

Chromatin as a potential carrier of heritable information

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Organisms with the same genome can inherit information in addition to that encoded in the DNA sequence — this is known as epigenetic inheritance. Epigenetic inheritance is responsible for many of the phenotypic differences between different cell types in multicellular organisms. Work by many investigators over the past decades has suggested that a great deal of epigenetic information might be carried in the pattern of post-translational modifications of the histone proteins, although this is not as well established as many believe. For example, it is unclear whether and how the histones, which are displaced from the chromosome during passage of the replication fork and are often exchanged from the DNA template at other times, carry information from one cellular generation to the next. Here, we briefly review the evidence that some chromatin states are indeed heritable, and then focus on the mechanistic challenges that remain in order to understand how this inheritance can be achieved.

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Introduction

Epigenetic inheritance

Because many definitions of the word ‘epigenetic’ can now be found in the literature, it is important to start this review by pointing out that we use the term in the Holliday [1] sense — traits that are *mitotically heritable* without a change in DNA sequence are called epigenetically heritable. Epigenetic inheritance is increasingly appreciated as a major contributor to processes from development to metabolism to oncogenesis. Whereas inheritance of DNA sequence is conceptually straightforward thanks to the elegance of the complementary base-pairing between paired DNA strands, the mechan-

istic basis underlying most epigenetic inheritance systems is less clear. Multiple epigenetic information carriers have been proposed including transcription factors, prions, cytosine methylation patterns, small RNAs, and chromatin structure [2–4]. Although some proposed information carriers, such as DNA methylation, are well established as heritable marks [1,5,6], the idea that post-translational histone modifications are the mechanistic basis for inheritance is more controversial. Here, we will explore the current information regarding the mechanism by which chromatin states *might* be inherited.

Chromatin as a carrier of epigenetic information

It is commonly believed that the packaging of eukaryotic genomes into chromatin provides a carrier of mitotically heritable information. This concept is not as well supported as many believe, as covered in detail in an excellent review from Ptashne [7•]. We will not reiterate those arguments, but agree with the central point that chromatin *per se* is *seldom* heritable. Instead we briefly review the evidence for cases in which chromatin might be the epigenetic information carrier. In general, it is worth noting that most epigenetic inheritance paradigms that implicate chromatin tend to involve heritable repression, rather than heritable activity, of downstream genes.

Two broad lines of evidence suggest that chromatin carries epigenetic information. First, stable epigenetically heritable cell types in multicellular organisms are frequently correlated with alternative chromatin states. For example, the β -globin gene is maintained in open chromatin in definitive erythroid cells, in closed chromatin in liver cells, and when each cell divides this packaging state is maintained; the converse is true for the albumin gene. These chromatin states are therefore epigenetically inherited (as are aspects of cell morphology and other defining features of cell type), but this does NOT necessarily mean that chromatin itself *carries* heritable information during cell division. The second major line of evidence for chromatin as an epigenetic information carrier is genetic — in *Drosophila*, Trithorax, and Polycomb class mutants that fail to epigenetically maintain cell identity encode proteins involved in histone modifying complexes [8,9]. Chromatin regulators are also genetically implicated in epigenetic inheritance systems in unicellular organisms — variegated repression of subtelomeric genes requires histone deacetylases in *Saccharomyces cerevisiae* and *P. falciparum* [10,11], and subtelomeric loci in the epigenetic OFF state are packaged into a distinctive chromatin structure that differs

significantly from the packaging state of the epigenetic ON state [12,13].

These types of evidence have led to the widespread idea that chromatin itself carries information during genomic replication. However, a situation in which a DNA-linked mark like cytosine methylation carried information at replication, but required downstream histone deacetylation as an effector function, would produce both types of observations described above. For example, yeast that have recently experienced galactose activate *GAL* genes more rapidly in response to galactose than do naïve yeast [14,15]. This heritable ‘memory’ genetically requires several chromatin regulators, but elegant heterokaryon analysis reveals that memory is carried in the cytoplasm [16]. Thus, in this system chromatin modifiers are genetically required for an epigenetic state (for unclear reasons), and chromatin structure differs in naïve and memory-induced cells, yet the information is transferable via the cytoplasm.

Beyond intellectual interest, the actual information carrier in epigenetic inheritance systems is of practical importance — there is widespread clinical interest in reversing pathological epigenetic states such as inappropriate silencing of tumor suppressors [17]. The appeal of such a therapeutic modality, beyond having yet another ‘druggable’ molecular target, is that in principle blocking a pathological epigenetic state need only be a transient chemotherapeutic regimen, thus allaying some of the drawbacks of long-term chemotherapy.

All this said, there are several scenarios where it seems chromatin has not been excluded as the actual epigenetic information carrier. First, in the case of *S. pombe* mating-type silencing, substantial evidence argues that chromatin state *per se* is heritable [18,19]. Most importantly, epigenetic transcriptional states at the *S. pombe* mating-type locus are heritable *in cis* during both mitosis and meiosis [20]. Histone modification-bound proteins such as heterochromatin protein 1 (HP1) homolog Swi6 are the likely information carriers (although absolutely ruling out an underlying DNA modification is still difficult).

The most stringent way to rule out an unidentified information carrier that re-establishes chromatin *de novo* every S phase is to recapitulate chromatin replication *in vitro* with purified components. Polycomb group proteins are involved in the maintenance of repression during development in many organisms [9]. In living cells, artificial targeting of the methyltransferase-containing PRC2 complex to a reporter construct has been used to explore inheritance of silencing [21]. Here, prior expression of GAL4-EED resulted in stable H3K27me3 and PRC1 binding and repression of the reporter gene, suggesting that, once recruited, the PRC1 complex was being retained at the reporter via a local positive

feedback. Importantly, Francis *et al.* have recently shown using an *in vitro* SV40/human cell extract replication system that the Polycomb PRC1 complex can remain associated with DNA as it undergoes coordinated DNA replication and chromatin assembly [22], even in the presence of replicating competitor templates. This key result proves that Polycomb association can be inherited during *in vitro* replication, thereby disfavoring a role for some cryptic underlying carrier.

Chromatin dynamics at the replication fork

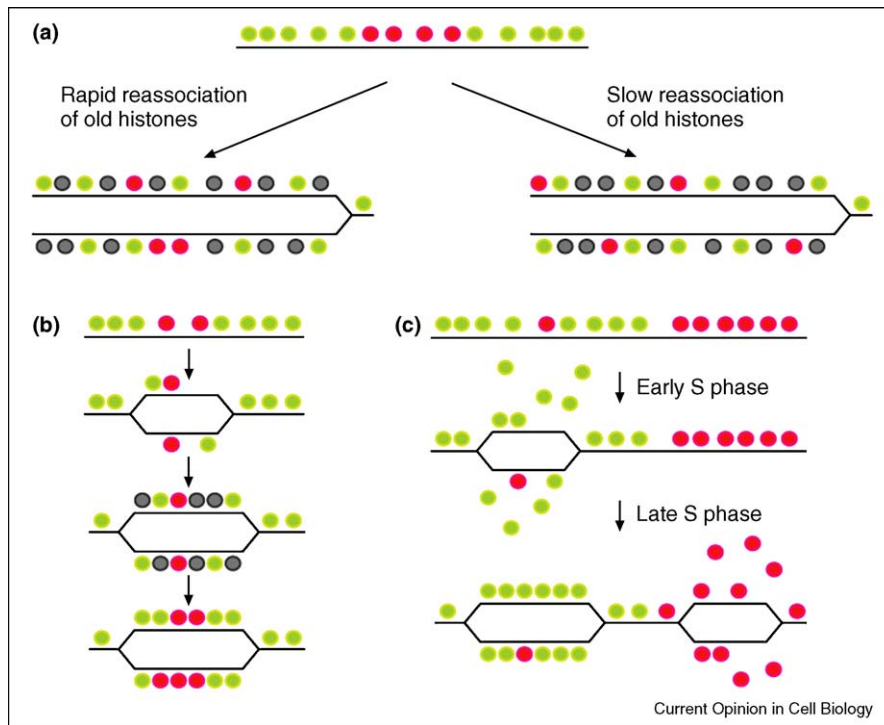
Inheritance of chromatin states is subject to several influences that do not affect standard genetic inheritance. *First*, during genomic replication the passage of the replication fork disrupts histone–DNA contacts, and old histones must reassociate with daughter chromosomes at a location close to their original location on the mother chromosome (Figure 1a). Otherwise, locus-specific epigenetic information would be randomly shuffled every generation. *Second*, old histones only account for one half of the histones on each new genome, and the remaining histones are newly synthesized during each S phase. This implies some information passage from old to new histones (Figure 1b,c), as otherwise old chromatin states would rapidly be diluted by new histones.

To make an analogy between chromatin replication and DNA replication, the first problem, that of histone dissociation, is akin to having DNA nucleotides diffuse relative to one another after the two strands are separated. The second issue, that of communication between old and new nucleosomes, is analogous to asking what the equivalent of base pairing is when nucleosomes are the ‘bases.’ Going further in this vein, one might ask what the ‘mutation’ rate is for inherited chromatin states, which might well translate into ON → OFF phenotypic epimutation rates. We discuss these issues in greater detail below.

What happens to nucleosomes during genomic replication?

After DNA replication, newly synthesized DNA must be packaged into chromatin [23]. However, in addition to providing the template for assembly of new nucleosomes, the passage of the replication fork also disrupts previously existing nucleosomes, dissociating H3/H4 tetramers from H2A/H2B dimers [24], and disrupting histone–DNA contacts [25]. Nucleosomes undergo some level of disassembly as the replication fork passes, but the exact results in terms of histone composition and position are yet undefined. Of course, if nucleosomes completely dissociated and became freely soluble then epigenetic inheritance of chromatin would become conceptually difficult. Evidence that nucleosomes do not completely dissociate comes from *in vitro* studies using the SV40 replication system — here, two competition studies using unlabeled DNA indicated that nucleosomes were not transferred to

Figure 1



Processes likely to tune the precision of inheritance of chromatin states. For all figures, green and red circles represent histones in different modification states, while gray represents free nucleosomes drawn from the nucleoplasmic pool. **(a)** Dispersal of old nucleosomes determines the precision of chromatin inheritance. Examples show two distinct dispersal patterns, using red histones to indicate a potentially heritable modification. If histones displaced by the replication machinery rapidly reassociate with the chromosome following DNA synthesis, then short domains of a given histone modification state will be heritable (left), while slow reassociation will result in spreading and dilution of the old histones in a particular state (right). **(b, c)** Two models for feedback from old nucleosomes to new. In **(b)**, new nucleosomes are modified by histone modifying enzymes recruited locally by old nucleosomes. In **(c)**, green represents acetylated nucleosomes and red represents poorly acetylated nucleosomes. Regions of the genome associated with highly acetylated nucleosomes are replicated early in S phase, concordant with the acetylation of newly synthesized histones. Late in S phase, hypoacetylated chromatin is replicated, and new histones recruited to those regions are deacetylated. These models may be distinguished based on how small domains of histone modification behave during S phase.

competitor DNA [26,27], although transfer of histones *in trans* was later observed at an 5–10-fold excess of competitor [25].

Following passage of the replication fork, parental H3/H4 molecules reassociate with one of the two daughter strands in a seemingly random fashion [28–31], accounting for half of the required nucleosome density over the two new genomes. *In vivo*, psoralen crosslinking studies show that nucleosomes associate with daughter chromosomes rapidly after the passage of the replication fork, with nucleosome-depleted DNA confined to ~1 kb surrounding the replication fork [31–33]. Translational positions of nucleosomes are established rapidly after passage of the replication fork at the yeast rDNA locus [33], although whether rapid positioning occurs globally or only at strong nucleosome positioning sequences [34] is unknown. Interestingly, in the absence of new histones, old nucleosomes appear to associate with daughter chromosomes in stretches, suggesting cooperative bind-

ing of nucleosomes to DNA [31,35]. Despite the rapid reassociation of nucleosomes with daughter genomes, it is unknown whether old nucleosomes normally reassociate close to their original positions on the mother chromosome. If nucleosomes do remain associated with their previous location, then how histones are kept from diffusing away is unclear (Figure 1a) — candidate factors include the histone chaperone CAF-1 (see below), or the Mcm proteins at the replication fork [36].

An important question regarding recruitment-based models is the relevant unit of chromatin inheritance. H2A/H2B are replaced during G1 much more extensively than H3/H4 [24,37–39], and since H3/H4-modifying enzymes are implicated in most putative chromatin inheritance paradigms, we simply consider H3/H4 here — do H3/H4 tetramers stay intact during genomic replication, or do H3/H4 dimers split apart? The vast majority of studies argue against tetramer splitting (reviewed in [40]). However, this debate has recently

been reopened by the observation that nascent H3/H4 complexes are dimers [41], in part because the histone chaperone Asf1 prevents H3/H4 tetramer formation [42,43]. A major open question is whether nucleosome diversity is generated based on dimeric intermediates. Do newly synthesized H3/H4 dimers ever mix with old ones? Crosslinking studies [29,30] suggest that this is not a common pathway, but could those bulk studies have missed mixing at critical loci? And if new dimers are only combined with new, how is mixing prevented?

Information transfer from old histones to new histones

After replication, old and new nucleosomes will often be neighbors [30]. Multiple models exist for the transfer of chromatin information from old histones to newly synthesized histones, of which we consider two. First, parental nucleosomes (carrying a given modification) could recruit chromatin modulators to adjacent newly synthesized nucleosomes, recreating even very small (~2–4 nucleosome) domains of histone modification [40,44,45]. Second, the ‘timing’ model (Figure 1c) rests on the observation that hyperacetylated chromatin is replicated early in S phase, while hypoacetylated chromatin is replicated late in S phase (see [46] for review). If hyperacetylated histones are loaded onto DNA replicated early in S phase and hypoacetylated histones are loaded late in S phase, then broad domains of acetylation could be replicated [47].

The idea that positive feedback in chromatin inheritance is provided by modifying enzymes that are recruited by the very modifications they create is a very popular one. In *S. cerevisiae*, subtelomeric silencing requires the Sir complex, which deacetylates H4K16 [45,48,49], and preferentially binds to deacetyl H4K16 [44,50–52]. In *S. pombe*, silencing of the heterochromatic loci requires the H3K9 methylase Clr4, and the HP1 homolog Swi6, which binds to H3K9me3 [18,19]. In flies and mammals, Polycomb group proteins play major roles in memory of repressive states, and mammalian PRC2 contains both a H3K27 methylase (Ezh2) and a H3K27me3-binding subunit (Eed) [53]. Consistent with the recruitment model, most well-characterized loci subject to epigenetic inheritance are associated with relatively long domains of histone modifications (see [54] for review). Inheritance of domains of many modified nucleosomes may provide cells with an error correction mechanism to decrease epimutation rates.

Despite the clear appeal of recruitment models, they are difficult to reconcile with some observations. For instance, the Eed component of the PRC2 H3K27 methylase complex binds not only to H3K27me3, but also to other ‘repressive’ histone marks such as H3K9me3 [53]. Thus, in the simplest model of recruitment, one would expect ‘crosstalk’ to result in co-occurrence of K9 and K27 methylation, yet this is

not observed in mapping studies. Furthermore, not every positive feedback loop that propagates chromatin marks laterally results in heritable chromatin states. For instance, a K4 methylase complex subunit, WDR5, directly associates with H3K4me3, and is required for global H3K4me3 [55]. Despite this positive feedback, typical H3K4me3 domains are quite short *in vivo*, and H3K4me3 patterns are generally quite plastic, correlating with the presence of RNA polymerase [54]. Therefore, the combination of modifying enzymes and modification-binding proteins is a common paradigm for building chromatin structures, but not unique to truly inherited modules.

PCNA-linked interactions at replication forks may foster epigenetic inheritance

Assembly of nucleosomes onto newly replicated DNA is carried out by a number of proteins, including the chromatin assembly complex CAF-1 which is recruited to the replication fork by the sliding clamp protein PCNA [56]. CAF-1 is required for fully efficient epigenetic silencing at the yeast subtelomeric and mating loci [57–59] while mutations in PCNA that diminish its interaction with CAF-1 also reduce heterochromatic repression [60]. It is not certain whether the primary role of CAF-1 is in the replication of silent chromatin, in the stable maintenance of silenced chromatin structures during cell cycle progression, or both. Evidence exists for both roles: in budding yeast, CAF-1 is required for stable silencing of the mating loci even in G1-arrested cells, demonstrating that its contribution to heterochromatin stability is not limited to S phase [61].

One hypothesis for CAF-1s role in epigenetic stability during S phase may be that this chaperone prevents excessive spreading of old histones from scrambling locus-specific epigenetic information (Figure 1a). An alternative hypothesis is that CAF-1 is required for feedback from old to new histones, and CAF-1 has specific protein interactions consistent with this idea. For example, during S phase, human SetDB1 associates with CAF-1 and mono-methylates H3–K9 before histone deposition [62]. After deposition, the monomethyl K9 is likely di-methylated and tri-methylated by Suv39 enzymes, creating binding sites for HP1. Notably, CAF-1 itself can recruit HP1 to pericentric heterochromatin [63,64], supporting the notion that CAF-1 contributes to a positive feedback network that favors maintenance of heterochromatic structures during replication. In another case of PCNA recruitment of positive feedback networks, the DNA methyltransferase DNMT1 is recruited to replication forks by binding to PCNA [65]. DNMT1 also interacts with G9a [66], a major H3K9-monomethyltransferase. This interaction results in monomethylation of H3K9 at sites of replication, with further methylation by Suv39H1 providing binding sites for HP1.

Timing model

A completely different idea for positive feedback from old to new histone states is a model based on replication timing. Here, the basic observation is that highly acetylated chromatin replicates early in S phase, and poorly acetylated chromatin replicates late in S phase. Elegant microinjection studies in tissue culture cells showed that reporter plasmids injected early during S phase were assembled into acetylated chromatin and were competent for transcription in the next cell cycle, whereas reporters injected late in S phase were assembled in repressive chromatin and were not transcribed [47,67]. Thus, regions designated for early replication timing could self-propagate if there was a limited pool of highly acetylated newly synthesized histones available each S phase. An implication of this model is that the unit of chromatin inheritance would be domains whose length would be set by the genomic distance replicated during each window of availability of a given histone modification.

Computational modeling of epigenetic inheritance of chromatin state

Computational modeling has proven a valuable tool for understanding key aspects of many signaling cascades, and for making predictions about key reactions in the pathway. Recently, systematic *in silico* modeling of chromatin inheritance has been carried out using the *S. pombe* mating locus as a specific model [68^{••}]. In this model, a region of 60 nucleosomes was modeled, with parameters such as feedback strength, cooperativity, distance over which feedback acts, and number of modifying cycles per cell cycle. At around the same time, a model treated Sir complex propagation in *S. cerevisiae* [69^{••}], focusing on the requirements for bistability of heterochromatin domains. A subsequent more general chromatin inheritance model [70^{••}] came to the conclusion that systems with multiple marks are naturally subject to nonlinearities, as was the case in the *S. pombe* model.

Together, these models made a variety of interesting predictions. First, all models agreed in a requirement for cooperativity in producing a bistable system. This is consistent with a general requirement for ‘ultrasensitivity’ in systems that transform graded stimuli into bistable all or none outputs [71–73]. The molecular basis for cooperativity likely differs between different candidate chromatin inheritance systems. For example, in the case of *S. pombe* silencing, cooperativity was inadvertently built in to the original model from Dodd *et al.* by including the competing acetylation and methylation of H3 lysine 9. In contrast, in *S. cerevisiae* there is no known competing modification on H4 lysine 16, although several features of Sir complex biochemistry provide candidates for the required cooperativity, such as the Sir2 deacetylation byproduct *O*-acetyl ADP-ribose [49,74], which enhances the affinity of Sir complex for chromatin [51,75]. Alternatively, competing modifications such as H3K4me3 or

H3K79me3 also provide a ‘three state’ solution (79+/Sir–, 79–/Sir–, 79–/Sir+) that eliminates the need for an explicit cooperativity term.

Another interesting prediction of the first model was that bistability was very difficult to achieve when feedback only occurred between adjacent nucleosomes — when nucleosomes only influenced their direct neighbors, resulting chromatin domains behaved as ‘patches’ of modified nucleosomes of varying size that grew and contracted as a random walk. The idea that lateral spreading of heterochromatin complexes might involve short loops or other interactions beyond adjacent nucleosomes has some support (reviewed in [76]). Additional predictions regarding the relationship between heterochromatin domain length, silencing factor concentration, and silencing stability differ depending on whether silencing is constrained between boundaries as at mating loci, or spreading into chromosomes as from telomeres [68^{••},69^{••}].

Perspective

Recent advances have illuminated much about the behavior of chromatin during replication, and have identified numerous histone modification systems that contain positive feedback loops of modifying enzymes linked to modification-binding modules. What remains to be determined is whether any of these feedback systems actually result in heritable epigenetic states (and why others do not), and the replication dynamics of chromatin that allow some old nucleosomes to influence the state of newly synthesized nucleosomes. Finally, a great deal of evidence not surveyed here points toward abundant crosstalk between epigenetic modalities such as cytosine methylation, small RNAs, and chromatin, and future work will be needed to determine the contribution of chromatin-mediated feedback loops in epigenetic stability and plasticity.

Acknowledgements

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