}

The genome-wide profiling of chromatin, a much higher resolution than chemical staining, became available after the development of microarray and Illumina sequencing, chromatin immunoprecipitation (ChIP-chip or ChIP-seq), and DNA adenine methyltransferase identification (DamID) experiments with antibodies targeting chromosomal proteins or various histone modifications. Such efforts have culminated in multiple international consortia that have collected massive ChIP-seq datasets of various histone modifications from human tissues, for example, the Encyclopedia of DNA Elements (ENCODE) project\textsuperscript{29} and the Roadmap Epigenomics project,\textsuperscript{30} or from model organisms, such as \textit{Caenorhabditis elegans} and \textit{D. melanogaster}, that is, the modENCODE project.\textsuperscript{31,32} A major aim of these consortia is to annotate the functions of noncoding genomes in the postgenomic era.

The modENCODE consortium first partitioned the \textit{Drosophila} genome into nine chromatin states by combinatorial genome-wide patterns of 18 histone modifications\textsuperscript{32} and, later, into 16 states on the basis of eight histone modifications.\textsuperscript{31} Before that, Filion \textit{et al.} used the DamID technique and partitioned the genome into five color-coded chromatin states, based on the binding maps of 53 chromatin proteins.\textsuperscript{33} The different chromatin state numbers between these works can be attributed to the differences in techniques (ChIP and DamID), antibodies (histone modifications versus chromatin proteins), and cell lines (S2 versus Kc167); the main differences are in the numbers of euchromatin states. Both identified genomic regions enriched for cHet-associated epigenetic markers H3K9me2/3 and
proteins HP1 and SU(VAR)3-9 (termed “green” chromatin in Filion et al.) and fHet markers H3K27me3 and PcG proteins (“blue” chromatin). In addition, Filion et al. characterized a third type of heterochromatin, termed “black” chromatin (bHet), which covers nearly half of the Drosophila genome and corresponds to the modENCODE states without any enrichment of measured histone modifications. bHet is enriched in the binding of SuUR protein, whose mutations influence the binding of H3K9me3 and H3K27me3. It shares many features with canonical cHet despite a lack of H3K9me2/3 enrichment, that is, it is late replicating, mainly contains low-expressed or tissue-specific genes, and transgenes inserted into bHet are much more likely to be silenced compared with those in the genomic background. Another bHet-associated protein is lamin, a major component of nuclear lamina, where canonical cHet is also concentrated. Since large genomic regions lacking histone modifications have also been found in plants and mammals, it remains to be elucidated whether bHet-associated proteins identified in Drosophila are also enriched in corresponding chromatin regions of other species. In summary, bHet together with cHet and fHet comprise the three major types of heterochromatin to our current knowledge.

Functions of heterochromatin

While fHet regions encompass genes and enhancers and can dynamically switch between the active and repressive states during development to regulate gene expression, cHet contains predominantly various types of repetitive sequences (satellite DNAs, ribosomal DNAs, and transposable elements (TEs)) and few functional genes that are usually silenced among most cell types. Such a genomic composition and the chromosomal distribution of cHet concentrated at centromeric and telomeric regions led to the assumption that the primary function of cHet is related to genome stability. Indeed, the disruption of cHet-related genes, such as HP1 of Drosophila or Swi6 of fission yeast, produces mutant phenotypes of telomere fusions, aberrant subtelomeric recombination or dysregulation of telomere lengths. Such mutations also result in the loss of pericentromeric cHet, which together with CENP-A–containing (a histone H3 variant conserved among euukaryotes) centromeric chromatin comprise the functional centromeres (reviewed in Refs. 41 and 42). The affected cells of fission yeast, Drosophila, and mice tend to show chromosome segregation errors and a higher chance of chromosome loss.

In addition to acting as chromosome structural components, cHet also guards the genome integrity by suppressing the activities, including both recombination and transcription, of repetitive sequences. In species other than yeast, cHet is also distributed on chromosome arms in between the euchromatic regions (called intercalary heterochromatin in Drosophila). Repetitive sequences at different chromosomal regions are isolated from each other by cHet domains to prevent their ectopic recombination to avoid chromosomal rearrangements, such as large insertions, deletions, and translocations. Another threat to genome integrity comes from DNA transposons and retrotransposons that can mobilize within the genome and disrupt gene functions by cut- or copy-and-paste mechanisms if not properly regulated (reviewed in Refs. 48 and 49). The suppression of these TE transcripts (posttranscriptional gene silencing) or mediate cHet formation at the TE loci to directly suppress their transcription (transcriptional silencing). The latter has been extensively studied in model organisms because it directly informs the mechanisms of cHet establishment. In fission yeast, small interfering RNAs (siRNAs) transcribed by RNA polymerase II from centromeric repeats (outer repeats) flanking the CENP-A/Cnp1–containing centromeric core regions form an RNAi machinery (RITs, including Argonaute protein AGO1) that recruits histone methyltransferase CLR4 to initiate H3K9m2/3, followed by the HP1 homolog Swi6 to establish and maintain cHet. The recruitment of HP1 to the H3K9me2/3 marks is realized by the characteristic chromodomain at the N-terminus of the HP1 protein, while its chromoshadow domain (CSD) at the C-terminus is responsible for the dimerization of HP1 proteins and further recruiting histone methyltransferase (i.e., spreading of heterochromatin). This classic paradigm of cHet assembly has been found in both plant and animal species with different types of small RNAs and proteins involved, associated with triggering DNA methylation or H3K9me2/3 modification.

The biogenesis pathway of the small RNAs related to cHet formation has been most extensively
studied in *Drosophila*, where a large portion of small RNAs interacting with Argonaute protein PIWI (PIWI-interacting RNAs, piRNAs) are transcribed from a few genomic clusters (piRNA clusters) that are predominantly composed of transposon relics. In *D. melanogaster*, transgenics carrying the 1360 DNA transposons or *invader 4* retroposons demonstrate *de novo* heterochromatin formation at mobile element euchromatic insertion sites, while deletions of their encompassed piRNA sequences compromise ectopic assembly of heterochromatin. This supports a model in which piRNAs recognize the complementary sequences of nascent TE transcripts that initiate cHet formation at the TE loci (cotranscriptional silencing). More mechanistic details of this model have been recently uncovered; it is now known in *D. melanogaster*, for example, that an HP1 paralog gene *Rhino* licenses the piRNA clusters to produce dual-strand primary piRNA transcripts by RNA polymerase II in a cHet environment, facilitated by the gene *Moonshiner*, a paralog of transcription factor TFIIA. The PIWI downstream effector gene *Panoramix* then recruits histone demethylase LSD1 and methyltransferase EGG to remove the active chromatin mark H3K4me2 and establish H3K9me3 on the transposons targeted by the piRNAs.

The case of transcribing piRNA clusters indicates that cHet is not completely silenced. Hundreds of protein-coding genes, in addition to the rRNA and small RNA loci, are also embedded in the cHet of diverse organisms, and active expression of many protein-coding genes is in fact dependent on cHet acting in either a *cis* or a *trans* manner (reviewed by Ref. 63). In the 1930s, some essential genes (e.g., *light*+) in *Drosophila* originally located in pericentromeric cHet were found to show PEV after being displaced to a novel euchromatin and heterochromatin (eu-het) boundary by chromosome rearrangements. The gene variegated expression patterns suggested that certain proteins enriched in cHet are required for their normal expression. Insights into the actual mechanisms of how cHet paradoxically (i.e., the heterochromatin paradox) regulates expression of cHet-encompassed genes were later gained from studies of the fourth chromosome pair of *D. melanogaster*.

This pair of unusual autosomes originated from a pair of ancestral heterochromatic sex chromosomes in *Diptera* species and still shares many features with canonical sex chromosomes after becoming autosomes (reviewed by Refs. 67 and 68). These chromosomes show many heterochromatic properties that were discovered as early as in the 1940s; they are small size and have a relatively low gene number (~80 genes and are thus also called dot chromosomes), a very low recombination rate, and are enriched in repetitive elements compared with other autosomes. Transcribed genes are simultaneously coated by the cHet mark H3K9me3 and the active gene mark H3K36me3, and by the protein POF (painting of fourth) that specifically binds to the dot chromosome. POF’s specific binding seems to derive from its ancestral function of specifically upregulating the hemizygous X chromosome in males of other *Diptera* species, similar to the MSL proteins of *Drosophila* species, the major protein responsible for dosage compensation. Depletion of either heterochromatin protein HP1a (an isoform of HP1 in *Drosophila*) or POF leads to a decrease in gene expression, and depletion of histone methyltransferase EGG leads to decreased binding of POF, HP1a, and H3K9me2/3 on the dot chromosomes except for the pericentromeric regions. These characteristics suggest that HP1 positively regulates the active expression of some genes, as opposed to its canonical role in gene silencing.

Such a function is not restricted to the dot chromosome genes already embedded in the cHet, but is also related to many euchromatin genes on the other autosomes of *Drosophila*. This was revealed by experiments disrupting or downregulating HP1 in *Drosophila* larvae or the Kc cell line in which many euchromatin genes, particularly cell-cycle regulatory genes, are transcriptionally affected. Although the actual mechanism of how HP1 positively regulates gene expression remains elusive, it has been reported that HP1 may facilitate transcript elongation by interacting with the RNA processing proteins hnRNP. Specifically, for the dot chromosome, HP1 is recruited to be concentrated at the body of active genes marked by H3K9me3, but not H3K9me2, in a POF/EGG-dependent manner, where POF probably binds to the nascent transcripts to upregulate gene expression. This contrasts with the repeat-enriched cHet regions of the dot chromosome or of any other chromosomes where the recruitment of HP1 is independent of...
POE, and the H3K9me2 and H3K9me3 marks show a high correlation with each other.

These studies suggested that HP1 can participate in the active transcription of some genes and that different cHet regions can be assembled by distinct mechanisms. In parallel, a recent PEV study also suggested that the telomeric regions of the *Drosophila* Y chromosome probably have a distinct type of cHet.76

Finally, the trans-acting regulatory function of cHet was originally indicated by the suppression of PEV found in flies carrying one extra Y chromosome (XYY) and the enhancement of PEV found in male flies without the Y chromosome in the 1980s.77,78 Such a pattern is more likely to be caused by the different dosages of cHet rather than the very few functional genes harbored by the Y chromosome. Given a fixed supply of proteins (e.g., satellite-binding factor D1 or HP1a) that are tightly regulated for packaging the cHet of the entire genome, it has been proposed that changes in the cHet content on one chromosome may cause redistribution of cHet and, hence, its enclosed or surrounding gene expression levels on other chromosomes (called the heterochromatin sink hypothesis).79,80 Experimental support later came from studies of *D. melanogaster* strains that differ only in their origin of Y chromosomes.81–83 These studies found that hundreds of X-linked and autosomal genes, particularly male-biased expressed genes or genes already residing in repressive chromatin regions (e.g., bHet) of other chromosomes,84 are affected by their gene expression levels.81 Although the underlying mechanisms of such Y-linked regulatory variations (YRV) remain unclear, it has been speculated that variations in repetitive sequences of the Y chromosome would either compete with other cHet regions for binding of the limited amount of heterochromatin-targeted proteins under the heterochromatin sink hypothesis or produce various amounts of small RNAs that can affect the chromatin configuration elsewhere in the genome.85 Consistent with this, recent characterization of *D. melanogaster* strains with varying numbers of Y chromosomes has found genome-wide changes in H3K9me2/3 binding, but not for the active histone mark H3K4me3,86 while Y-linked noncoding RNAs have a regulatory role in autosomal gene expression in mice.87

**Heterochromatin in space and time**

The segregation and mutual exclusion between heterochromatin and euchromatin is one of the major mechanisms that drives the genome to spatially fold into separate regulatory domains. Such a self-assembly model88 is supported in *Drosophila*89–91 and plants,92,93 where there is a strong correspondence between the chromatin state and the three-dimensional (3D) folding of the genome. The latter is measured by the recently developed chromosome conformation capture methods, particularly the high-throughput version called Hi-C. Hi-C captures genomic regions that are proximal to, and thus potentially interacting with each other, in 3D space in interphase nuclei and allows quantification of the frequency of such interactions by the numbers of normalized read pairs that span the regions in contact. At a megabase-level scale, a given chromosome can be divided into two types of compartments that have preferential long-range interactions within the regions of the same type, with the A compartment largely corresponding to euchromatin and the B compartment largely to heterochromatin.94,95 Consistent with their chromatin types, B compartment regions have a higher frequency of interactions, that is, a more condensed chromatin configuration, than that of A compartment regions. At a finer scale, depending on the experimental and bioinformatic protocols and sequencing coverage, Hi-C data can define topologically associated domains (TADs) spanning tens to hundreds of kilobase-long genomic sequences or subTADs (several kb) within a TAD.96–98 In mammals, TAD boundaries are enriched for cohesion complex and insulator proteins, such as CTCF.98,99 Removal of the CTCF/cohesion protein or CTCF binding sites leads to the disruption or shift in TAD boundaries.100–103 This indicates that in addition to self-assembly, a loop-extrusion mechanism (i.e., chromatin being extruded by the cohesion complex until it encounters the TAD boundary elements) also plays a crucial role in TAD formation.104

Different parts of the folded genome occupy different territories within the nuclei; early electron microscopy studies found that the condensed heterochromatin is mostly concentrated near the nuclear periphery or around nucleoli.95,105 Genome-wide mapping using DamID or ChIP and targeting proteins of the nuclear lamina distributed
throughout the nuclear inner membrane revealed lamina-associated domains (LADs) that can comprise over 40% of the mammalian genome (reviewed in Ref. 106). Purification of nucleoli allowed the identification of nucleolus-associated domains (NADs).\textsuperscript{107,108} While TADs reflect the interactions within the genome, LADs and NADs reflect the architectural genome organization within the nuclei. Consistent with the microscopic observations, compartment B or large TADs of heterochromatin largely overlap with LADs and NADs.\textsuperscript{96,109} Such a conventional nuclear architecture is inverted in the rod nuclei (inverted nuclei) of nocturnal mammals or mouse cells with disrupted lamina proteins (lamina B receptor), where heterochromatin is dissociated from the lamina and localized to the nuclear interior. Interestingly, in such inverted nuclei, genomic compartments and TADs are still preserved compared with the other cell types with conventional nuclear architecture of lamina–cHet associations. This suggests that the compartmentalization of the genome is not strictly dependent on the lamina. By further polymer simulations of chromatin models that fit the microscopic observation of heterochromatin distribution in the nuclei, a recent study further suggested that interactions between heterochromatic regions, but not euchromatic regions, are crucial for establishing the genomic compartments.\textsuperscript{110}

This indicated that the nuclear position of cHet can vary between different cell types. Additionally, LAD- or NAD-associated cHet can switch positions after mitosis.\textsuperscript{111} These dynamic features of cHet and the spatial formation of its membrane-less domain cannot be explained solely by local compaction of chromatin facilitated by associated proteins. An alternative model of phase separation has been proposed from the characterization of HP1 in both \textit{Drosophila} and humans.\textsuperscript{112,113} Studies found that HP1 proteins form liquid droplets \textit{in vitro} in a reversible manner dependent on either temperature or the phosphorylation state of the protein. Although it remains to be determined whether such phase-separated HP1 droplets correspond to the actual genomic compartment B or to heterochromatic TADs identified by Hi-C, they clearly share important features of heterochromatin domains, that is, nucleosomes and DNA, but not transcriptional factors preferentially partition into such droplets. Under the phase-separation model, heterochromatin regulates gene expression by precluding transcription factors from its phase boundary, while still maintaining certain flexibilities of forming larger domains by dynamically fusing with other droplets or dissolving itself if HP1 is released from the chromatin.

Such dynamic changes in cHet domains can be observed during the establishment of cHet in early \textit{Drosophila} embryos. Overall, epigenetic information—including histone modifications, DNA methylation, small RNAs, and 3D chromatin conformation—undergoes dramatic reprogramming to reset the embryo to acquire totipotency after fertilization (Fig. 2). \textit{Drosophila} embryos initially remain in a naive state without zygotic transcription and chromatin architecture and rely on maternally deposited transcripts and histone modifications that are required to activate the zygotic transcription at the later stage of mitotic cycle 14. Histone acetylation, but almost no methylation (including the canonical H3K27me3 and H3K9me2/3 marks), can be detected before zygotic activation, as early as mitotic cycle 8.\textsuperscript{116} The deposition of some acetylation marks (e.g., H3K18ac, H3K27ac, and H4K8ac) at TAD boundaries in mitotic cycle 12 suggests that they are associated with early establishment of chromatin conformation.\textsuperscript{115,117} Demethylation of another active chromatin marker H3K4me2 is required for the proper establishment of H3K9me2 marks.\textsuperscript{118} From mitotic cycle 11 on, HP1 foci have been observed to grow, fuse, and dissolve according to the progression of cell cycles; the percentage of immobile HP1 without liquid properties gradually increases before the onset of zygotic expression. Such a maturation process of heterochromatic domains may correspond to either the inclusion of more DNA/nucleosomes or the formation of contacts with the lamina during early embryogenesis.\textsuperscript{112} Counterintuitively, the aggregation of HP1 droplets is probably realized by concentration of HP1 increasing rather than by decreasing of accessibility of buried histone residues within the octamer core through its CSD. The consequential change in the conformation of nucleosomes may further promote the interactions between nucleosomes to form phase-separated liquid condensates.\textsuperscript{119} However, it remains largely unclear how heterochromatin is initially established in early embryos.
Among the three H3K9 methyltransferases in the *Drosophila* genome, only EGG has recently been found to be required for the de novo establishment of cHet, while the other two (G9A and SU(VAR)3-9) are probably required for the maintenance of cHet. PIWI also seems to play an important role, as maternal depletion of PIWI leads to the suppression of PEV in both somatic and gonadal tissues of later developmental stages, consistent with the piRNA-guided model of heterochromatin establishment in early embryos. A small subset of piRNAs might participate in this process; live imaging shows that at the onset of zygotic transcription, some (e.g., 359 satellite sequences), but not all, repetitive sequences recruit HP1 to form cHet. However, depletion of PIWI after the onset of zygotic expression has a much smaller impact on PEV, suggesting that the establishment and maintenance of cHet involve distinct pathways. Once heterochromatin is established, it seems to be generally conserved across different cell types with respect to its boundary with euchromatin.

**Evolution of heterochromatin and its associated proteins**

Genomic sequences embedded in cHet, mainly TEs and satellite sequences, usually show rapid turnover among and within species. This can be partially explained by the mutagenic effect of DNA methylation in heterochromatic regions, where methylated CpG dinucleotides have a higher rate of forming TpG by oxidative deamination. Repetitive sequences can also readily have replication slippage and unequal crossovers, resulting in the expansion or contraction of copy numbers. In addition, a compact chromatin structure may restrict the accessibility of enclosed DNAs to repair complexes. Consistent with these explanations, a multivariate study of a cancer genome characterizing the association between the mutation rate and 46 genomic...

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**Figure 2.** Establishment of heterochromatin during *Drosophila* embryogenesis. After fertilization, *Drosophila* embryos undergo 13 rapid cleavage divisions (cycles 2–13) with little zygotic transcription, as transcripts and proteins (e.g., EGG and PcG proteins) and histone modifications (e.g., H3K27me3) are mainly maternally inherited. During these early embryonic stages, both the state and topology of chromatin remain in a naive state, and there are more mobile HP1 proteins than immobile ones observed in the embryos. The establishment of the heterochromatin marker H3K9me2/3 has been recently shown to involve the histone methyltransferase EGG. At the onset of zygotic transcription at mitotic cycle 14, H3K9me2/3 can be detected by immunostaining, and there are significantly more structured TADs than in previous stages. Once the constitutive heterochromatin and chromatin topology are established, they remain largely stable across later developmental stages. The dynamics of mobile and immobile HP1a are adopted from Strom *et al.*

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features, including various histone modifications, base compositions, and the recombination rate, found that the H3K9me3 binding level showed a positive correlation and alone could account for over 40% of the mutation rate variation. In addition to these mutational effects, purifying natural selection is expected to be low on the mostly silenced junk DNA in heterochromatin and to evolve mainly by genetic drift; however, emerging evidence supports a regulatory role of transposons and satellite DNAs in many cases.

Evidence for the great interspecific variation of heterochromatic sequences comes from extensive studies in closely related species of plants, birds, and mammals. A recent characterization of six tunicate genomes revealed that TE divergence can contribute as much as a 12-fold difference in genome size between related species.

At the chromosome-wide level, the mammalian Y chromosomes and Drosophila dot chromosomes afford good examples of the effects of turnover of heterochromatic sequences. While the dot chromosome is approximately 5 Mb long in D. melanogaster, it has expanded to over 18 Mb in Drosophila ananassae because of the massive accumulation of retroposons. By contrast, the dot chromosome of Drosophila willistoni has fused to an autosome and evolved a comparable level of recombination rate and codon usage as that of euchromatic autosomes, suggesting a possible transition from a heterochromatic ancestor to euchromatin. A parallel case is the fusion between the ancestral Y chromosome shared by all Drosophila species and the dot chromosome in Drosophila pseudoobscura, with reduction in the size of intron and intergenic regions and some evidence of selective sweep on the Y chromosome after its transition to an autosome.

The ancestrally heterochromatic configuration changed on the Y chromosome after the fusion?

Mammalian Y chromosomes are more characterized than those of other species, with sequences of at least eight species having become available. The human Y chromosome contains 23 Mb of euchromatin and approximately 40 Mb of heterochromatin; the euchromatin contains a region that recently transposed from the X chromosome (X-transposed) after the divergence of human and chimpanzee, a region with 20 single-copy genes that are shared with the X chromosome (their common autosomal origin (X-degenerate)) and ampliconic regions that contain massive palindrome sequences. The Y chromosome heterochromatin is shared between humans and gorillas on the Y long-arms but absent on the Y chromosome of chimpanzees. In contrast to the expectation that Y chromosomes are highly heterochromatic and gene poor owing to relatively low recombination, over 95% of the mouse Y chromosome contains euchromatic and ampliconic regions that contain 100–300 copies of three gene families with nearly identical sequences within each family and predominant expression in the male germline. Such interspecific heterochromatic changes occurred over 5 million years ago, thus affording few trackable clues indicating the actual mechanisms or functional consequences of heterochromatin turnovers.

There have generally been many more reported cases of euchromatin changing to heterochromatin (e.g., Ref. 144) than the opposite. Genetic manipulations in model organisms, such as Drosophila and yeast that cause euchromatin-to-heterochromatin transition, or comparative studies within populations or between closely related species, have provided important insights into the molecular mechanisms of heterochromatin evolution. Based on the previous work, we summarize three models of transition from euchromatin to heterochromatin (Fig. 3).

The first model (Fig. 3A) involves de novo establishment of the heterochromatin domain in a euchromatin background. This is demonstrated by the transgenic study of 1360 DNA transposons in Drosophila, which were targeted for heterochromatin formation after being inserted into euchromatin. A similar pattern has been observed for mouse embryonic stem cell lines carrying polymorphic retroposon insertions, where the H3K9me3 and H4K20me3 marks have been observed in some, but not all, types of retroposon insertions. In D. melanogaster populations, polymorphic TE insertions have been recently shown to cause epigenetic state changes in nearby genes due to the spreading effect of heterochromatin. Such a deleterious effect on gene expression seems to be strongly selected against by natural selection; thus, heterochromatin-induced TEs (mainly long terminal repeat retroposons) are more likely to segregate at a low frequency in the population. Genomic
Figure 3. Transitions from euchromatin to heterochromatin. Three proposed models of euchromatin to heterochromatin transition. Heterochromatin is usually distributed close to the nuclear periphery and tethered to the lamina or around the nucleoli, while euchromatin is located in the nuclear interior. (A) The de novo formation of heterochromatin domains induced by TE insertions. A TE insertion into the euchromatic region may trigger heterochromatin formation mediated by small RNA pathways. This may further impact the expression of nearby genes by the spreading effect of newly formed heterochromatin domains. (B) Change in euchromatin/heterochromatin balance. The expansion of heterochromatin or ectopic formation of heterochromatin can be caused by upregulation of heterochromatin-associated proteins (e.g., histone methyltransferase SU(VAR)3-9, shown as dark blue circles) or downregulation of euchromatin-associated proteins (red circles). This has been demonstrated in Drosophila and yeast. Such chromatin boundaries form without the participation of boundary elements, such as CTCF proteins, and are thus called negotiable borders. (C) Mutations of TAD boundary sequences (green bars between the two TADs) between euchromatin and heterochromatin domains. The TAD boundary sequences are usually CTCF binding sites or transcriptionally active genes or TEs. Removal or inversion of such boundary sequences may lead to the expansion of heterochromatin domains into euchromatin. The newly formed heterochromatin domains through the TE insertions, A, or expanded heterochromatin domains through B or C will convergently interact with other preexisting heterochromatin domains at the lamina through fusions of phase-separated droplets. Such interactions may impact the nearby genes or genes on a different chromosome by reshaping the genome-wide folding. Het, heterochromatin; Eu, euchromatin; TE, transposable element.

regions with low levels of or no homologous recombination—for example, polymorphic inversions or sex chromosome regions—and insertion hotspots of TEs in euchromatin are vulnerable to transposon invasions and thus can readily form heterochromatin under this model.145,147,148

The second model (Fig. 3B) does not involve cis element–inducing heterochromatin, but rather changes in the balance between neighboring euchromatin and heterochromatin domains that shift the boundary between them. Such fluid borders between the two chromatin states form without the participation of boundary proteins, such as CTCF, and can change according to the dosage of heterochromatin- or euchromatin-associated proteins92,149 that act in trans. For example, in Drosophila, overexpression of cHet-associated protein SU(VAR)3-9 drives the expansion of cHet domains.149 Similarly, disruption of the euchromatin mark H4K16ac-associated gene Sas2 or overexpression of the gene Sir3, which participates in the deacetylation in budding yeast,150,151 leads to heterochromatin spreading. It has been demonstrated that the copy numbers and expression patterns of cHet-associated genes dramatically vary between species,152,153 probably in response to the rapid evolution of repetitive DNA sequences (see below). This may result in the shift in boundaries between euchromatin and heterochromatin, reshaping the chromatin architecture of the entire genome. Interestingly, different cHet- and euchromatin-associated proteins may form a
feedback loop to prevent promiscuous heterochromatin assembly. In budding yeast, disruption of histone demethylase EPE1 or histone acetyltransferase MST2 leads to heterochromatin spreading. EPE1/MST2 double mutant yeast, although initially deleteriously affected, quickly recovers through epigenetic mutations, which reduce the expression of another cHet-associated gene, Clr4, (a homolog of the Drosophila Su(var)3-9) to mitigate the negative effects of ectopic heterochromatin formation.\textsuperscript{154}

The third model (Fig. 3C) involves the disruption of the original boundaries between euchromatin and heterochromatin through either mutations involving the TAD boundary sequences, for example, CTCF-binding sites or transcriptionally active genes,\textsuperscript{96} or large genomic rearrangements, such as inversions (e.g., PEV), deletions, and duplications. Such changes may occur more frequently at a sub-TAD level rather than involving two neighboring TADs. The removal or shift in TAD boundaries not only affects the local chromatin states near the boundary, but also erroneously creates contacts between enhancers and promoters of different TADs, leading to the misexpression of numerous genes. Consistent with this, comparison of chromatin architectures across several mammals has found that TAD boundaries are highly conserved in syntenic regions. Genomic rearrangements, if any, preferentially occur at the boundaries or within open chromatin regions, rather than disrupting the individual TAD structure as a regulatory module.\textsuperscript{96,155} This model requires experimental test because the boundaries of TADs do not necessarily overlap with those between different types of chromatin. It is possible that in some cases, the euchromatin/heterochromatin boundary is regulated in \textit{trans} and thus resilient to the change in TAD boundaries in \textit{cis}.

In a broader sense, the impact of the evolutionary transition of euchromatin to heterochromatin is probably not restricted to their enclosed regions because of the fusion propensity of the phase-separated heterochromatin droplets.\textsuperscript{112,113,119} The tethering of the newly evolved heterochromatin droplets formed through the first model or of the expanded droplets formed through the second or third model to the nuclear lamina and their subsequent fusions with other preexisting droplets, sometimes located on a different chromosome, can influence the expression of genes near these droplets in space if their spatial positions in the nucleus are altered. This may explain the \textit{cis}-spreading effect of heterochromatin and the \textit{trans}-regulatory function of the heterochromatic Y chromosomes or the YRV.\textsuperscript{81,85} It has been recently shown that heterochromatin clustering is essential for the spatial compartmentalization of the entire genome.\textsuperscript{110} How the evolutionary transition of euchromatin to heterochromatin would influence heterochromatin clustering and what the consequences are for genome folding are intriguing questions that remain to be addressed.

Rapid turnover of heterochromatin does not simply reflect a passive evolutionary process driven by a faster mutation rate and genetic drift, but rather can have significant effects on the host genome. Species-specific heterochromatin can function as a barrier for reproductive isolation. Taking the 359 satellite sequences mentioned above, a “pioneer” satellite recruits HP1 for establishing heterochromatin during early embryogenesis in \textit{D. melanogaster},\textsuperscript{121} such satellites are concentrated on the X chromosome of \textit{D. melanogaster} but absent in \textit{Drosophila simulans}. Female hybrid offspring between the two species die during embryogenesis because of lagging X-linked chromatin derived from \textit{D. melanogaster} and a resulting mitotic defect.\textsuperscript{156} This type of speciation process can be caused by the incompatibility between fast-evolving heterochromatin and its regulatory proteins or RNAs in the hybrid genome. Consistent with this scenario, in hybrids between \textit{Drosophila mauritiana} and \textit{D. simulans}, the heterochromatin-binding allele of OdsH from \textit{D. mauritiana} erroneously decondenses Y chromosome heterochromatin of \textit{D. simulans} and thereby causes male sterility.\textsuperscript{157} The rapid evolution of heterochromatin must be contained to avoid impairing its important structural and regulatory functions. This is manifested as an “arms race” of sorts between heterochromatin and its regulatory proteins and RNAs, including those involved in heterochromatin packaging,\textsuperscript{152,153} telomere protection,\textsuperscript{158} and small RNA pathways.\textsuperscript{159,160} These proteins either show an excess of amino acid change, that is, have the signature of positive selection,\textsuperscript{161,162} or undergo rampant gene birth and death processes accompanied by newly evolved expression patterns that are usually restricted in the germline.\textsuperscript{152,153} To date, a recent characterization of 64 \textit{Diptera} genomes found a total of 121 HP1 duplications, but almost
no duplications in SU(VAR)3-9 and PcG proteins. Interestingly, the loss of a male-specific HP1 family protein HP1E correlates with the transformation of the ancestral Y chromosome to an autosome in D. pseudoobscura, suggesting a relaxed selective constraint on HP1E due to the loss of Y-linked heterochromatin.\textsuperscript{153}

**Sex chromosomes as a unique paradigm to study heterochromatin evolution**

Sex chromosomes have been associated with heterochromatin ever since its discovery\textsuperscript{2,3} (Fig. 1), as the Y or W chromosomes of most species are much more heterochromatic than other autosomes. This is because the evolution of sex necessitates the suppression of recombination between X and Y chromosomes (or Z and W chromosomes in species such as birds and butterflies), to prevent the male-determining genes or male-beneficial but female-detrimental genes (so-called sexually antagonistic genes) from appearing in females through recombination.\textsuperscript{163} The costs of sex on the Y chromosome include the massive accumulation of repetitive elements and loss of functional genes due to the reduced efficiency of natural selection in a non-recombining environment. To balance the expression level resulting from Y gene loss, fHet is also involved in various dosage compensation mechanisms to downregulate X chromosome gene expression (e.g., C. elegans and eutherian mammals).\textsuperscript{17}

The major difficulty in studying sex chromosomes is shared with that of studying heterochromatin: the highly repetitive sequence nature poses tremendous challenges for genome sequencing, assembly, and alignment, as well as gene mapping and manipulation. For example, the Y chromosome sequence of D. melanogaster is still not complete but received some significant improvement recently by PacBio sequencing,\textsuperscript{17,164} nearly 20 years after the release of its first draft genome.\textsuperscript{165}

Young sex chromosomes have initiated the heterochromatinization process very recently and still contain large portions of unique sequences, thus providing a paradigm to study the mechanisms and consequences of heterochromatin evolution. Such systems can include species that have evolved sex-determining regions very recently or those that have recently undergone fusions between the ancestral sex chromosome and autosome (called neo-sex chromosomes). The former are more concentrated in plant,\textsuperscript{166} fish,\textsuperscript{167} and amphibian\textsuperscript{168} species, while the latter have been extensively studied in Drosophila species,\textsuperscript{163} though similar systems have also been reported in birds\textsuperscript{169} and mammals.\textsuperscript{170}

It seems that the heterochromatinization process can occur very quickly on the Y chromosome after recombination is suppressed. For example, the X and Y chromosomes of papaya have been estimated to diverge from each other 2–3 million years ago. The Y-linked male-specific region without recombination only accounts for 13% of the entire chromosome, but has already exhibited several heterochromatin knobs by cytogenetic staining and is also associated with an elevated level of DNA methylation.\textsuperscript{171} A more systematic study of heterochromatin evolution comes from Drosophila miranda, which formed a neo-Y chromosome through a chromosome fusion between an autosome and an older Y chromosome that originated over 10 million years ago. The fused autosome is homologous to chr3 of D. pseudoobscura and only appears in males that do not have recombination, thus evolving like a true Y chromosome. The divergence time between D. miranda and D. pseudoobscura sets the maximum age of this neo-Y to be within 1.5 million years.\textsuperscript{172}

Previous cytogenetic studies showed that neo-Y has already accumulated an excessive amount of retroposons relative to its homolog neo-X.\textsuperscript{173,174} This probably caused the chromosome-wide increase in the H3K9me2 binding level of neo-Y, indicated by both immunostaining and ChIP-seq, relative to the neo-X and autosomes. Indeed, the binding level of H3K9me2 shows a positive correlation with the copy numbers of TEs surrounding the neo-Y linked genes. Interestingly, genes with a D. melanogaster ortholog located in black heterochromatin are much more likely to have become decorated by heterochromatin than any other genes on the neo-Y. This suggests that the ancestral chromatin configuration affects the evolution propensity of heterochromatin.\textsuperscript{175}

A similar state has been observed in another neo-Y system of Drosophila busckii, where the homologous chromosome of the dot chromosome has fused to the ancestral Y chromosome within the last 1 million years. The active genes of the D. melanogaster dot chromosome are associated with the enrichment of H3K9me3 and the silencing of genes with H3K9me2, while the two heterochromatin marks coincide with each other in the rest
of the genome. The neo-Y in *D. busckii* seems to have adopted such a unique heterochromatin configuration and only has become more enriched for H3K9me2 on silent genes, but there was no difference for H3K9me3.\textsuperscript{176}

**Conclusions and perspective**

A persistent interest in heterochromatin since its first description in the 1920s\textsuperscript{1} has been refueled by the development of sequencing techniques and new findings regarding developmental changes\textsuperscript{114,121} and biophysical properties\textsuperscript{112,113} of heterochromatin. Heterochromatic sequences had been previously assumed to be selfish and non-functional genomic parasites that only exist because of over-replication. It is clear now that heterochromatin contributes a variety of important structural and regulatory functions to the host genome. Because of the formidable cost of acquiring the complete heterochromatic sequences and the technological limits of Illumina sequencing, the burst of genomic resources for various species in the past 10 years contributed little to advancing our understanding of the diversity and evolution of heterochromatin. Most sequencing projects tended to choose the homogametic sex (e.g., a female mammal or a male bird) and sometimes intentionally omitted the heterochromatin-enriched Y or W chromosomes. High-quality heterochromatic sequences were a privilege of a few model organisms or primate species.\textsuperscript{177,178}

The development of third-generation sequencing, that is, PacBio or the nanopore platform, and its recent applications in *Drosophila*,\textsuperscript{173,179} gorillas,\textsuperscript{180,181} and humans\textsuperscript{179} have promised new discoveries regarding the diversity and functions of heterochromatin in the future. For example, centromere sequences of most species are not known except for yeast and humans, and they are epigenetically determined by CENP-A chromatin without consensus genomic sequences across species and embedded in the heterochromatin.\textsuperscript{182} A recent study in *D. melanogaster* combined third-generation sequencing and CENP-A ChIP and discovered sequences that are not present in the published *Drosophila* genome and are also enriched for CENP-A binding. It turns out that *D. melanogaster* centromeres are unexpectedly enriched for non-LTR retroelements G2/Jockey-3, instead of satellite sequences\textsuperscript{185} like humans or yeast.\textsuperscript{184} In addition, G2/Jockey-3 elements seem to be only restricted to *D. melanogaster* subgroup species, indicating the rapid evolution of centromere genomic sequences.

In addition to the new findings regarding the composition of heterochromatin sequences, we also expect more studies characterizing the spatiotemporal changes in heterochromatin in the context of chromatin topologies. Previous epigenomic or chromatin conformation studies in ENCODE or modENCODE consortiums mostly used adult tissue with pooled cell types,\textsuperscript{29,31,183} restricted by the requirement of the large amount of starting materials. However, chromatin domains and TAD structures seem to be dynamically established during embryogenesis, as shown by recently developed single-cell ChIP and Hi-C techniques.\textsuperscript{114,185} Once established, the TAD structures seem to remain stable during development\textsuperscript{117} and conserved during evolution.\textsuperscript{96,155,186}

For developmental and molecular biologists, the critical questions remaining to be answered include what factors are involved, and how do they interact with each other to recognize the genomic *cis* elements, probably with a different priority during early embryogenesis to establish the chromatin states and architectures? What are the distinct mechanisms for establishing and maintaining heterochromatin in different genomic regions? For evolutionary biologists, new questions emerge from these new discoveries: What is the resolution for the paradox of fast-evolving heterochromatin sequences and conserved TAD structures, sometimes even between humans and *Drosophila*?\textsuperscript{217} How would the arms race between heterochromatin and its regulatory proteins and RNAs drive other parts of the host genome to change? Do the genomic sequences evolve with their respective regulatory compartments as a module? Answers to these questions will greatly benefit from studying the systems that recently evolved heterochromatin, that is, polymorphic inversions or young sex chromosomes, as well as genetic manipulations that introduce or perturb the heterochromatin, with more insights to come that illuminate the genomic and epigenomic dark matter.

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Competing interests
The authors declare no competing interests.

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