Shaping epigenetic memory via genomic bookmarking

Davide Michieletto^{1,*,†}, Michael Chiang^{1,†}, Davide Colì^{2,†}, Argyris Papantonis³, Enzo Orlandini², Peter R. Cook⁴ and Davide Marenduzzo^{1,*}

¹School of Physics and Astronomy, University of Edinburgh, Peter Guthrie Tait Road, Edinburgh EH9 3FD, UK, ²Dipartimento di Fisica e Astronomia and Sezione INFN, Università di Padova, Via Marzolo 8, Padova 35131, Italy, ³Centre for Molecular Medicine, University of Cologne, Robert-Koch-Str. 21, D-50931, Cologne, DE, Germany and ⁴The Sir William Dunn School of Pathology, South Parks Road, Oxford OX1 3RE, UK

Received October 09, 2017; Revised November 06, 2017; Editorial Decision November 16, 2017; Accepted November 19, 2017

ABSTRACT

Reconciling the stability of epigenetic patterns with the rapid turnover of histone modifications and their adaptability to external stimuli is an outstanding challenge. Here, we propose a new biophysical mechanism that can establish and maintain robust yet plastic epigenetic domains via genomic bookmarking (GBM). We model chromatin as a recolourable polymer whose segments bear non-permanent histone marks (or colours) which can be modified by 'writer' proteins. The threedimensional chromatin organisation is mediated by protein bridges, or 'readers', such as Polycomb Repressive Complexes and Transcription Factors. The coupling between readers and writers drives spreading of biochemical marks and sustains the memory of local chromatin states across replication and mitosis. In contrast, GBM-targeted perturbations destabilise the epigenetic patterns. Strikingly, we demonstrate that GBM alone can explain the full distribution of Polycomb marks in a whole Drosophila chromosome. We finally suggest that our model provides a starting point for an understanding of the biophysics of cellular differentiation and reprogramming.

INTRODUCTION

Cells belonging to distinct tissues in a multi-cellular organism possess exactly the same genome, yet the DNA sequence is expressed differently. This is made possible by the establishment of lineage-specific epigenetic patterns (or 'landscapes')—the heritable marking of posttranslational modifications (PTM) on histones and of methylation on DNA (1–8). Epigenetic patterns are robust, as they can be remembered across many rounds of cell division (1,2,7,9-11). At the same time, they are plastic and dynamic. They can adapt in response to external stimuli (1,9,12-14), and they are affected by disease and ageing (15,16). Additionally, many biochemical marks encoding the epigenetic information can turn over rapidly and are lost during DNA replication (17,18). For example, acetyl groups on histones have half-lives <10 min (17,19), methyl groups on histones change during the period of one cell cycle (17,20,21) and DNA methylation is modified during development (16). The turnover may originate from histone replacement/displacement during transcription (7,17,22,23), replication (7,18,24) or from stochastic PTM deposition and removal (25-27).

Our goal is to develop a biophysical model that can reconcile the reproducible and robust formation of heritable yet plastic epigenetic landscapes across cell populations in the face of the rapid turnover of the underlying histone marks. In particular we will be interested in models which can yield 'epigenetic domains', by which we mean 1D stretches of similarly-marked histones which tend to be colocalised in 3D and co-regulated (28–32). [Note that in the context of our model, the terms histone marks, chromatin states and PTM will be used interchangeably.]

Existing models describe changes of PTMs in onedimension (1D) or through effective long-range contacts; they yield smooth transitions between stable states and weak (transient) bistability (25,26,30,33–39). In contrast, our model explicitly takes into account the realistic structure and dynamics of the chromatin fibre in 3D (Figure 1)—crucial elements for the spreading of histone marks *in vivo* (11,40–45).

From the physical perspective, accounting for realistic 3D interactions (e.g. the formation of loops and trans-contacts driven by the binding of bi- and multi-valent transcription factors) triggers 'epigenetic memory' (7,8), i.e. stability of

*To whom correspondence should be addressed. Tel: +44 131 6 517742; Email: davide.michieletto@ed.ac.uk

Correspondence may also be addressed to Davide Marenduzzo. Email: dmarendu@ph.ed.ac.uk

© The Author(s) 2017. Published by Oxford University Press on behalf of Nucleic Acids Research.

[†]These authors contributed equally to this work as first authors.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com

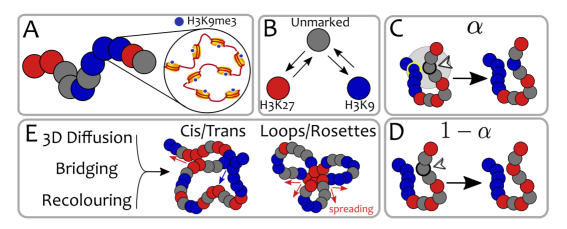


Figure 1. Polymer model with dynamic epigenetic patterns. (A) In our coarse-grain polymer model, each bead represents a group of nucleosomes and its colour captures the predominant epigenetic mark. (B) Epigenetic marks are dynamic. They can change between red, blue or grey (no mark) according to biophysical rules. For example, a red bead can be thought of as an inactive Polycomb state (marked by H3K27me3) while a blue bead as a heterochromatic segment (marked by H3K9me3). The precise nature of the marks does not affect the qualitative behaviour of this generic model. In the Voter-like dynamics, each bead must go through the unmarked state (gray) before changing to the opposite colour (26). Each bead is selected at rate k_R (see text and SM) and, (C) with probability α , it changes its colour 'closer' to that of a randomly chosen 3D-proximal bead (in this case the one circled in yellow, see also SM). (D) The same bead has probability $1 - \alpha$ to undergo a random colour conversion (in this case to red, see SM). (E) The synergy between 3D chromatin dynamics, bridging due to (implicit) binding-proteins/TFs and epigenetic recolouring gives rise to dynamic structures such as loop/rosettes and cis/trans contacts which drive (cis and trans) epigenetic spreading (indicated by red/blue arrows, see text).

the epigenetic patterns against extensive perturbations such as DNA replication (46). Within this framework, the possible 'epigenetic phases' of the system are either disordered (no macroscopic epigenetic domain is formed) or homogeneous (only one histone mark spreads over the whole chromosome). Thus, no existing biophysical model can currently predict the spontaneous emergence of multiple heritable epigenetic domains starting from a 'blank' chromatin canvas (46).

Here, we propose a model for the de novo formation, spreading and inheritance of epigenetic domains that relies solely on three elements. First, we assume a positive feedback between multivalent PTM-binding proteins ('readers') and other proteins which replace such marks ('writers'). This captures the well-known observations that, for instance, HP1 (a reader binding to heterochromatin) recruits SUV39h1 (a writer for H3K9me3 (47)), and that the Polycomb-Repressive-Complex PRC2 (a reader) contains the enhancer-of-zeste EZH2 (a writer) that spreads H3K27me3 (9,17,46,48,49). Second, we assume the presence of genomic bookmarking (GBM) factors, typically transcription factors that can bind to their cognate sites and remain dynamically associated with chromatin through mitosis (50). Examples of such GBMs include Polycomb-Group-Proteins (PcG) (11,51-53), and Posterior-Sex-Combs (PSC) (54) bound to Polycomb-Response-Elements (PREs) in Drosophila (11,41,53,54), GATA (55,56) and UBF (57) in humans and Esrbb (23,58) and Sox2 (50,59) in mouse. Here, we will use the term transcription factor (TF) to include both activators and repressors. Third, we assume that the recruitment of read-write machineries is coupled to specific GBM binding. These three assumptions allow our model to reconcile short-term turnover of PTM with long-term epigenetic memory and plasticity. Finally, we show that our model can quantitatively recapitulate the distribution of H3K27me3 mark seen in Drosophila cells *in vivo*.

MATERIALS AND METHODS

A polymer model for dynamic epigenetic patterns

To capture the dynamic nature of epigenetic landscape due to PTM turnover and histone displacement (17,58), we enhance the (semi-flexible) bead-spring polymer model for chromatin (62–70) by adding a further degree of freedom to each bead. Specifically, each bead – corresponding to one or few nucleosomes (choosing a different coarse-graining leaves our result qualitatively unaffected) – bears a 'colour' representing the instantaneous local chromatin state (e.g., H3K9me3, H3K27me3, H3K27ac, etc., see Figure 1(A)), which can dynamically change in time according to realistic biophysical rules (25,26,46) (see Figure 1(B)). This is in contrast with previous works that only accounted for static epigenetic patterns via co-polymer modelling (30,65,71,72).

We first consider a toy model in which beads may be found in one of three possible states: grey (unmarked), red (e.g. Polycomb-rich) and blue (e.g. heterochromatin-rich). [A more realistic model will be discussed later]. Beads bearing the same histone mark are mutually 'sticky', indicating the presence of implicit bridging proteins (17,65,67), and can thus bind to each other with interaction energy ε (see Figure 1(E)). All other interactions are purely repulsive. The natural time-scale for our simulations is the Brownian time $\tau_{Br} = \sigma^2/D$ which is the typical diffusion time for a bead of size σ . As discussed in the SM, this time can be estimated as $\tau_{Br} \simeq 10$ ms which is equivalent to considering a nucleoplasm viscosity of $\eta = 150$ cP and a bead of size $\sigma = 30$ nm (46).

The action of writer proteins is modelled through 'recolouring' moves occurring at rate $k_{\rm R}$; here, we set $k_{\rm R} = 0.1 \text{ s}^{-1}$ which is close to typical timescales for acetylation marks (19). In selected cases, we have also employed a faster recolouring rate of $k_{\rm R} = 10 \, {\rm s}^{-1}$ to ensure faster convergence to steady state (see SM for details on simulations and time-mapping).

Our model couples reading and writing as follows. First, a bead is selected randomly. Next, with probability α , it recruits a neighbour from spatially-proximate beads (within $r_c = 2.5\sigma$, where σ is bead size). The colour of the first bead is then shifted one step 'closer' to the colour of the second (Figure 1B and C). Otherwise (with probability 1- α), the bead undergoes a noisy conversion to one of the other colours (see Figure 1D and SM for further details).

This re-colouring scheme encodes a purely nonequilibrium process and it is akin to a 'voter' or 'infectiontype' model (25,26). In SM, we describe a 'Potts' recolouring scheme which can be arbitrarily tuned either in- or out-of-equilibrium (46). Both schemes couple 1D epigenetic information along the chromatin strand to 3D folding. Both drive a positive feedback loop between readers (which bind and bridge chromatin segments) and writers (which can change the underlying epigenetic pattern). Strikingly, both strategies lead to qualitatively similar behaviours, in which cis/trans contacts, globules and rosettes (Figure 1E) spontaneously emerge and drive the spreading of histone modifications. To simplify the presentation of our results, and because the observed behaviours are similar, we choose to report in the main text the findings obtained via the 'infection-type' model. This model may better capture the one-to-one nature of the chemical reactions required for the deposition (or writing) of histone marks (see SM for more details).

RESULTS

The phase diagram of the system entails epigenetic memory

We first map the phase diagram obtained by varying the 'feedback' parameter $f = \alpha/(1 - \alpha)$ and the attraction energy $\varepsilon/k_{\rm B}T$ between any two like-coloured beads. A more realistic model accounting for different attractions between 'Polycomb-rich' and 'heterochromatin-rich' beads is considered later.

Figure 2A shows that there are four distinct phases predicted by our minimal model. First, at small α and ε/k_BT , the fibre is swollen and epigenetically disordered (SD). At large α and ε/k_BT , the system is in the compact epigenetically ordered (CO) phase. These two states are separated by a discontinuous transition, signalled by the presence of hysteresis and coexistence (see SM). The discontinuous nature of the transition is important because it confers metastability to the two phases with respect to perturbations in the parameter space. In addition, perturbing a compact heterochromatin-rich state by extensively erasing PTM marks (e.g. during replication) fails to drive the system out of that epigenetic state (46); in other words, the global epigenetic state is remembered across genome-wide re-organisation (9,46).

The two remaining regions of the phase diagram (Figure 2A) are (i) an ordered-swollen phase (SO), observed at large α but small or moderate $\varepsilon/k_{\rm B}T$, and (ii) a compactdisordered phase (CD), found at small α and large $\varepsilon/k_{\rm B}T$. Our simulations suggest that the transitions from, or to, these states are smooth and unlike that between the SD and CO phases.

We highlight that the first order line (black thick line in Figure 2A) entails hysteresis (see SM, Supplementary Figure S3) and robustness of the states against small perturbations in the parameter space. On the other hand, a pathway that brings, for instance, a CO state into a SD one passing through the SO region, crosses continuous lines. Such a pathway in the parameter space may be a valid model to describe a change of identity of a cell, for instance during reprogramming. While this is an appealing avenue, we leave its exploration for future work as it requires a more detailed mapping between the recolouring rules of real systems and our parameter space.

Polymer simulations of the minimal model capture realistic chromatin conformations

Intriguingly, some of the phases in the phase diagram in Figure 2 correspond to structures seen in eukaryotic chromosomes. Most notably, the compact-ordered phase provides a primitive model for the structure of the inactive copy of the X chromosome in female mammals; this is almost entirely transcriptionally silent, and this state is inherited through many cell divisions (2).

The compact-disordered phase is reminiscent of 'gene deserts' (or black chromatin (28,60)). This is a state without a coherent epigenetic mark which tends to co-localise in 3D, possibly due to the linker histone H1 (28,60,73). Finally, the swollen-ordered phase is reminiscent of open and transcriptionally-active chromatin (61,74,75).

In this simplified model, feedback between readers and writers leads to unlimited spreading of a single histone mark in both ordered phases (CO and SO, see Figure 2) (46,76). Although near-unlimited spreading of silencing marks is seen in telomere position effects in yeast (40) and position-effect variegation in *Drosophila* (77)), this minimal model cannot recapitulate the existence of multiple epigenetic domains, or 'heterogeneous' epigenetic patterns.

A biophysical model for genomic bookmarking

We now introduce genomic bookmarking (GBM) to account for heterogeneous epigenetic patterns, coexistence of heritable epigenetic domains and active/inactive (A/B) compartments (31,32). A bookmark is here considered as a TF (activator or repressor) that binds to a cognate site and recruits appropriate readers or writers (see Figure 3A).

A mechanistic model of how bookmarks might guide the re-establishment of the previous epigenetic patterns after mitosis remains elusive (16,50,56,78). Here, we assume that GBMs are expressed in a tissue-specific manner and remain (dynamically) associated to chromatin during mitosis (50,54). Then, on re-entering into inter-phase, they can recruit appropriate read/write machineries and re-set the previous transcriptional programme.

In our polymer model, we account for bookmarks by postulating that some of the beads cannot change their chromatin state (Figure 3A). Thus, a red (blue) bookmark is a red (blue) bead that cannot change its colour, and otherwise behaves like other red (blue) beads. In Figure 3A,

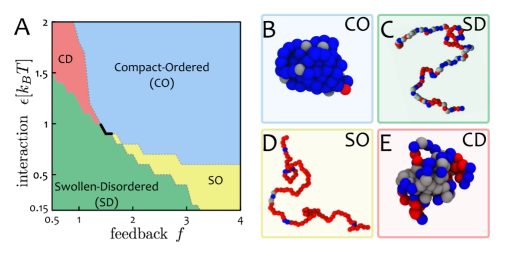


Figure 2. Phase Diagram: Chromatin states and epigenetic memory. (A) The phase diagram of the system in the space (ϵ , $f \equiv \alpha/(1 - \alpha)$) displays four distinct regions: (i) swollen-disordered (SD); (ii) compact-ordered (CO); (iii) swollen-ordered (SO) and (iv) compact-disordered (CD). The thick solid line represents a first-order transition between the SD and CO phases, whereas the dashed lines signal smoother transitions between the regions. (**B**–**E**) Representative snapshots of the stable states, which resemble conformations of chromatin seen *in vivo*. The CO phase may be associated to globally-represent heterochromatin, the SO phase to open transcriptionally-active euchromatin, and the the CD phase to 'gene deserts' characterised by low signal of PTMs and collapsed 3D conformations (28,29,60,61). The first-order nature of the SD–CO transition entails 'epigenetic memory' (8), and the CO phase is robust against extensive perturbations such as the ones occurring during replication (46).

a bookmark is indicated by an orange square that binds to DNA (rather than a PTM) and recruits read/write machineries (e.g. PRC2), which then spread a histone mark (e.g. H3K27me3) to the neighbours (2,5,17,79).

It is important to stress that, in these polymer simulations, spreading of a colour is driven by the local increase in the density of that color. Indeed, bridging drives likecolour attractions and increases the probability that a random bead will be 'infected' by a 3D-proximal bead bearing that mark. The choice of which mark dominates the local spreading is decided via symmetry breaking and we thus bias the local concentration of marks by introducing DNAbound enzymes, i.e. bookmarks (see Supplementary Movie 1).

GBM drives stable coexistence of 1D epigenetic domains and shapes the 3d chromatin organisation

We now consider a chromatin fibre where a fraction ϕ of beads are 'bookmarks' and analyse how their spatial distribution affects the epigenetic patterns in steady state. We consider three possible GBM distributions as follows: (i) *Clustered*: bookmarks are equally spaced along the fibre; the colour alternates after every n_c consecutive bookmarks $(n_c > 1$ defines the cluster size). (ii) *Mixed*: same as clustered, but now colours alternate every other bookmark $(n_c = 1)$. (iii) *Random*: random bookmarks are placed along the fibre while the fraction ϕ is kept constant.

Figure 3B–D shows the results for $\phi = 0.1$ and a chromatin fibre L = 1000 beads long. This correspond to about 3 Mb, or 1.5×10^4 nucleosomes, for a coarse graining of 3 kb per bead, i.e. a fibre with approximately one bookmark every 150 nucleosomes. Simulations are initialised with the chromatin fibre in the swollen-disordered phase and non-bookmarked regions contain equal numbers of red, blue and grey beads.

The clustered distribution of bookmarks (Figure 3B) reaches a stable epigenetic pattern with blocks of alternating colours (domains). On the contrary, the mixed bookmark distribution hinders domain formation, and the fibre remains in the SD state (Figure 3C). Remarkably, random bookmarks also yield domains in 1D (Figure 3D), even in the absence of any correlation between the location of bookmarks.

Importantly, we highlight that the bookmarking pattern affects 3D structure. Thus, in Figure 3C and D, both the random and mixed patterns yield swollen or partiallycollapsed fibres, even though the parameters used normally drive the system to a collapsed phase. [Note that our parameter choice accounts for the fact that the critical $\varepsilon(f)$ marking the SD-CO transition decreases with L.] For the random distribution, the contact map exhibits locally compact structures with coherent epigenetic marks (see arrowhead in Figure 3D) while long-range interactions between likecoloured domains are supressed. This result is in marked contrast with equilibrium models with static epigenetic pattern (30,72)). On the other hand, for clustered bookmarks. red and blue domains separately coalesce in 3D (macrophase-separation), to give a checker-board appearance of the contact map (Figure 3B) reminiscent of the pattern formaed by A/B compartments in Hi-C maps after suitable normalisation (31, 80).

We highlight that these patterns are achieved independently of the chosen initial configuration. As shown in the SM (Supplementary Figure S4), a system initialised from deep into the collapsed-disordered phase (reminiscent of condensed mitotic chromosomes) leads to the same 1D pattern of marks and 3D organisation found in Figure 3 at large times.

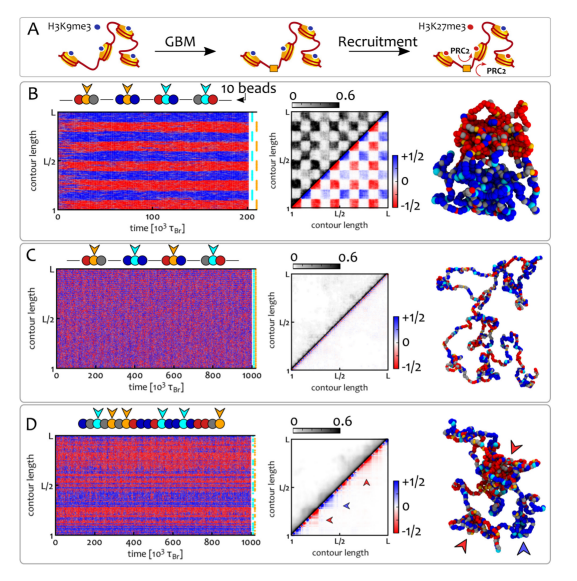


Figure 3. GBM shapes the 1D epigenetic pattern and the 3D chromatin conformation. (A) At the nucleosome level, GBM is mediated by a TF that binds to its cognate site and recruits read/write machineries that spread the respective histone mark to 3D-proximal histones (here PRC2 spreads H3K27me3). (**B**–**D**) We consider a chromatin fibre L = 1000 beads long, starting from an epigenetically random and swollen condition with $\phi = 0.1$, equivalent to one bookmark in 150 nucleosomes at 3 kb resolution and we fix f = 2 and $\varepsilon/k_B T = 0.65$. GBM is modelled by imposing a permanent colour to some beads along the fibre. Cyan and orange beads denote bookmarks for blue and red marks, respectively. Plots show kymographs (left column), average contact maps (central column) and typical snapshots (right column) for different bookmarking patterns (shown at the end of kymographs and cartoons above). Contact maps are split into two: the upper triangle shows a standard heat-map quantifying the normalised frequency of contacts between segments *i* and *j*, whereas the lower triangle shows an 'epigenetically-weighted' one in which each contact is weighted by the type of beads involved (+1 for blue-blue contacts, -1 for red-red and 0 for mixed or grey–grey). (**B**) A clustered GBM pattern yields well-defined epigenetic domains which coalesce into A/B compartments ($k_R = 0 \text{ s}^{-1}$). (**D**) Random GBM creates stable and coexisting locally-compacted structures (indicated by the arrowheads) without generating long-range contacts ($k_R = 10 \text{ s}^{-1}$). (**D**) Random GBM creates stable and coexisting locally-compacted structures (indicated by the arrowheads) without generating long-range contacts ($k_R = 10 \text{ s}^{-1}$). (**D**) Random GBM creates stable and coexisting locally-compacted structures (indicated by the arrowheads) without generating long-range contacts ($k_R = 10 \text{ s}^{-1}$). (**D**) Readom GBM creates stable and coexisting locally-compacted structures (indicated by the arrowheads) without generating long-range contacts

A critical density of bookmarks is required to form stable domains

We now ask what is the minimum density of like-coloured bookmarks needed to form stable domains. To address this question we systematically vary bookmark density and perform simulations with clustered patterns (Figure 3B) as these are the most effective way to create domains. Here, ϕ varies from 0.01 to 0.1 for a chain with L = 1000. To facilitate the analysis, we fix the domain size at 100 beads (300 kb), which is in the range of typical HiC domains (31,32,80).

We set the system to be in the collapsed-ordered phase, i.e. $\epsilon/k_{\rm B}T = 1$ and f = 2 and quantify the efficiency of domain formation by measuring the probability that bead i $(1 \le i \le L)$ is in a 'red' state, $P_{\rm red}(i)$. If ideal regular domains are formed along the fibre (i.e. if all beads have the intended colour, that of the closest bookmarks) then $P_{\rm red}(i)$ would be a perfect square wave $\Pi(i)$ (Figure 4, caption). The fidelity of domain formation can then be estimated as $\chi = 1 - \Delta^2$, where Δ^2 is the mean square deviation (variance) between $P_{\rm red}(i)$, measured in simulations, and $\Pi(i)$, i.e.

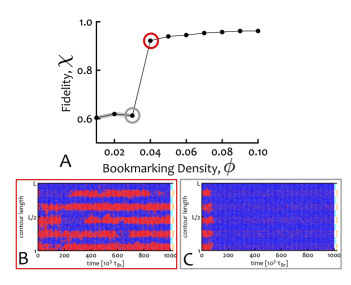


Figure 4. A critical density of bookmarks is required for stable domain formation. (A) Using the clustered pattern of bookmarks at different densities ϕ , we quantify the deviation from a 'perfect' block-like epigenetic pattern. To do this we define the 'fidelity', χ , as $1 - \Delta^2$ where $\Delta^2 = Var[P_{red}(i), \Pi(i)]$, i.e. the variance of the probability $P_{red}(i)$ of observing a red bead at position *i* with respect to the perfect square wave $\Pi(i) = 0.5 [sgn(sin(\pi i/n_d)) + 1]$, where n_d is the number of beads in a domain (here $n_d = 100$). The fidelity χ jumps abruptly from a value near its lower bound of 1/2 towards unity, at the critical $\phi_c \simeq 0.04$. (**B** and **C**) Kymographs representing the behaviour of the system at the points circled in red and grey in (A).

 $\Delta^2 = \sum_{i=1}^{L} [P_{\text{red}}(i) - \Pi(i)]^2 / L.$ The fidelity parameter is $\chi \simeq 1/2$, when the epigenetic pattern is far from the ordered block-like state and is dominated by a single colour, whereas $\chi \simeq 1$ for ideal block-like domain formation.

We observe (Figure 4A) that the system displays a phase transition near the critical density $\phi_c \simeq 0.04$. For $\phi > \phi_c$, stable domains are seen in kymographs and $\chi \simeq 1$. For $\phi < \phi_c$ instead, a single mark takes over the whole fibre. Near $\phi = \phi_c = 0.04$ there is a sharp transition between these two regimes in which domains appear and disappear throughout the simulation (see kymograph in Figure 4B).

The critical density ϕ_c corresponds to about 1 or 10 nucleosomes in about 400 as not all nucleosomes coarse-grained in a 'bookmark bead' need to be bookmarked. We argue that, crucially, not all the genome must have this critical density of bookmarks, but only regions required to robustly develop a specific domain of coherent histone marks in a given cell-line.

Biasing epigenetic landscapes with asymmetric interactions

Thus far, we have considered symmetric interactions between like-coloured beads. In other words, red-red and blue-blue interaction strengths were equal. However, such binding energies may differ if mediated by distinct proteins. Consider the case where red and blue marks encode Polycomb repression and constitutive heterochromatin, respectively. If the blue-blue interaction is larger than the redred one, the thermodynamic symmetry of the system is broken and the blue mark eventually takes over all nonbookmarked regions (Figure 5A). However, if there are bookmarks for the red mark, they locally favour the red state, whereas the stronger attraction globally favours the blue mark. This competition creates an additional route to form stable domains as exemplified in Figure 5A and B. Here, red bookmarks (identified by orange beads) are concentrated in the central segment of a chromatin fibre. Starting from a swollen and epigenetically disordered fibre, where red, blue and grey beads are equal in number, we observe that blue marks quickly invade non-bookmarked regions and convert red beads into blue ones (a process mimicking heterochromatic spreading *in vivo* (47)). However, the central segment containing the bookmarks displays a stable red domain (Figure 5A and B).

Bookmark excision but not DNA replication destabilizes the epigenetic landscape

We next asked whether the epigenetic pattern established through GBM is also stable against extensive perturbations such as DNA replication. In order to investigate this scenario we simulated semi-conservative replication of the chromatin fibre by replacing half of the (non-bookmarked) beads with new randomly coloured beads (27). In Figure 5C and D, we show that our model can 'remember' the established epigenetic pattern through multiple rounds of cell division. Importantly, the combination of 'memory' and local epigenetic order (via bookmarks) may allow cells to display 'epialleles', i.e. alleles with different transcriptional behaviours thus explaining local (or '*cis*-') memory (27,81).

We next considered a set-up relevant in light of recent experiments in *Drosophila* (53,82), where the role of polycomb-response-elements (PREs) in epigenetic memory was investigated. In these works, polycomb-mediated gene repression was perturbed as a consequence of artificial insertion or deletion of PREs. In Figure 5 we thus performed a simulated dynamic experiment where replication was accompanied by random excision of bookmarks (53) (Figure 5E and F); in practice, we remove one-fourth of the initial number of bookmarks at each replication event. Then each 'cell cycle' successively dilutes the bookmarks which at some point can no longer sustain the local red state and the region is consequently flooded with blue marks.

Importantly, the system does not display immediate loss of the red domain as soon as $\phi < \phi_c$; on the contrary, this domain is temporarily retained through local memory (see Figure 5F, LM) (9,27,81). This originates from an enhanced local density of marks together with the positive read/write feedback (see SM). [The persistence of the local memory can be tuned via the parameters of our polymer model.] These results are again consistent with experiments, as regions of the *Drosophila* genome marked with H3K27me3 are only gradually lost after PRE excision (53). Similarly, epialleles have been observed to be temporarily remembered across cell division (81).

We finally highlight that the results presented in Figure 5 are independent on the chosen initial configuration. In SM (Supplementary Figures S4 and S5) we show that starting from a collapsed and epigenetically disordered chromatin (CD phase), resembling heavily condensed and sparsely marked mitotic structures, leads to the same behaviour and strongly supports the robustness of our findings.

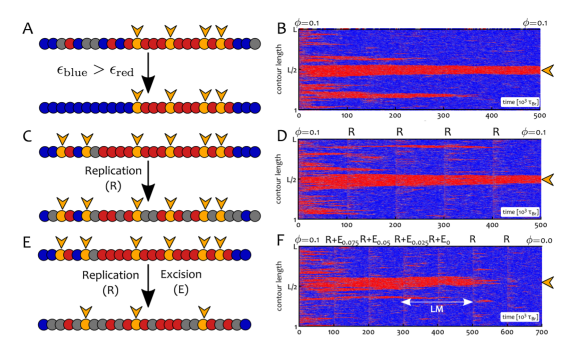


Figure 5. Asymmetric interactions and bookmark excision but not DNA replication affect the epigenetic landscape. (**A** and **B**) Here we consider the case in which blue-blue interactions are stronger than red-red ones. We set $\varepsilon_{blue} = 1k_BT$ and $\varepsilon_{red} = 0.65k_BT$ with f = 2. The central region of a chromatin segment L = 2000 beads long is initially patterned with bookmarks at density $\phi = 0.1 > \phi_c$ (this region is indicated in the kymograph by an orange arrowhead). Blue beads invade non-bookmarked regions thanks to the thermodynamic bias whereas the local red state is protected by the bookmarks. (**C** and **D**) The chromatin fibre undergoes replication cycles which extensively perturb the pattern of PTM of histones on chromatin. A semi-conservative replication event (**R**) occurs every $10^5 \tau_{Br}$ and half of the (non-bookmarked) beads become grey. The epigenetic pattern is robustly inherited. (**E** and **F**) The chromatin fibre undergoes semi-conservative replication of bookmarks (**R**+E). At each time, 1/4 of the initial bookmarks are removed and turned into grey (recolourable) beads. The epigenetic pattern is inherited until $\phi < \phi_c$. At this point, the central red domain is either immediately lost (not shown) or it can be sustained through some replication cycles (F) by local memory (LM). See also Suppl. Movie 5 for a direct comparison of the behaviour with and without bookmarks.

Chromosome-wide simulations predict the epigenetic landscape in *drosophila*

Simplified models considered thus far are useful to identify generic mechanisms; we now aim to test our model in a realistic scenario. To do so, we perform polymer simulations of the whole right arm of chromosome 3 in *Drosophila* S2 cells.

Bookmarks (orange, in Figure 6) are located on the chromosome using PSC ChIP-Seq data (54), as PSC binds to PREs during inter-phase and mitosis (54) as well as recruiting PRC2 (via molecular bridging). Some other beads are permanently coloured according to the '9-state' Hidden Markov Model (HMM, (60)). If they correspond to gene deserts (state 9), promoter/enhancers (state 1) or transcriptionally active regions (states 2-4) they are coloured grey, red and green, respectively. We further introduce an interaction between promoter and enhancer beads to favour looping, plus, an attractive interaction between gene desert (grey) beads mimicking their compaction by H1 linker histone (28) (see SM for full list of parameters). The remaining 20% of the polymer is left blank and these 'unmarked' beads are allowed to dynamically change their chromatin state into heterochromatin (blue) or polycomb (purple) according to our recolouring scheme.

We evolve the system to steady state and we evaluate the probability of finding a Polycomb mark at a certain genomic position. [To determine these probability, a bookmarked bead is counted as bearing the H3K27me3 mark when it is near beads with polycomb marks, or within large stretches of bookmarked beads.] This provides us with an *in silico* ChIP-seq track for Polycomb marks which can be compared with *in vivo* ChIP-Seq data (60) (see Figure 6 B). The two are in excellent agreement (Pearson correlation coefficient $\rho = 0.46$, against $\rho = 0.006$ for a random dataset).

Remarkably, not all bookmarked segments (orange) are populated by Polycomb marks; instead we observe that H3K27me3 spreading requires appropriate 3D folding (Figure 6B and C, insets). Bookmarks which do not contact other bookmarks due to the local epigenetic landscape do not nucleate H3K27me3 spreading. Again, this is consistent with 3D chromatin conformation being crucial for the spreading and establishment of epigenetic patterns (11,42,45).

DISCUSSION

We proposed and investigated a new biophysical mechanism for the *de novo* establishment of epigenetic domains and their maintenance through interphase and mitosis. Our simplest model requires only one element: a positive feedback between readers (e.g. binding proteins HP1, PRC2, etc.) and writers (e.g. methyltransferases SUV39, EzH2, etc.).

We performed large-scale simulations in which chromatin is modelled as a semi-flexible bead-and-spring polymer chain overlaid with a further degree of freedom representing a dynamic epigenetic patterning. Specifically, each bead is assigned a colour corresponding to the local instantaneous

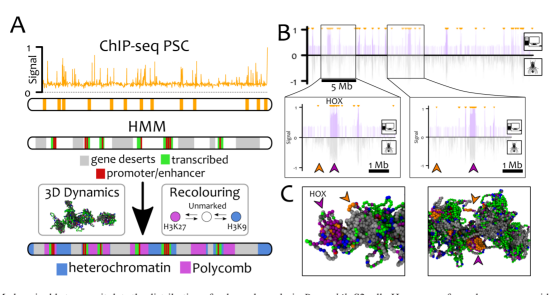


Figure 6. GBM alone is able to recapitulate the distribution of polycomb marks in *Drosophila* S2 cells. Here, we perform chromosome-wide simulations of Ch3R of Drosophila S2 cells at 3 kb resolution (L = 9302) with GBM. (A) The location of PSC/PRE (bookmarks) are mapped onto beads using ChIP-Seq data from Ref. (54). Using the '9-states' HMM data (60), gene deserts (regions lacking any mark in ChIP-seq data, state 9), promoter/enhancers (state 1) and transcriptionally active regions (states 2–4) are permanently coloured grey, red and green, respectively. The remaining beads ($\sim 20\%$) are initially unmarked (white) and may become either heterochromatin (blue) or polycomb (purple). (B) *In silico* ChIP-seq data for H3K27me3 (top half, purple lines) is compared with *in vivo* ChIP-seq (60) (bottom half, grey line). Small orange arrows at the top of the profile indicate the location of the bookmarks. The excellent quantitative agreement between the datasets is captured by the Pearson correlation coefficient $\rho = 0.46$ – to be compared with $\rho = 0.006$ obtained between a random and the experimental datasets. We highlight that not all the bookmarked beads foster the nucleation of H3K27me3 domains (see big purple/orange arrowheads in the insets, corresponding to the HOX cluster). The reason can be found by analysing the 3D conformations of the chromosome (C). The non-nucleating bookmarks (orange arrowheads), although near in 1D, are found far from potential target beads in 3D space (purple arrowheads) and so fail to yield large H3K27me3 domains. See also Suppl. Movie 6 for a direct comparison of the results with and without bookmarks.

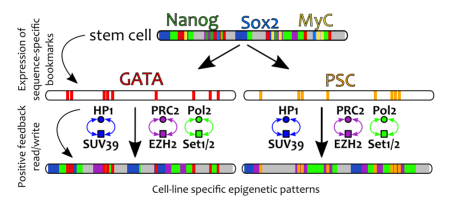


Figure 7. Model for Cellular Differentiation. We speculate that cellular differentiation may be driven by a two-step process. First, sequence-specific factors (bookmarks) are expressed as a consequence of environmental and positional cues. Second, the positive feedback set up by read/write machineries drives the establishment and maintenance of tissue-specific epigenetic patterns. As a consequence, genomic bookmarks are key targets to understand cellular differentiation and reprogramming.

epigenetic state. Readers are implicitly included by setting an attraction between like-coloured beads (65,72), whereas writers are modelled by performing re-colouring moves according to realistic and out-of-equilibrium rules (26,34) (see Figure 1).

We find that, if read-write positive feedback is sufficiently strong, a single histone mark can spread over the whole fibre and drives a discontinuous transition to a collapsedordered state (see Figure 2). This state is stable and robust against extensive perturbations such as those occurring during replication (5,17,40), when most histones are removed or displaced (2,17,23). In other words, our model displays 'epigenetic memory'. The main limitation of this simple model is that epigenetic order in real chromosomes is local, rather than global. Distinct epigenetic domains coexist on a chromosome, thereby forming an 'heterogeneous' epigenetic pattern. Our main result is that this feature of real chromosomes can be reproduced by our model when we include genomic bookmarking (GBM).

Here, we envisage bookmarks which can perform functions typical of many TFs: they recruit read/write machineries, and hence nucleate the spreading of epigenetic marks and the establishment of epigenetic domains. We also assumed that bookmarking TFs are permanently bound to DNA, however our conclusions should hold even for dynamic bookmarks that switch between bound and unbound state (50,83).

We find that stable domains can be formed with only one type of bookmark when the competing epigenetic mark is thermodynamically favoured (Figure 5). This result rationalises the common understanding that heterochromatin can spread at lengths (blue mark in Figure 5A and B) and it is stopped by actively transcribed (bookmarked) regions. Further, it is in agreement with recent genome editing experiments in Drosophila: when PRE is inserted into the genome, it provides a bookmark for H3K27me3 which leads to spreading of that mark (53), whereas PRE excision leads to (gradual) loss of the mark (53) (Figure 5). Additionally, the expression of HOX and other Polycomb-regulated genes (which contain multiple PREs) is predicted by our model to be less sensitive to deletion of single PREs (84). We suggest that this is because domains remain stable if bookmark density is kept above the critical threshold (Figure 4).

Our results strongly suggest that bookmarks can establish specific epigenetic domains by exploiting the local diffusion of chromatin and thereby 'infecting' 3D-proximal chromatin segments. The local increase in the density of a mark is then stopped either by thermodynamics (Figure 5A) or competition with other bookmarks (Figure 3B). Crucially, our model does not require any boundary element to stop the spreading of marks, which is instead self-regulated.

Losing bookmarks (via artificial excision or DNA mutation) will thus impair the ability of cells to inherit the cellline-specific epigenetic patterns. In addition, we argue that newly activated bookmarks (for instance subsequently to inflammation response or external stimuli (13,14,85)) may drive the *de novo* formation of transient epigenetic domains which allow the plastic epigenetic response to environmental changes.

We show that our model can recreate the pattern of H3K27me3 in *Drosophila* S2 cells starting solely from the position of PSC proteins acting as Polycomb bookmarks Intriguingly, our simulations show that not all bookmarks end up in H3K27me3 domains; whether or not they do, depends on their network of chromatin contacts in 3D. This is agreement with recent experiments (11,42,45) and it is also reminiscent of the well-known position effect according to which the activity of a gene depends on its local environment (14).

While our framework can be directly applied to model competition between repressive epigenetic marks, the deposition of active marks may be better modelled as resulting from a co-transcriptional positive feedback loop. In light of this, in the SM we show that a model with thermodynamically favoured heterochromatin competing with local recolouring due to transcription leads to results that are qualitatively similar to those presented in the previous sections, as long as promoters are seen as bookmarks for active marks (see SM for more details).

Our results also prompt several further questions. First, starting from a stem cell, how might different cell lineages be established? We suggest that environmental and morphological cues trigger production of lineage-specific bookmarks such as GATA (56) and PSC (54), which nucleate the positive feedback between readers and writers to generate and sustain new cell-line specific epigenetic patterns (Fig-

ure 7). Thus, bookmarks are here envisaged as key elements that should be targeted in order to understand, and manipulate, cellular differentiation. Second, how might reprogramming factors like Nanog work? We argue that their binding can 'mask' the action of pre-existing bookmarks, thereby allowing the establishment of new epigenetic patterns (58) (see also BioRxiv: https://doi.org/10.1101/127522).

In conclusion, we have extended the existing notion of GBM to include the ability of nucleating the spreading of epigenetic marks by triggering *local* read/write feedback loops. This model predicts the *de novo* establishment of heterogeneous epigenetic patterns which can be remembered across replication and can adapt in response to GBM-targeted perturbations.

Within our framework, architectural elements such as CTCF (2), Cohesins (63) and SAF-A (75) may provide the initial 3D chromatin conformation upon which the GBM-driven establishment of epigenetic landscape takes place.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR online.

ACKNOWLEDGEMENTS

The authors thank C.A. Brackley, A. Buckle, N. Gilbert, J. Allan and G. Cavalli for insightful remarks on the manuscript.

FUNDING

European Research Council for funding [Consolidator Grant THREEDCELLPHYSICS, Ref. 648050]; Papantonis lab is supported by CMMC core funding. Funding for open access charge: ERC [648050].

Conflict of interest statement. None declared.

REFERENCES

- 1. Waddington, C.H. (1942) Canalization of development and the inheritance of acquired characters. *Nature*, **150**, 563–565.
- 2. Alberts, B., Johnson, A., Lewis, J., Morgan, D. and Raff, M. (2014) Molecular Biology of the Cell. Taylor & Francis.
- Strahl, B. and Allis, C. (2000) The language of covalent histone modifications. *Nature*, 403, 41–45.
- 4. Jenuwein, T. and Allis, C.D. (2001) Translating the histone code. *Science*, **293**, 1074–1081.
- Cavalli,G. and Misteli,T. (2013) Functional implications of genome topology. *Nat. Struct. Mol. Biol.*, 20, 290–299.
- Pal,B., Bouras,T., Shi,W., Vaillant,F., Sheridan,J.M., Fu,N., Breslin,K., Jiang,K., Ritchie,M.E., Young,M. *et al.* (2013) Global changes in the mammary epigenome are induced by hormonal cues and coordinated by Ezh2. *Cell Rep.*, 3, 411–426.
- Probst,A.V., Dunleavy,E. and Almouzni,G. (2009) Epigenetic inheritance during the cell cycle. *Nat. Rev. Mol. Cell. Biol.*, 10, 192–206.
- Ng,R.K. and Gurdon,J.B. (2008) Epigenetic memory of an active gene state depends on histone H3.3 incorporation into chromatin in the absence of transcription. *Nat. Cell Biol.*, **10**, 102–109.
- Angel,A., Song,J., Dean,C. and Howard,M. (2011) A Polycomb-based switch underlying quantitative epigenetic memory. *Nature*, 476, 105–108.
- Kouskouti, A. and Talianidis, I. (2005) Histone modifications defining active genes persist after transcriptional and mitotic inactivation. *EMBO J.*, 24, 347–357.

- Ciabrelli, F., Comoglio, F., Fellous, S., Bonev, B., Ninova, M., Szabo, Q., Xuéreb, A., Klopp, C., Aravin, A., Paro, R. *et al.* (2017) Stable Polycomb-dependent transgenerational inheritance of chromatin states in Drosophila. *Nat. Genet.*, 49, 876–886.
- Stern, S., Fridmann-Sirkis, Y., Braun, E. and Soen, Y. (2012) Epigenetically heritable alteration of fly development in response to toxic challenge. *Cell Rep.*, 1, 528–542.
- 13. Wood,S. and Loudon,A. (2014) Clocks for all seasons: Unwinding the roles and mechanisms of circadian and interval timers in the hypothalamus and pituitary. *J. Endocrinol.*, **222**, R39–R59.
- Feuerborn,A. and Cook,P. R. (2015) Why the activity of a gene depends on its neighbors. *Trends Genet.*, 31, 483–490.
- 15. Pal,S. and Tyler,J. (2016) Epigenetics and aging. Sci. Adv., 2, 253-254.
- Heard, E. and Martienssen, R.A. (2014) Transgenerational epigenetic inheritance: Myths and mechanisms. *Cell*, 157, 95–109.
- Zentner, G.E. and Henikoff, S. (2013) Regulation of nucleosome dynamics by histone modifications. *Nat. Struct. Mol. Biol.*, 20, 259–266.
- Klosin, A., Reis, K., Hidalgo-Carcedo, C., Casas, E., Vavouri, T. and Lehner, B. (2017) Impaired DNA replication derepresses chromatin and generates a transgenerationally inherited epigenetic memory. *Sci. Adv.*, 3, e1701143.
- Barth, T.K. and Imhof, A. (2010) Fast signals and slow marks: the dynamics of histone modifications. *Trends Biochem. Sci.*, 35, 618–626.
- Kheir, T.B. and Lund, A.H. (2010) Epigenetic dynamics across the cell cycle. *Essays Biochem.*, 48, 107–120.
- Alabert, C., Barth, T.K., Reverón-Gómez, N., Sidoli, S., Schmidt, A., Jensen, O., Imhof, A. and Groth, A. (2015) Two distinct modes for propagation of histone PTMs across the cell cycle. *Genes Dev.*, 29, 585–590.
- Skene, P.J. and Henikoff, S. (2013) Histone variants in pluripotency and disease. *Development*, 140, 2513–2524.
- Festuccia, N., Gonzalez, I. and Navarro, P. (2017) The epigenetic paradox of pluripotent ES cells. J. Mol. Biol., 429, 1476–1503.
- Scharf,A.N.D., Barth,T.K. and Imhof,A. (2009) Establishment of histone modifications after chromatin assembly. *Nucleic Acids Res.*, 37, 5032–5040.
- Arnold, C., Stadler, P.F. and Prohaska, S.J. (2013) Chromatin computation: epigenetic inheritance as a pattern reconstruction problem. J. Theor. Biol., 336, 61–74.
- Dodd,I.B., Micheelsen,M.A., Sneppen,K. and Thon,G. (2007) Theoretical analysis of epigenetic cell memory by nucleosome modification. *Cell*, **129**, 813–822.
- 27. Berry, S., Dean, C. and Howard, M. (2017) Slow chromatin dynamics allow polycomb target genes to filter fluctuations in transcription factor activity. *Cell Syst.*, **4**, 445–457.
- Sexton, T., Yaffe, E., Kenigsberg, E., Bantignies, F., Leblanc, B., Hoichman, M., Parrinello, H., Tanay, A. and Cavalli, G. (2012) Three-dimensional folding and functional organization principles of the drosophila genome. *Cell*, **148**, 458–472.
- Dixon, J.R., Selvaraj, S., Yue, F., Kim, A., Li, Y., Shen, Y., Hu, M., Liu, J.S. and Ren, B. (2012) Topological domains in mammalian genomes identified by analysis of chromatin interactions. *Nature*, 485, 376–380.
- Jost, D., Carrivain, P., Cavalli, G. and Vaillant, C. (2014) Modeling epigenome folding: formation and dynamics of topologically associated chromatin domains. *Nucleic Acids Res.*, 42, 1–9.
- Rao,S.S.P., Huntley,M.H., Durand,N.C., Stamenova,E.K., Bochkov,I.D., Robinson,J.T., Sanborn,A.L., Machol,I., Omer,A.D., Lander,E.S. *et al.* (2014) A 3D map of the human genome at kilobase resolution reveals principles of chromatin looping. *Cell*, 159, 1665–1680.
- Dixon, J.R., Jung, I., Selvaraj, S., Shen, Y., Antosiewicz-Bourget, J.E., Lee, A.Y., Ye, Z., Kim, A., Rajagopal, N., Xie, W. *et al.* (2015) Chromatin architecture reorganization during stem cell differentiation. *Nature*, 518, 331–336.
- Micheelsen, M.A., Mitarai, N., Sneppen, K. and Dodd, I.B. (2010) Theory for the stability and regulation of epigenetic landscapes. *Phys. Biol.*, 7, 026010.
- 34. Dodd,I.B. and Sneppen,K. (2011) Barriers and silencers: A theoretical toolkit for control and containment of nucleosome-based epigenetic states. *J. Mol. Biol.*, **414**, 624–637.

- Anink-Groenen, L.C.M., Maarleveld, T.R., Verschure, P.J. and Bruggeman, F.J. (2014) Mechanistic stochastic model of histone modification pattern formation. *Epigenet. Chromatin*, 7, 30.
- Obersriebnig, M.J., Pallesen, E.M.H., Sneppen, K., Trusina, A. and Thon, G. (2016) Nucleation and spreading of a heterochromatic domain in fission yeast. *Nat. Commun.*, 7, 11518.
- Erdel, F. and Greene, E.C. (2016) Generalized nucleation and looping model for epigenetic memory of histone modifications. *Proc. Nat. Acad. Sci. U.S.A.*, 113, E4180–E4189.
- Erdel, F., Müller-Ott, K. and Rippe, K. (2013) Establishing epigenetic domains via chromatin-bound histone modifiers. *Ann. N. Y. Acad. Sci.*, 1305, 29–43.
- Erdel, F. (2017) How communication between nucleosomes enables spreading and epigenetic memory of histone modifications. *BioEssays*, 1700053, 1700053.
- 40. Talbert, P.B. and Henikoff, S. (2006) Spreading of silent chromatin: inaction at a distance. *Nat. Rev. Genet.*, **7**, 793–803.
- Lanzuolo,C., Roure,V., Dekker,J., Bantignies,F. and Orlando,V. (2007) Polycomb response elements mediate the formation of chromosome higher-order structures in the bithorax complex. *Nat. Cell Biol.*, 9, 1167–1174.
- 42. Engreitz, J.M., Pandya-jones, A., Mcdonel, P., Shishkin, A., Sirokman, K., Surka, C., Kadri, S., Xing, J., Goren, A., Lander, E.S. *et al.* (2013) The Xist lncRNA exploits three-dimensional genome architecture to spread across the X chromosome. *Science*, **341**, 1–9.
- Pinter,S.F., Sadreyev,R.I., Yildirim,E., Jeon,Y., Ohsumi,T.K., Borowsky,M. and Lee,J.T. (2012) Spreading of X chromosome inactivation via a hierarchy of defined Polycomb stations. *Genome Res.*, 22, 1864–1876.
- 44. Schauer, T., Ghavi-Helm, Y., Sexton, T., Albig, C., Regnard, C., Cavalli, G., Furlong, E.E. and Becker, P.B. (2017) Chromosome topology guides the Drosophila Dosage Compensation Complex for target gene activation. *EMBO Rep.*, e201744292.
- Deng, W., Rupon, J.W., Krivega, I., Breda, L., Motta, I., Jahn, K.S., Reik, A., Gregory, P.D., Rivella, S., Dean, A. and Blobel, G.A. (2014) Reactivation of developmentally silenced globin genes by forced chromatin looping. *Cell*, **158**, 849–860.
- 46. Michieletto, D., Orlandini, E. and Marenduzzo, D. (2016) Polymer model with epigenetic recolouring reveals a pathway for the de novo establishment and 3D organisation of chromatin domains. *Phys. Rev. X*, 6, 041047.
- Hathaway, N.A., Bell, O., Hodges, C., Miller, E.L., Neel, D.S. and Crabtree, G.R. (2012) Dynamics and memory of heterochromatin in living cells. *Cell*, 149, 1447–1460.
- Hauri,S., Comoglio,F., Seimiya,M., Gerstung,M., Glatter,T., Hansen,K., Aebersold,R., Paro,R., Gstaiger,M. and Beisel,C. (2016) A high-density map for navigating the human polycomb complexome. *Cell Rep.*, **17**, 583–595.
- Collinson,A., Collier,A.J., Morgan,N.P., Sienerth,A.R., Chandra,T., Andrews,S. and Rugg-Gunn,P.J. (2016) Deletion of the polycomb-group protein EZH2 leads to compromised self-renewal and differentiation defects in human embryonic stem cells. *Cell Rep.*, 17, 2700–2714.
- 50. Teves, S.S., An, L., Hansen, A.S., Xie, L., Darzacq, X. and Tjian, R. (2016) A dynamic mode of mitotic bookmarking by transcription factors. *Elife*, **5**, 1–24.
- 51. Kassis, J.A. and Brown, J.L. (2013) Polycomb group response elements in Drosophila and vertebrates. *Adv. Genet.*, **81**, 83.
- Schuettengruber, B., Oded Elkayam, N., Sexton, T., Entrevan, M., Stern, S., Thomas, A., Yaffe, E., Parrinello, H., Tanay, A. and Cavalli, G. (2014) Cooperativity, specificity, and evolutionary stability of polycomb targeting in Drosophila. *Cell Rep.*, 9, 219–233.
- Laprell, F., Finkl, K. and Müller, J. (2017) Propagation of Polycomb-repressed chromatin requires sequence-specific recruitment to DNA. *Science*, 8266, eaai8266.
- Follmer, N.E., Wani, A.H. and Francis, N.J. (2012) A polycomb group protein is retained at specific sites on chromatin in mitosis. *PLoS Genet.*, 8, e1003135.
- 55. Kadauke, S., Udugama, M.I., Pawlicki, J.M., Achtman, J.C., Jain, D.P., Cheng, Y., Hardison, R.C. and Blobel, G.A. (2012) Tissue-specific mitotic bookmarking by hematopoietic transcription factor GATA1. *Cell*, **150**, 725–737.
- Kadauke, S. and Blobel, G.A. (2013) Mitotic bookmarking by transcription factors. *Epigenet. Chromatin*, 6, 6.

- Grob, A., Colleran, C. and McStay, B. (2014) Construction of synthetic nucleoli in human cells reveals how a major functional nuclear domain is formed and propagated through cell division. *Genes Dev.*, 28, 220–230.
- Festuccia, N., Dubois, A., Vandormael-Pournin, S., Tejeda, E.G., Mouren, A., Bessonnard, S., Mueller, F., Proux, C., Cohen-Tannoudji, M. and Navarro, P. (2016) Mitotic binding of Esrrb marks key regulatory regions of the pluripotency network. *Nat. Cell Biol.*, 18, 1139–1148.
- Deluz, C., Friman, E.T., Strebinger, D., Benke, A., Raccaud, M., Callegari, A., Leleu, M., Manley, S. and Suter, D.M. (2016) A role for mitotic bookmarking of SOX2 in pluripotency and differentiation. *Genes Dev.*, **30**, 2538–2550.
- Kharchenko, P.V., Alekseyenko, A.A., Schwartz, Y.B., Minoda, A., Riddle, N.C., Ernst, J., Sabo, P.J., Larschan, E., Gorchakov, A.A., Gu, T. *et al.* (2011) Comprehensive analysis of the chromatin landscape in Drosophila melanogaster. *Nature*, 471, 480–485.
- Gilbert, N. and Bickmore, W.A. (2006) Structure and Transcription. Biochem. Soc. Symp., 73, 59–66.
- 62. Rosa, A. and Everaers, R. (2008) Structure and dynamics of interphase chromosomes. *PLoS Comp. Biol.*, **4**, 1.
- Fudenberg, G., Imakaev, M., Lu, C., Goloborodko, A., Abdennur, N. and Mirny, L.A. (2016) Formation of chromosomal domains by loop extrusion. *Cell Rep.*, 15, 2038–2049.
- 64. Mirny,L.A. (2011) The fractal globule as a model of chromatin architecture in the cell. *Chromosom. Res.*, **19**, 37–51.
- 65. Brackley, C.A., Johnson, J., Kelly, S., Cook, P.R. and Marenduzzo, D. (2016) Simulated binding of transcription factors to active and inactive regions folds human chromosomes into loops, rosettes and topological domains. *Nucleic Acids Res.*, 44, 3503–3512.
- 66. Brackley, C.A., Michieletto, D., Mouvet, F., Johnson, J., Kelly, S., Cook, P.R. and Marenduzzo, D. (2016) Simulating topological domains in human chromosomes with a fitting-free model. *Nucleus*, 7, 453–461.
- Brackley, C.A., Taylor, S., Papantonis, A., Cook, P.R. and Marenduzzo, D. (2013) Nonspecific bridging-induced attraction drives clustering of DNA-binding proteins and genome organization. *Proc. Natl. Acad. Sci. U.S.A.*, **110**, E3605–E3611.
- Cheng, T.M., Heeger, S., Chaleil, R.A., Matthews, N., Stewart, A., Wright, J., Lim, C., Bates, P.A. and Uhlmann, F. (2015) A simple biophysical model emulates budding yeast chromosome condensation. *Elife*, 4, 1–22.
- 69. Sanborn,A.L., Rao,S.S.P., Huang,S.C., Durand,N.C., Huntley,M.H., Jewett,A.I., Bochkov,I.D., Chinnappan,D., Cutkosky,A., Li,J. *et al.* (2015) Chromatin extrusion explains key features of loop and domain formation in wild-type and engineered genomes. *Proc. Natl. Acad. Sci. U.S.A.*, **112**, 201518552.
- 70. Rosa, A., Becker, N.B. and Everaers, R. (2010) Looping probabilities in model interphase chromosomes. *Biophys. J.*, **98**, 2410–2419.
- Di Pierro, M., Zhang, B., Aiden, E.L., Wolynes, P.G. and Onuchic, J.N. (2016) Transferable model for chromosome architecture. *Proc. Natl. Acad. Sci. U.S.A.*, **113**, 201613607.

- Barbieri, M., Chotalia, M., Fraser, J., Lavitas, L.M., Dostie, J., Pombo, A. and Nicodemi, M. (2012) Complexity of chromatin folding is captured by the strings and binders switch model. *Proc. Natl. Acad. Sci. U.S.A.*, **109**, 16173–16178.
- 73. Filion,G.J., van Bemmel,J.G., Braunschweig,U., Talhout,W., Kind,J., Ward,L.D., Brugman,W., de Castro,I.J., Kerkhoven,R.M., Bussemaker,H.J. *et al.* (2010) Systematic protein location mapping reveals five principal chromatin types in Drosophila cells. *Cell*, **143**, 212–224.
- Gilbert, N., Gilchrist, S. and Bickmore, W.A. (2004) Chromatin Organization in the Mammalian Nucleus. *Int. Rev. Cytol.*, 242, 283–336.
- Nozawa,R.S., Boteva,L., Soares,D.C., Naughton,C., Dun,A.R., Ramsahoye,B., Bruton,P.C., Saleeb,R.S., Arnedo,M., Hill,B. *et al.* (2017) SAF-A regulates interphase chromosome structure through oligomerisation with chromatin- associated RNAs. *Cell*, 169, 1214–1227.
- Michieletto, D., Orlandini, E. and Marenduzzo, D. (2017) Epigenetic transitions and knotted solitons in stretched chromatin. *Sci. Rep.*, 7, 14642.
- 77. Schotta,G., Ebert,A., Krauss,V., Fischer,A., Hoffmann,J., Rea,S., Jenuwein,T., Dorn,R. and Reuter,G. (2002) Central role of Drosophila SU(VAR)3-9 in histone H3-K9 methylation and heterochromatic gene silencing. *EMBO J.*, **21**, 1121–1131.
- Sarge, K.D. and Park-Sarge, O.K. (2005) Gene bookmarking: keeping the pages open. *Trends Biochem. Sci.*, 30, 605–610.
- Cheutin, T. and Cavalli, G. (2012) Progressive polycomb assembly on H3K27me3 compartments generates Polycomb bodies with developmentally regulated motion. *PLoS Genet.*, 8, e1002465.
- Lieberman-Aiden, E., van Berkum, N.L., Williams, L., Imakaev, M., Ragoczy, T., Telling, A., Amit, I., Lajoie, B.R., Sabo, P.J., Dorschner, M.O. *et al.* (2009) Comprehensive mapping of long-range interactions reveals folding principles of the human genome. *Science*, **326**, 289–293.
- Berry, S., Hartley, M., Olsson, T.S.G., Dean, C. and Howard, M. (2015) Local chromatin environment of a Polycomb target gene instructs its own epigenetic inheritance. *Elife*, 4, 1–11.
- Coleman, R.T. and Struhl, G. (2017) Causal role for inheritance of H3K27me3 in maintaining the OFF state of a Drosophila HOX gene. *Science*, **356**, eaai8236.
- Brackley, C.A., Liebchen, B., Michieletto, D., Mouvet, F., Cook, P.R. and Marenduzzo, D. (2017) Ephemeral protein binding to DNA shapes stable nuclear bodies and chromatin domains. *Biophys J.*, **112**, 1085–1093.
- De,S., Mitra,A., Cheng,Y., Pfeifer,K. and Kassis,J.A. (2016) Formation of a polycomb-domain in the absence of strong polycomb response elements. *PLoS Genet.*, **12**, 1–22.
- Kirmes, I., Szczurek, A., Prakash, K., Charapitsa, I., Heiser, C., Musheev, M., Schock, F., Fornalczyk, K., Ma, D., Birk, U. et al. (2015) A transient ischemic environment induces reversible compaction of chromatin. *Genome Biol.*, 16, 246.