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**Citation:** Boettiger, Alistair N. et al. "Super-Resolution Imaging Reveals Distinct Chromatin Folding for Different Epigenetic States." Nature 529, 7586 (January 2016): 418–422 © 2016 Macmillan Publishers Limited

**As Published:** http://dx.doi.org/10.1038/NATURE16496

Publisher: Nature Publishing Group

Persistent URL: http://hdl.handle.net/1721.1/112795

**Version:** Author's final manuscript: final author's manuscript post peer review, without publisher's formatting or copy editing

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Author manuscript *Nature*. Author manuscript; available in PMC 2016 July 13.

Published in final edited form as:

Nature. 2016 January 21; 529(7586): 418–422. doi:10.1038/nature16496.

# Super-resolution imaging reveals distinct chromatin folding for different epigenetic states

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#### Abstract

Metazoan genomes are spatially organized at multiple scales, from packaging of DNA around individual nucleosomes to segregation of whole chromosomes into distinct territories<sup>1-5</sup>. At the intermediate scale of kilobases to megabases, which encompasses the sizes of genes, gene clusters and regulatory domains, the three-dimensional (3D) organization of DNA is implicated in multiple gene regulatory mechanisms $^{2-4,6-8}$ , but understanding this organization remains a challenge. At this scale, the genome is partitioned into domains of different epigenetic states that are essential for regulating gene expression 9-11. Here, we investigate the 3D organization of chromatin in different epigenetic states using super-resolution imaging. We classified genomic domains in Drosophila cells into transcriptionally active, inactive, or Polycomb-repressed states and observed distinct chromatin organizations for each state. Remarkably, all three types of chromatin domains exhibit power-law scaling between their physical sizes in 3D and their domain lengths, but each type has a distinct scaling exponent. Polycomb-repressed chromatin shows the densest packing and most intriguing folding behaviour in which packing density increases with domain length. Distinct from the self-similar organization displayed by transcriptionally active and inactive chromatin, the Polycomb-repressed domains are characterized by a high degree of chromatin intermixing within the domain. Moreover, compared to inactive domains, Polycomb-repressed

RNA-Seq data have been deposited at Gene Expression Omnibus (GEO) under the accession code GSE75060. The authors declare no competing financial interests.

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Supplementary Information (including Supplementary Methods and Supplementary Table) is linked to the online version of the paper.

Author contributions A.N.B. and X.Z. conceived the project. A.N.B., B.B., and X.Z. designed the experiments and simulations with input from C.W. and L.M. A.N.B. and B.B. performed the experiments, data analysis and simulations. J.R.M. conceived the high yield probe synthesis method, and J.R.M. and A.N.B developed this method. B.J.B. assisted with initial probe synthesis. S.W. assisted with live-cell STORM experiments. G.F. and M.I. assisted with methods for simulation. A.N.B., B.B., L.M., C.W., and X.Z. interpreted the results. A.N.B., B.B., J.R.M. and X.Z. wrote the manuscript, with inputs from B.J.B., G.F., M.I., L.M. and C.W.

domains spatially exclude neighbouring active chromatin to a much stronger degree. Computational modelling and knockdown experiments suggest that reversible chromatin interactions mediated by Polycomb-group proteins plays an important role in these unique packaging properties of the repressed chromatin. Taken together, our super-resolution images reveal distinct chromatin packaging for different epigenetic states at the kilobase-to-megabase scale, a length scale that is directly relevant to genome regulation.

> Multiple lines of evidence suggest that the spatial organization of chromatin at the kilobaseto-megabase is important for genomic functions<sup>2–4,6–8,10–12</sup>. The sizes of genes, gene clusters and regulatory domains all occur in this range; in addition, physical interactions between genomic elements separated by this distance range, such as promoter-enhancer interactions, are important for gene activity<sup>2–4</sup>. Recent high-throughput chromatin conformation capture measurements revealed that individual chromosomes are partitioned into contact domains or topologically associating domains with lengths ranging from tens of kilobases (kb) to multiple megabases (Mb), and that this structural organization could be relevant to a variety of genome functions<sup>3,6–8,10–12</sup> At this length scale, chromatin is also demarcated into domains of distinct epigenetic states characterized by biochemical modifications and DNA-binding proteins<sup>9–11</sup>. Yet, how the 3D spatial organization of chromatin differs amongst these different epigenetic states is largely unknown.

> Direct imaging of the spatial organization of chromatin in different epigenetic states requires the ability to specifically label genomic DNA *in situ* and to resolve chromatin structures beyond the diffraction-limit resolution of ~200 nm. Here, we used fluorescent *in situ* hybridization (FISH) to label specific regions of the genome with complementary oligonucleotide probes tagged with fluorescent dyes. We adopted and modified a previously reported Oligopaint approach<sup>13,14</sup> to produce tens of thousands of unique, oligonucleotide probes to label kilobase-to-megabase long genomic regions using massively parallel oligo synthesis<sup>13–16</sup>. High-yield probe synthesis was achieved with a recently described enzymatic amplification method<sup>17</sup> (Extended Data Fig. 1 and Supplementary Methods). We used osmotically balanced fixation conditions that minimized shrinkage effect (Supplementary Methods) and observed no detectable chromatin shrinkage (Extended Data Fig. 2). We then imaged the labelled chromatin regions using three-dimensional stochastic optical reconstruction microscopy (3D-STORM)<sup>18,19</sup>, a single-molecule-based super-resolution imaging method. This approach yielded images of specific genomic regions in cells with 20nm *xy* and 50-nm *z* resolution<sup>14</sup>.

> We imaged 46 epigenetically defined genomic domains (Extended Data Fig. 3a; Extended Data Table) in *Drosophila* Kc<sub>167</sub> cells. We classified these regions into three major epigenetic states — transcriptionally active, inactive, and Polycomb-repressed (hereafter referred to as *active, inactive* and *repressed*, respectively) — based on enrichment of histone modifications and regulatory proteins from ChIP-seq and DamID data (Fig. 1a), as described previously<sup>10,11,20</sup> and in Supplementary Methods. *Active* chromatin domains were selected based on the enrichment of the histone modifications H3K4me2 or H3K79me3. *Repressed* domains were selected based on enrichment for H3K27me3 or Polycomb Group (PcG) proteins. *Inactive* domains were selected based on the predominance of unmodified histones

and a depletion of PcG proteins and transcriptional activators. The lengths of the selected domains span those observed for all three epigenetic types in *Drosophila* ( $\sim$ 10–500 kb).

Compared to conventional fluorescence images, STORM images of these domains revealed substantially more structural information (Fig. 1b and Extended Data Fig. 4). From these super-resolution images, we first measured the physical volume occupied by each domain, as a measure of the compactness of the DNA in the domain (Fig. 1c; Extended Data Fig. 3b, c). The volume measurements showed cell-to-cell variations (Extended Data Fig. 5a, b), which may reflect differences in expression state or cell cycle. However, the values averaged over ~50 randomly selected cells for each domain exhibited a clear difference among the three types of epigenetic domains. Across the nearly two orders of magnitude of domain lengths that we studied, the median volumes of the *active* domains were always larger than those of the *repressed* domains (Fig. 1c, solid circles). These results are in line with previous data showing that PcG proteins can lead to chromatin compaction<sup>2,21–23</sup> and that actively transcribed chromatin regions tend to be more open than non-transcribing regions<sup>2</sup>.

Notably, the volume (V) of the chromatin domains exhibited a power-law scaling behaviour with the domain length (L), i.e.  $V \sim L^b$ , and the scaling exponent b was distinct for the three different epigenetic states (Fig. 1c, solid circles; Extended Data Fig. 3b, c). Inactive chromatin domains had a scaling exponent of  $b = 1.00 \pm 0.04$  (± standard error), indicating that the 3D density of the chromatin was constant over different domain lengths. Active domains had a scaling exponent significantly greater than 1 ( $b = 1.26 \pm 0.05$ ), indicating increasingly less dense packaging for larger domains. Repressed domains exhibited the most intriguing scaling behaviour, with a scaling exponent that was notably less than 1 (b = 0.76 $\pm$  0.03), indicating that the packaging density increased with increasing domain length. As an alternative measure of the physical sizes of chromatin domains, we determined the radius of gyration ( $R_{\sigma}$ ), defined as the root-mean square distance of molecule positions measured by STORM in each domain from the centroid of these positions in the domain (Supplementary Methods). Power-law scaling was also observed for  $R_g$  as a function of L, i.e.  $R_{\sigma} \sim L^{c}$ , with the scaling exponents  $c = 0.37 \pm 0.02$ ,  $0.30 \pm 0.02$ , and  $0.22 \pm 0.02$ , for active, inactive and repressed domains, respectively (Fig. 1d, solid circles; Extended Data Fig. 3d). These scaling behaviours were conserved across different genomic regions on multiple chromosomes (Extended Data Fig 3a, b), suggesting that the different packaging behaviours are characteristic of the epigenetic states. Epigenetic states also influence the scaling of contact frequencies measured by chromosome conformation capture<sup>24</sup>, but how contact frequency is related to the size measurements here remains to be understood. In addition to different size-scaling properties, these different types of epigenetic domains also tend to have different 3D shape characteristics (Extended Data Fig. 3e, f).

Next, we probed how chromatin was folded within epigenetic domains. To this end, we selected two large chromatin domains for each epigenetic type and measured the  $R_g$  of internal regions of varying lengths within these domains, hereafter referred to as subdomains (Fig. 2a, b; Extended Data Fig. 5c; Extended Data Table). Interestingly, both *inactive* and *active* domains showed a self-similar organization, in which the internal subdomains exhibited scaling behaviours that were similar to those observed for the whole epigenetic

domains (Fig. 2b, left and middle). In stark contrast, we did not observe such a self-similar organization for either of the *repressed* chromatin domains investigated (the Bithorax (Fig. 2b, right) and Antennapedia (Extended Data Fig. 6) complexes). Instead, the  $R_g$  values grew rapidly as a function of subdomain length and quickly saturated such that subdomains longer than  $1/5^{\text{th}}$  of the length of the parent domain essentially all exhibited the same  $R_g$  values.

The observation that even a small subdomain traversed nearly the entire volume of the parent domain predicts that two such small subdomains would occupy the same physical space, suggesting a high degree of intermixing of chromatin within these *repressed* domains. We tested this hypothesis by simultaneously labelling two subdomains within the same repressed domain with two distinct sets of FISH probes conjugated to spectrally distinct photoswitchable dyes and imaged these subdomains with two-colour STORM (Fig. 2c, right panel). Indeed, images of these subdomain pairs showed a high degree of overlap. This highly intermixed state was markedly distinct from the behaviours observed for the active and *inactive* chromatin regions that we examined, which were characterized by only moderate levels of intermixing between subdomains (Fig. 2c; left and middle panels). Quantitatively, the pairs of subdomains within each repressed domain showed ~60-80% overlap in space (median 68%, 3 different pairs of subdomains investigated,  $n \approx 150$  cells) (Fig. 2d, light blue). In contrast, the neighbouring subdomains of *inactive* chromatin only showed ~10–30% spatial overlap (median 26%, 3 different pairs of subdomains,  $n \approx 150$ cells) (Fig. 2d, black) and the neighbouring subdomains of active chromatin only showed ~15–25% spatial overlap (median 20%, 2 different pairs of subdomains,  $n \approx 100$  cells) (Fig. 2d, red). The difference observed between active (or inactive) subdomains and repressed subdomains is statistically highly significant ( $p < 1 \times 10^{-10}$ , Wilcoxon test). These results indicate that the degree of intermixing of chromatin within individual epigenetic domains depends strongly on the epigenetic state.

We then probed how these different epigenetic domains interacted with one another across epigenetic boundaries. Notably, the *repressed* domains did not show any appreciable overlap with neighbouring *active* domains, whereas the neighbouring *inactive* and *active* domains appeared to be partially intermixed with each other (Fig. 3a, b). We quantified four different *repressed::active* boundaries and three different *inactive::active* boundaries. The *repressed* domains typically showed less than 3% overlap with their neighbouring *active* domains (median 1.5%,  $n \approx 150$  cells), whereas *inactive* domains exhibited up to 15% overlap with neighbouring *active* domains (median 9.8%,  $n \approx 150$ ) (Fig. 3c). The difference between these two types of domain boundaries was statistically highly significant ( $p < 1 \times 10^{-14}$ , Wilcoxon test). Therefore, the degree of spatial separation between neighbouring domains of different epigenetic types also depends strongly on the epigenetic state.

The different packaging and intermixing behaviour observed for these chromatin types point to distinct mechanisms involved in chromatin folding in the different epigenetic states. It is remarkable that the two types of non-transcribing chromatin — *inactive* and *repressed* states — exhibited such distinct packaging behaviours. The *repressed* chromatin appeared to exhibit a substantially more compact packing, higher degree of chromatin intermixing within domains, and stronger tendency to spatially exclude neighbouring domains. To explore the potential mechanisms underlying these different packaging behaviours, we employed

stochastic polymer simulations. It has been suggested that genomic DNA can be approximated as a "fractal globule", which arises when an unknotted polymer is confined into a volume substantially smaller than the relaxed volume of the polymer<sup>25–27</sup>. Our experimentally observed power-law scaling ( $R_g \sim L^{0.3}$ ) and self-similar organization of the *inactive* chromatin are roughly consistent with the expected properties of the fractal globule, as confirmed by our simulations of a polymer confined to a small volume (Supplementary Methods) (Fig. 4a, b).

Polycomb-repressed chromatin, on the other hand, exhibited distinct folding behaviours that could not be explained by the fractal globule model. Prompted by the observations that some PcG proteins can bridge nucleosomes<sup>21,22</sup>, we used self-interacting monomers<sup>28</sup> to simulate such PcG-mediated chromatin interactions. We then embedded such a "sticky" polymer domain containing self-interacting monomers within a large non-sticky polymer (to emulate the surrounding non-repressed domains), and simulated the polymer in a confined volume (Supplementary Methods). This model reproduced the packaging behaviours that we observed for repressed chromatin domains provided that the monomer-interaction strength was not too strong to cause irreversible monomer binding and that the simulation was sufficiently long to achieve intermixing (Supplementary Methods). First, the scaling exponent of  $0.22 \pm 0.01$  ( $R_g \sim L^{0.22}$ ) derived from the simulation agreed quantitatively with the experimentally observed value (Fig. 4c). Second, subdomains of these simulated polymers showed a saturation behaviour where the subdomains had nearly the same  $R_{g}$ values as the full domain until they became less than one-third the length of the parent domain (Fig. 4d), also similar to our experimental observation (Fig. 2b, right panel). Third, subdomains within the sticky polymer domains exhibited substantially more intermixing than neighbouring subdomains of the non-sticky polymer, and the sticky polymer domains spatially excluded neighbouring non-sticky regions (Fig. 4e, f) – both behaviours also consistent with the differences that we observed experimentally between repressed and inactive chromatin domains (Fig. 2c, d and Fig. 3b, c). Overall, our simulation results suggest that interactions between PcG proteins could provide an explanation for the distinct folding behaviour observed for the *repressed* chromatin, though it is possible that other mechanisms also contribute to the observed folding behaviour.

To test if these structural features for *repressed* domains indeed depend on PcG proteins, we used RNAi to knockdown a Polycomb Repressive Complex I (PRC1) component, Polyhomeotic (Ph), which contains a SAM motif that is capable of self-interaction<sup>29</sup>. qPCR experiments indicated a ~80% knockdown efficiency of the two copies of *ph, ph-p* and *ph-d* (Extended Data Fig. 7a). Functionally, Ph-knockdown led to a substantial increase in expression of many known PcG-target genes but did not affect the expression of the hundreds of genes from the *inactive* and *active* domains that we imaged (Extended Data Fig. 7a, b), consistent with previous work showing the important role of PcG proteins in transcriptional repression<sup>2,21–23</sup>. Structurally, Ph-knockdown abolished all of the three unique packaging properties that we observed for the *repressed* domains. First, Ph-knockdown caused substantially decompaction of the *repressed* domains to an extent that their 3D sizes approached those of *active* domains, and the power-law scaling coefficients of the *repressed* domains increased substantially such that the domains no longer became more densely packed with increasing domain length (Fig. 1c, d, hollow circles, Extended Data

Fig. 7c). Second, Ph-knockdown abolished the saturation effect that we observed for the subdomains of the *repressed* domains (Extended Data Fig. 7d). Finally, the spatial overlap between neighbouring *repressed* and *active* domains in Ph-knockdown cells increased dramatically to 20–30% (Fig. 3b, c), becoming similar to the extent of overlap observed between neighbouring active subdomains. The increase in intermixing of *repressed* and *active* domains upon Ph-knockdown is statistically highly significant (Wilcoxon test  $p = 1 \times 10^{-18}$ ,  $n \approx 150$  cells). These data indicate that the unique structural properties that we observed for the *repressed* domains indeed depend on PcG proteins, potentially through the self-interactions of the Ph component of PRC1.

Although we primarily focused on the relationship between chromatin structures and epigenetic states, our experimental data also revealed significant locus-specific variations in chromatin packaging, particularly for the *active* domains (Extended Data Fig. 8). After normalization for different domain lengths based on our observed scaling law, 42% of the pairs of examined *active* domains showed statistically distinct degrees of compaction (p < 0.05), while this fraction was only 16% for the examined *repressed* domains (Extended Data Fig. 8b). These variations may be linked to local genomic context, such as different activating histone modifications or transcription factor binding<sup>20</sup>, or different levels of transcriptional activity. Indeed, our analysis suggested that domains with higher transcriptional activity (measured by RNA-seq) appeared to have less dense packaging, as did domains with higher than average density of CTCF binding, though the degrees of correlation were only moderate (Extended Data Fig. 9).

Taken together, our results showed that direct visualization of DNA in the nucleus with nanometre-scale resolution can reveal previously unseen structural details of chromatin. We observed three distinct spatial organizations for chromatin in Drosophila cells, each corresponding to a different epigenetic state. Transcriptionally inactive chromatin, lacking substantial post-translational modification or transcription factor binding, adopts a 3D organization that resembles the fractal globule state of a polymer. On the other hand, Polycomb-repressed chromatin adopts a substantially more compact packaging configuration. Within these repressed domains, chromatin is highly intermixed, much more so than within the constitutively *inactive* chromatin. These Polycomb-repressed domains harbour genes encoding key developmental transcription factors, whose misexpression can have detrimental consequences in differentiated cells. Such a high-density packaging and high degree of chromatin intermixing may contribute to the inaccessibility of DNA to the transcription machinery, and thereby may help ensure that transcription is repressed in this chromatin state<sup>21,22</sup>. Moreover, our observation that Polycomb-repressed domains become increasingly more compact as the domain length increases could explain why PcG-targeted genes (such as the well-known Hox genes) tend to cluster in the genome. Concatenating PcG domains would increase domain length and therefore could lead to more compact packaging and potentially a more complete repression of gene expression. Notably, these Polycombrepressed domains also strongly exclude intermixing with nearby (active) chromatin, which may help prevent "bystander" activation<sup>30</sup> caused by transcription activators and enhancers accidentally brought into proximity of PcG-targeted genes by the active chromatin. We note that the Polycomb-repressed domains investigated here all have relatively high density of bound PcG proteins (Supplementary Methods); it is thus possible that the structural

properties that we observed may not apply to all Polycomb-enriched domains and that a sufficiently high PcG density is needed to establish these properties. Finally, we observed a far less compact packaging for transcriptionally active domains as compared to transcriptionally inactive domains; such an open structure could help transcription factors and polymerase machinery gain access to the encoded genes and facilitate transcription. Thus, our data suggest that epigenetic factors not only partition the genome into distinct, one-dimensional domains of different biochemical properties but can also shape the three-dimensional nanoscale structures of these domains, which may work in concert with the biochemical properties to regulate gene expression.

#### Extended Data



**Extended Data Figure 1. Schematic of oligonucleotide probe design and synthesis** A unique pair of index primers are used in a PCR reaction to selectively amplify the templates for the probe set of interest from a complex pool of custom, array-derived oligonucleotides. These templates are then amplified and converted to RNA in an *in vitro* transcription reaction. The RNA products are converted back to DNA in a reversetranscription reaction using a primer labelled with an activator dye, Alexa 405, which incorporates the dye into the resulting single-stranded DNA probe. Finally, a 32-nt

oligonucleotide attached to Alexa 647 is hybridized to all of the probes. The photoswitchable dye, Alexa 647, is used for STORM imaging. The activator dye, Alexa 405, facilitates the 405-nm light induced reactivation of the Alexa 647 dye.



#### Extended Data Figure 2. Effect of cell fixation on chromatin sizes

**a**, Top panels: Example images of DNA in a  $Kc_{167}$  cell, visualized with the viable DNA dye Hoechst 33342, both in the live cell before fixation and in the same cell after applying our fixation buffer (osmotically balanced methanol-free formaldehyde in PBS). Bottom panels: Same as the top panels but for a  $Kc_{167}$  cell before and after fixation with methanol, a fixative that is known to cause a shrinkage effect. **b**, Quantifications of the distances between chromatin features in live and fixed cells. Corresponding chromatin features were identified in the live and corresponding fixed cell images through scale-invariant feature transform

(SIFT) registration<sup>40</sup>. We measured distances between pairs of identified SIFT features in each cell, and calculated the ratio between the median inter-feature distances before and after fixation for each cell. Plotted here are the histograms of ratios determined from many cells for fixation with our osmotically balanced methanol-free formaldehyde fixation buffer (magenta), for a "mock fixed" condition in which the growth media was replaced with fresh media without any fixation reagent (cyan), and for fixation with methanol (red),  $n \approx 80$  cells in each cases. The average ratios are  $1.009 \pm 0.003$  and  $1.008 \pm 0.003$  for fixation with our fixation buffer and the mock fixation, respectively, indicating a lack of shrinkage effect. In contrast, the average ratio for the methanol-fixation case ( $0.868 \pm 0.005$ ) is appreciably less than one, indicating a chromosome shrinkage induced by methanol. c, STORM images of TRF1-mMaple3 labelled telomeres in live and fixed HEK293 cells. mMaple3 is a photoactivatable fluorescent protein<sup>42</sup>. Cells are fixed with our osmotically balanced methanol-free formaldehyde in PBS. Two examples of telomere STORM images are shown for each condition. d, Quantifications of the radius of gyration of the telomeric domains in live and fixed cells. We determined the radius of gyration for each telomere structure and plotted here are the histograms of the radii of gyration across ~150 telomeres from ~30 cells for live (Cyan) and fixed (Magenta) cells. The average radius of gyration is  $77 \pm 3$  nm for live cells and  $78 \pm 2$  nm for fixed cells, again indicating that there is no significant chromatin shrinkage effect upon fixation. The telomere size measurement is not limited by our image resolution with mMaple3 (~30 nm).

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Extended Data Figure 3. Volume, radius of gyration and other shape characteristics for chromatin domains of various domain lengths in three different epigenetic states a, Scheme of Drosophila chromosomes (X, 2L, 2R, 3L, and 3R) with the position of the imaged epigenetic domains marked (Red: *active* domains A-01 to A-23; Black: *inactive* domains I-01 to I-14; Blue: *repressed* domains R-01 to R-11). b, Log-log plot of the median domain volume as a function of domain contour length reproduced from Fig. 1c but with the domain ID labelled. c, As in Fig. 1c but plotted on a linear-linear scale. d, Linear plot of the median radius of gyration as a function of domain contour length. e, Coefficient of variation

(CV) in density per voxel for all domains as a function of domain length. CV in density is defined as the ratio of the standard deviation of density to the average density within the domain-occupied volume, which characterizes how uniformly the chromatin is distributed in space within these domains (Supplementary Methods). **f**, Ratio of surface area to volume<sup>2/3</sup> for all domains as a function of domain length. This surface-to-volume parameter characterizes the complexity of the physical shapes taken by the domains in 3D (Supplementary Methods). Error bars represent 95% confidence intervals derived from resampling.



### Extended Data Figure 4. Conventional images of chromatin domains and domain volume characterization based on conventional images

**a**, Blow-up view of the conventional images of chromatin domains shown in Fig. 1b. The left column shows the raw conventional, wide-field images, with pixel size defined by our camera. The right column shows the corresponding anti-aliased and de-noised images. **b**, Quantification of the median domain volume determined from conventional images (foreground symbols), overlaid on the median volume determined from STORM data plotted in Fig. 1c (faint background symbols and lines). Error bars represent 95% confidence intervals derived from resampling. Note that the conventional images may not only cause an artificial increase in domain size, especially severe for those domains whose physical sizes are smaller than the image resolution, but can also lead to an apparent decrease in domain size in some cases when the thin protrusions was too dim to detect by conventional imaging.

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Radius of gyration (nm)

# Extended Data Fig 5. Distributions of domain volume and radius of gyration of different epigenetic domains and subdomains over all imaged cells

a, Histograms of domain volume for all imaged cells for each of the domains shown in Extended Data Fig. 2a. Red: *active* domains; Black: *inactive* domains; Light blue: *repressed* domains. The domain IDs are indicated in the upper right corner of each plot. The x-axis (volume) range has been adjusted for each domain to ensure the readability of the histogram.
b, Histograms of the radius of gyration for each of the imaged domains in all cells. c, Histograms of the radius of gyration for subdomains of *active* (red), *inactive* (black) and *repressed* (blue) chromatin, shown in Fig. 2b and Extended Data Fig. 6, for all imaged cells. The subdomain IDs are indicated in the upper right corner of each plot.

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Extended Data Figure 6. Additional data on the scaling behaviour of subdomains of repressed chromatin

**a**, Enrichment profile of H3K4me2 (red), H3K27me3 (light blue) and unmodified H3 (black) in a genomic region harbouring the *repressed* domain R-11 (Antennapedia complex). **b**, The radius of gyration of subdomains (green triangles) of R-11 as a function of subdomain length compared to the scaling of whole *repressed* domains (light blue circles). The data shown in Fig. 2b, right panel are for the R-10 domain (Bithorax complex). The light blue dashed line indicates the power-law fit for the whole domain data. Green lines are to guide the eye. Error bars represent 95% confidence intervals ( $n \approx 50$  cells).

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#### Extended Data Figure 7. Effect of polyhomeotic (Ph) knockdown

**a**, Quantification of relative change in gene expression by qPCR (mean +/– s.e.m., n = 3 biological replicates) upon *ph-p* and *ph-d* double knockdown. Grey bars: Expression fold change of *ph-p* and *ph-d* upon the double knockdown. Light blue bars: Expression fold change of five Polycomb target genes, *Ubx, Abd-B, Dfd, Antp, en.* Red bars: Expression fold change of three control genes, *Act5c, alphatub84b* and *Gapdh1*, that are not targeted by Polycomb. Expression fold change was determined as the ratio between the signal detected in *ph-p* and *ph-d* double knockdown cells and that detected in wild-type cells. **b**, Average expression fold change upon *ph-p* and *ph-d* double knockdown for all genes in all of the *active* (red), *inactive* (black) and *repressed* (light blue) domains included in our study. The expression fold change is defined as the ratio of expression level measured in Ph-knockdown cells to that measured in the wild-type control cells determined by next generation RNA sequencing (mean ±s.e.m., n = 45, 89 and 532 genes for Repressed, Inactive and Active domains, respectively, 2 biological replicates). Expression level was measured in units of

read fragments per kilobase per million reads (FPKM). Note, some genes (9 from Repressed regions, 11 from Inactive regions and 2 from Active regions) are excluded from the average expression fold change calculation because they received zero counts in the wild-type control cells. **c**, Example images of the R-10 domain (Bithorax complex) in wild-type (left) and Ph-knockdown (right) cells. **d**, Radius of gyration vs. domain length for subdomains of R-10 in wild-type cells (solid green triangles) and Ph-knockdown cells (hollow green triangles).



Extended Data Figure 8. Locus-to-locus variation observed for the three types of epigenetic domains after normalization based on the observed scaling law over domain length

**a**, the normalized volume for domains of *active* (left), *inactive* (middle), and *repressed* (right) chromatin. Normalized volume is defined as the ratio of median volume of the domain to the expected volume calculated from the power-law scaling fits shown in Fig. 1c. Error bars represent 78% confidence intervals, such that there is a less than 5% chance that domains with non-overlapping error bars are not distinct. **b**, Volcano plots of the relative differences in volume between all pairs of *active* domains (left), *inactive* domains (middle) or *repressed* domains (right), after the normalization shown in (a). Each data point represents one pair of domains with their ratio of the normalized volumes plotted on the x-axis and the p-value of their normalized volume difference plotted on the y-axis. The dashed line is at a p-value of 0.05. All dots above this line represent pairs of domains in which the normalized volume of one domain is statistically distinguishable that of the other domain. **c**, Standard deviation of the normalized volumes for each domain type. Error-bars represent 95% confidence intervals.



Factor	Pearson Correlation Coefficient	p-value
TSS density	0.0	0.926
RNA-seq read density	0.6	0.008
Ave. gene expression	0.5	0.011
Max. gene expression	0.1	0.575
CTCF peak density	0.7	0.001
Su(HW) peak density	-0.4	0.069
BEAF32 peak density	-0.2	0.449

# Extended Data Figure 9. Additional factors correlating with the domain volume after normalizing the effect of domain length for *active* domains

To normalize for the effect of domain length, we determined the percent deviation of the volumes of *active* domains from the power law scaling trendline shown in Fig. 1c, and hereafter refer to this value as percent deviation from trendline. **a**, Correlation of the percent deviation from trendline with the binding density of the insulator proteins BEAF32 (left) and CTCF (right). Binding density was determined from the density of peaks per kb in Dam-ID data<sup>20</sup>. Peaks were defined as local maxima at least 2 standards deviation above the mean. **b**, As in (a) but for correlation with transcription start site (TSS) density (left) and RNA-seq total read density (right). The TSS density is defined as the average number of TSSs per kb in the domain and the RNA-seq total read density is defined as the total number of reads mapping to the domain measured using RNA sequencing divided by the domain length in kb. **c**, Pearson Correlation coefficients and corresponding p-values for the correlation of percent deviation from trendline with the indicated genomic factors. Average gene

expression is referring to the average expression value in FPKM (read fragments per kilobase per million reads) of all genes in the domain. Maximum gene expression is referring to the FPKM of the most highly expressed gene in the domain. Su(Hw) is an insulator protein like BEAF32 and CTCF. We noticed a weak trend in which domains with higher binding densities of the insulators BEAF32 or Su(Hw) are slightly more compact. Although this trend is consistent with the hypothesis that insulator proteins may function as loop forming factors and that loops may lead to more compact domains<sup>12</sup>, the correlation detected here was not statistically significant. Further analysis with improved sensitivity in detection of BEAF32 or Su(Hw) binding sites might uncover a stronger affect, so our data do not rule out the insulator loop hypothesis. Similarly, we caution that the positive correlation observed with the density of CTCF binding sites might reflect the preference of CTCF to bind open chromatin regions (such as enhancers and promoters), and does not necessarily suggest that CTCF binding induces a more open chromatin state.

# Extended Data Table 1

# Epigenetic domains and subdomains probed

List of the domain IDs, locus coordinates, lengths (in base pairs) and representative harboured genes of all epigenetic domains and subdomains imaged in this study.

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Doma	ins			Subdom	ains		
Ð	Genome coordinate	Length (bp)	Selected genes	Ð	Genome coordinate	Length (bp)	Selected genes
A-01	chr2L:2489241-2499536	10295	oaf	A-19.1	chrX:1907754-1977754	70000	(Many genes)
A-02	chr2R:7338881-7351765	12884	E(Pc)	A-19.2	chrX: 1977754-2047754	70000	(Many genes)
A-03	chr3R:12464802-12479673	14871	Cctgamma	A-19.3	chrX: 1907754-2047754	140000	(Many genes)
A-04	chr3R:2470393-2487998	17605	Taf1	A-22.1	chr3R:19929199-20029199	100000	(Many genes)
A-05	chr2L:15254575-15274520	19945	yuri	A-23.1	chr2R:19726615-19787585	02609	(Many genes)
A-06	chr3R:2890667-2916994	26327	alphaTub84b, Dpck, Aly	A-23.2	chr2R:19809874-19888410	78536	(Many genes)
A-07	chr2L:2210964-2237817	26853	mio, Uch, rti,	A-23.3	chr2R:19906491-19976552	70061	(Many genes)
A-08	chr2L:21153308-21183571	30263	Acon, bur, Mpp6, Nbr, Mcm10, Ret	A-23.4	chr2R:20000481-20092780	92299	(Many genes)
A-09	chr2L:5517877-5549514	31637	Lam, Cap-D3, tomb, Oscillin, Hel25E, cl	A-23.5	chr2R:19726615-19888410	161795	(Many genes)
A-10	chr2L:8353587-8391384	37797	Aats-ala, RpS13, CSNS, Pp2A-29B, Rcd4, Mur29b	A-23.6	chr2R:19809874-20092780	282906	(Many genes)
A-11	chr2L:13363559-13411277	47718	Vm34Ca.RpL24, loqs. beta'Cop, Tap42.Nnp-1	I-13.1	chr3R:1786654-1886654	100000	Pbprp3,Os-E
A-12	chr2R:4972056-5054981	82925	(Many genes)	I-13.2	chr3R:1686653-1786653	100000	(Contains only genes of little known function)
A-13	chr2R:13979742-14072371	92629	(Many genes)	I-14.1	chr2R:15602615-15702615	100000	Or56a,Obp56g
A-14	chr3L:167832-265882	98050	(Many genes)	I-14.2	chr2R:15702616-15802615	66666	Obp56h,Toll-7,Obp56i
A-15	chr2R:3328439-3429969	101530	(Many genes)	I-14.3	chr2R:15802616-15852615	49999	(Contains only genes of little known function)
A-16	chr2R:9019018-9125165	106147	(Many genes)	I-14.4	chr2R:15852616-15952615	66666	(Contains only genes of little known function)
A-17	chr3R:12811417-12951671	140254	(Many genes)	I-14.5	chr2R:15952616-16128463	175847	18w
A-18	chr3L:9344016-9505324	161308	(Many genes)	I-14.6	chr2R:15702616-15852615	149999	Obp56h,Toll-7,Obp56i
A-19	chrX:1907754-2083414	175660	(Many genes)	I-14.7	chr2R:15802616-15952615	149999	(Contains only genes of little known

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Doma	ins			Subdom	ins			
A	Genome coordinate	Length (bp)	Selected genes	Ð	Genome coordinate	Length (bp)	Selected genes	
							function)	_
A-20	chrX:16161627-16351359	189732	(Many genes)	I-14.8	chr2R:15852616-16128463	275847	18w	
A-21	chr2L:10203488-10443191	239703	(Many genes)	R-06.1	chr3R:3967588-4043731	76143	grn	_
A-22	chr3R:19929199-20187132	257933	(Many genes)	R-10.1	chr3R:12571412-12586412	15000	(Ubx regulatory region)	_
A-23	chr2R:19726615-20092780	366165	(Many genes)	R-10.2	chr3R:12571412-12616414	45002	(Ubx regulatory region)	_
I-01	chr2L:455203-465638	10435	(Contains only genes of little known function)	R-10.3	chr3R:12481406-12619389	137983	Ubx	
I-02	chr2L:18991061-19001768	10707	(Contains only genes of little known function)	R-10.4	chr3R:12616227-12683098	66871	abd-A	
I-03	chr3L:1250391-1295624	45233	(Contains only genes of little known function)	R-10.5	chr3R:12683230-12810708	127478	Abd-B	
I-04	chr3R:14754844-14804163	49319	(Containts only genes of little known function)	R-10.6	chr3R:12481406-12683098	201692	Ubx, abd-A	
I-05	chr3L:7154799-7231065	76266	(Contains only genes of little known function)	R-11.1	chr3R:2487143-2570646	83503	zen2, pb, lab	
I-06	chr3L:18903046-18983014	79968	(Contains only genes of little known function)	R-11.2	chr3R:2568667-2653159	84492	Dfd, bcd, zen, zen2	
I-07	chrX:16351360-16440669	89309	para(DmNav1)	R-11.3	chr3R:2641835-2721984	80149	Scr, ftz	_
I-08	chr3R:12312771-12438748	125977	Scp2, Decay, Gr89a, npf	R-11.4	chr3R:2721063-2827254	106191	Antp	
60-I	chrX:6269580-6417829	148249	(Contains only genes of little known function)	R-11.5	chr3R:2825712-2889707	63995	Sodh-1, (Antp regulatory region)	
I-10	chr3R:2287031-2473421	186390	sunz, glob2, dpR-11	R-11.6	chr3R:2487143-2653159	166016	Scr, Dfd, bcd, zen, zen2, pb, lab	
I-11	chr3R:12977555-13217431	239876	beat-IIa, beat-IIb, Brf	R-11.7	chr3R:2568667-2827254	258587	Antp, ftz, Scr, Dfd, bcd, zen, zen2	
I-12	chrX:9679192-10103054	423862	Yp1, Yp2, Cht6, Ser7, Idgf4					
I-13	chr3R:1686653-2165126	478473	Or83c, Os-E, Osi3-Osi19, Pbprp3, Obp83cd, Obp83ef, Obp83g					
I-14	chr2R:15602615-16128463	525848	18w, Toll-7, Obp56h,Obp56i					
R-01	chr2R:5080631-5092410	11779	Bdun					
R-02	chr2R:5860081-5876517	16436	eve					
R-03	chr2R:8770090-8803242	33152	٨g					
R-04	chr3R:11833445-11870730	37285	pni, GATAe					
R-05	chr2R:1593366-1640664	47298	ap					_

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ID         Genome coordinate         Length (bp)         Selected genes         ID         Genome coordinate           R-06         chr3R:3953538-4054361         1008233         gm         Genome coordinate         ID         Genome coordinate           R-07         chr3R:3953538-4054361         100823         gm         gm         Figure coordinate         Figure coordinate	Don	ains			Subdom	ains		
R-06chr3R:3953538-4054361100823gmR-07chr2R:7350135-7469804119669en, invR-08chr3R:17205633-17399201193568slou, lbe, lbl, bapR-09chr3R:17205633-17399201193568slou, lbe, lbl, bapR-10chr3R:120553918-16160086206168disco, disco-r, TobR-10chr3R:12481406-12810708329302Abd-B, abd-A, Ubx (Bithorax-complex)R-11chr3R:12481405-12810708402564Antp.fiz, Scr. Dfd, bcd, zen, zen2, pb,	Ð	Genome coordinate	Length (bp)	Selected genes	ID	Genome coordinate	Length (bp)	Selected genes
R-07       chr2R:7350135-7469804       119669       en, inv         R-08       chr3R:17205633-17399201       193568       slou, lbe, lbl, bap         R-09       chrX:15953918-16160086       206168       disco, disco-r, Tob         R-10       chr3R:12481406-12810708       329302       Abd-B, abd-A, Ubx (Bithorax-complex)         R-11       chr3R:2487143-2889707       402564       Antp, fiz, Scr, Dfd, bcd, zen, zen2, pb,	R-0	5 chr3R:3953538-4054361	100823	gm				
R-08         chr38:17205633-17399201         193568         slou, lbe, lbl, bap           R-09         chrX:15953918-16160086         206168         disco, disco-r, Tob           R-10         chr38:12481406-12810708         329302         Abd-B, abd-A, Ubx (Bithorax-complex)           R-11         chr38:2487143-2889707         402564         Antp.fiz, Scr. Dfd, bcd, zen, zen2, pb,	R-0	7 chr2R:7350135-7469804	119669	en, inv				
R-09         chrX:15953918-16160086         206168         disco, disco-r, Tob           R-10         chr3R:12481406-12810708         329302         Abd-B, abd-A, Ubx (Bithorax-complex)           R-11         chr3R:2487143-2889707         402564         Antp, fiz, Scr, Dfd, bcd, zen, zen2, pb,	R-0;	s chr3R:17205633-17399201	193568	slou, lbe, lbl, bap				
R-10         chr3R:12481406-12810708         329302         Abd-B, abd-A, Ubx (Bithorax-complex)           R-11         chr3R:2487143-2889707         402564         Antp, fiz, Scr. Dfd, bcd, zen, zen2, pb,	R-0	) chrX:15953918-16160086	206168	disco, disco-r, Tob				
R-11 chr3R:2487143-2889707 402564 Antp. fiz. Scr. Dfd. bcd, zen, zen2, pb.	R-1(	) chr3R:12481406-12810708	329302	Abd-B, abd-A, Ubx (Bithorax-complex)				
Iao (Amenopedia-comprex)	R-1.	l chr3R:2487143-2889707	402564	Antp, fitz, Scr, Dfd, bcd, zen, zen2, pb, lab (Antenopedia-complex)				

#### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

#### Acknowledgments

We thank Hazen Babcock for help with instrumentation and technical advice, Jané Kondev for insightful discussions about the data, Anton Goloborodko for sharing custom OpenMM force-field, and Sonny Nguyen for discussions on RNAi knockdown protocol. This work was supported in part by the National Institutes of Health (X.Z., C.W., L.M.). A.N.B. acknowledges support by the Damon Runyon Foundation postdoctoral fellowship. J.R.M. acknowledges support of the Helen Hay Whitney Foundation postdoctoral fellowship. S. W. acknowledges support of the Jane Coffin Childs Foundation postdoctoral fellowship. X.Z. is a Howard Hughes Medical Institute Investigator.

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Figure 1. Chromatin in different epigenetic states exhibits distinct packaging and power-law scaling

**a**, Enrichment profile of H3K4me2 (red), H3K27me3 (light blue) and unmodified H3 (black) in three genomic regions, each harbouring an example *active, inactive* or *repressed* domain (indicated by brackets). Marker enrichment, as defined in Supplementary Methods, was determined from ChIP-seq data<sup>20</sup>. **b**, 3D-STORM images of the three distinct epigenetic domains in (a), labelled by *in situ* hybridization with DNA probes conjugated to the photoswitchable dye Alexa-647, shown with their corresponding conventional images in the

inset. Each epigenetic domain appears as a single region in nearly all cells due to homologous pairing in the tetraploid Kc<sub>167</sub> cells. **c**, Log-log plot of the median domain volume as a function of domain length for *active* (red solid circles), *inactive* (black solid circles) and *repressed* (light blue solid circles) domains, as well as for *repressed* domains in Ph-knockdown cells (light blue hollow circles). Error bars represent 95% confidence intervals derived from resampling ( $n \approx 50$  cells). The lines indicate power-law fits, with the scaling exponent *b* shown in the legend. **d**, as in (c) but for the radius of gyration as a function of domain length with the scaling exponent *c* shown in the legend.

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## Figure 2. Different types of epigenetic domains exhibit distinct subdomain scaling and intermixing

**a**, Marker enrichment profile of three genomic regions with the example epigenetic domains marked by brackets and imaged subdomains marked by green and magenta lines. **b**, Linear plot of the radius of gyration as a function of the subdomain length (green symbols), compared to those for the whole domain data (red, black or light blue circles), for *active* (left panel), *inactive* (middle panel) and *repressed* chromatin (right panel). Different green symbols (triangle and squares) represent subdomains of two different parent domains.

Power-law fits of subdomains (green solid lines) and whole domains (red, black and light blue dashed lines) are shown with the scaling exponent *c* given in the legends. The green lines in the right panel are to guide the eye. **c**, Two-colour, 3D-STORM images of example pairs of subdomains within *active* (left), *inactive* (middle) and *repressed* (right) domains. Portions of the two subdomains that overlap in 3D are shown in white. The two subdomains are labelled with Alexa-647 and Alexa 750 tagged DNA probes, respectively. **d**, Quantification of overlap fraction between the subdomains for *active* (red), *inactive* (black), or *repressed* (light blue) chromatin (Supplementary Methods). Error bars represent 95% confidence intervals derived from resampling ( $n \approx 50$  cells).

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# Figure 3. Neighbouring chromatin domains show different amount of intermixing for different types of epigenetic boundaries

**a**, Marker enrichment profile of a genomic region harbouring three epigenetic domains marked by the magenta and green lines. **b**, Two-colour, 3D-STORM images for neighbouring epigenetic domains with *repressed::active* and *inactive::active* boundaries in wildtype cells, as well as for the *repressed::active* boundaries in Ph-knockdown (Ph KD) cells. **c**, Quantification overlap fraction between the indicated neighbouring domains (Supplementary Methods). Error bars represent 95% confidence intervals derived from resampling ( $n \approx 50$  cells).

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Figure 4. Computational modelling of chromatin packaging for *inactive* and *repressed* domains **a**, Radius of gyration of polymer domains as a function of the domain length for simulated polymers confined in a small volume to emulate *inactive* chromatin. Error bars indicate 95% confidence intervals derived from resampling ( $n \approx 20$  simulations). The line indicates the power-law fit with the scaling exponent c = 0.33. **b**, Radius of gyration of subdomains of a parent *inactive* domain (green triangles) compared to the whole-domain scaling data (black circles) with power-law fits and scaling exponents shown. **c**, Radius of gyration as a function of the domain length for the simulated sticky polymer domain, embedded in a non-stick,

confined polymer, to emulate *repressed* chromatin. The light blue line represent the powerlaw fit with the scaling exponent c = 0.22. The black dotted line indicates the point at which the closest possible packing of monomers is reached, causing c to deviate from 0.22 and approach 0.33 beyond this point. c also deviates from 0.22 at the small monomer number end because such a short polymer chain cannot sufficiently bend. **d**, Radius of gyration of the subdomains (green triangles) of the parent sticky domain in comparison with the wholedomain scaling data (light blue circles). **e**, Snapshots of simulations showing adjacent subdomains in the *inactive* (non-sticky) chromatin, adjacent subdomains of the *repressed* (sticky) chromatin, and adjacent *repressed* and *inactive* chromatin domains. **f**, Quantification of the overlap fraction (Supplementary Methods) between the adjacent polymer regions illustrated in (e).