1 Live-cell 3D single-molecule tracking reveals how NuRD modulates enhancer dynamics

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31 Abstract

32 Enhancer-promoter dynamics are crucial for the spatiotemporal control of gene expression, but it remains 33 unclear whether these dynamics are controlled by chromatin regulators, such as the nucleosome remodelling 34 and deacetylase (NuRD) complex. The NuRD complex binds to all active enhancers to modulate transcription 35 and here we use Hi-C experiments to show that it blurs TAD boundaries and increases the proximity of 36 intermediate-range (~1 Mb) genomic sequences and enhancer-promoter interactions. To understand whether 37 NuRD alters the dynamics of 3D genome structure, we developed an approach to segment and extract key 38 biophysical parameters from 3D single-molecule trajectories of the NuRD complex determined using live-cell 39 imaging. Unexpectedly, this revealed that the intact NuRD complex decompacts chromatin structure and 40 makes NuRD-bound enhancers move faster, increasing the overall volume of the nucleus that these key 41 regulatory regions explore. Interestingly, we also uncovered a rare fast-diffusing state of NuRD bound 42 enhancers that exhibits directed motion. The NuRD complex reduces the amount of time that enhancers 43 remain in this fast-diffusing state, which we propose might otherwise re-organise enhancer-promoter 44 proximity. Thus, we uncover an intimate connection between a chromatin remodeller and the spatial dynamics 45 of the local regions of the genome to which it binds.

46 Introduction

3D genome organisation and chromatin dynamics are thought to be crucial for the spatiotemporal control of gene expression. However, little is known about the multi-scale chromatin dynamics of cis-regulatory elements or how they relate to genome organisation. To probe the dynamics and organisation of these elements at a single-cell level two complementary methods are typically used – either live-cell imaging¹⁻⁵ or single nucleus versions of chromosome conformation capture experiments⁶⁻¹³ (such as Hi-C), which reveal snapshots of the structure of a dynamic 3D genome in different individual fixed cells⁶⁻¹⁷. Whether chromatin regulators modulate these dynamics or how these dynamics relate to genome structure remains unclear.

54 The nucleosome remodelling and deacetylase (NuRD) complex is a highly conserved 1 MDa multisubunit protein complex which binds to all active enhancers¹⁸. NuRD combines two key enzymatic activities – 55 56 nucleosome remodelling via its helicase-containing ATPase (predominantly CHD4 in embryonic stem (ES) cells and lysine deacetylation via its HDAC1/2 subunits¹⁷⁻²³: these activities are thought to be present in two 57 58 sub-complexes. HDAC1/2 associate, along with the histone chaperones RBBP4/7, with the core scaffold proteins MTA1/2/3 to form a stable sub-complex with deacetylase activity²⁴. The nucleosome remodeller 59 60 CHD4 interacts with chromatin by itself and also forms a second sub-complex with GATAD2A/B and DOC1 (CDK2AP1)²⁴⁻²⁶. The methyl-CpG DNA binding domain proteins MBD2/3 interact directly with both the 61 deacetylase sub-complex and GATAD2A/B^{24,27-29}, and thus play a critical role in linking the CHD4 62 63 remodeller and HDAC sub-complexes together to assemble the intact holo-NuRD complex (Figure 1a).

Assembly of the intact holo-NuRD complex is critical for controlling cell fate transitions. Knockout of *Mbd3*, which disrupts intact NuRD complex assembly, leads to only moderate up- or down-regulation ('fine-tuning') of transcription levels, but this modulation prevents ES cell lineage commitment³⁰⁻³³. Nucleosome remodelling by NuRD regulates transcription factor and RNA polymerase II binding at active enhancers¹⁸, but whether this impacts enhancer dynamics or enhancer-promoter interactions has remained unclear. Here, to understand whether enhancer dynamics are regulated by this crucial chromatin remodeller, we combine Hi-C and live-cell single-molecule tracking. Specifically, we exploit the ability to unlink the

71 chromatin remodelling and deacetylase subunits of intact NuRD by deleting *Mbd3* to explore the function of

the intact NuRD complex.

73 NuRD complex assembly increases contacts between intermediate-range genomic sequences

74 We initially set out to understand whether assembly of the intact NuRD complex, mediated by MBD3, might 75 affect genome architecture by carrying out chromosome conformation capture (in-nucleus Hi-C) experiments 76 on wild-type and previously characterised³¹ *Mbd3* knockout (*Mbd3-ko*) ES cells. This strategy, which exploits 77 the fact that MBD2 (and thus the MBD2-linked holo-NuRD complex) is expressed at low levels in ES cells 78 and cannot rescue the *Mbd3* deletion^{34,35}, allowed us to specifically perturb NuRD complex function. We 79 obtained high-quality Hi-C contact maps for both wild-type and Mbd3-ko ES cells after combining our wildtype data with previously published³⁶ (and consistent) datasets (**Extended Data Figure 1a**). As previously 80 81 observed^{6,8-17,37}, these contact maps showed that the genome is segregated into: 1) A and B compartments 82 (broadly regions containing more or fewer genes respectively); 2) megabase-scale topologically associating 83 domains (TADs), which have a higher frequency of intra-domain chromatin interactions; and 3) loops where 84 specific genomic regions contact each other, mediated e.g. via CTCF/Cohesin binding.

85 Comparison of the Mbd3-ko and wild-type Hi-C data showed that assembly of the intact NuRD 86 complex leads to a ~30 % genome-wide increase in the probability of intermediate-range contacts at the scale of TADs (500 kb to 3 Mb) ($p < 10^{-17}$, Figure 1b, Extended Data Figure 1b). We found that ~77% and 17% 87 of NuRD-regulated genes³⁸ remain in the A and B compartments, respectively, with only a small proportion 88 89 (~3%) moving from the A to the B compartment or vice versa. The genome-wide increase in mean contact 90 length/region binned (Extended Data Figure 1c) was most noticeable for regions containing NuRD-regulated 91 genes that are within the A compartment in both *Mbd3-ko* and wild-type cells (KO-A, WT-A; see **Extended** 92 Data Figure 1d). We also found that promoters that are down-regulated in the presence of intact NuRD are 93 enriched (chi-squared p-value: < 1e-10, odds-ratio: 1.16) amongst genes that remain in the A compartment 94 and significantly depleted (chi-squared p-value: < 1e-10, odds-ratio: 0.43) amongst those that remain within 95 the B compartment. Further classification showed that promoters displaying significant MBD3 (i.e. NuRD) 96 ChIP-seq signal [which our unpublished experiments (RR, NR & BDH) suggest represents association with a 97 NuRD-bound enhancer] are also significantly enriched amongst genes that remain in the A compartment (p < p

98 10^{-10} , odds-ratio: 1.27) and significantly depleted amongst those that remain within the B compartment (p <

99 10⁻¹⁰, odds-ratio: 0.41) (Extended Data Figures 1e,f). Thus, NuRD-bound enhancers tend to contact

100 promoters to down-regulate genes in the A (euchromatic) compartment.

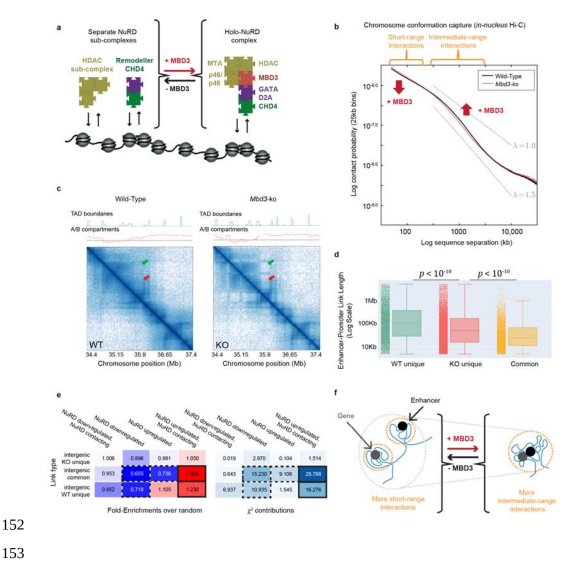
101 Comparison of the Hi-C contact maps for *Mbd3-ko* and wild-type cells showed that some boundaries 102 between TAD's and A/B compartments are weaker in cells containing the intact NuRD complex (Figure 1c 103 and Extended Data Figure 1g), with promoters that contact NuRD-bound enhancers having an increased 104 number of cross-compartment contacts (Extended Data Figure 1h) thereby decreasing TAD insulation 105 (Extended Data Figure 1i). Although we could not detect a significant correlation between the A/B 106 compartmentalisation of a gene and levels of transcription (Extended Data Figure 1j), these results show that NuRD-regulated genes tend to be at A/B compartment boundaries^{39,40}. The blurring of TAD and A/B 107 108 compartment boundaries suggested that NuRD might alter CTCF/Cohesin binding^{40,41}. We therefore carried 109 out Cut&Run experiments to study chromatin binding of CTCF and SMC3 (a subunit of the Cohesin complex) – which play a key role in loop and TAD formation^{13,42-47}. On assembly of the intact NuRD complex 110 111 we found there was a redistribution of both CTCF and SMC3 binding - the effect on SMC3 being more 112 pronounced – with both CTCF and SMC3 peaks appearing and disappearing (Extended Data Figure 2b; see 113 also Extended Data Figure 2e for an example gene). We found that NuRD-regulated genes are more likely to 114 be found near to CTCF/Cohesin binding sites that form when the intact NuRD complex assembles – this was 115 more noticeable for genes that were upregulated in wild-type vs Mbd3-ko cells and whose promoters contact 116 NuRD bound enhancers (Extended Data Figure 2d). Interestingly, we also found that genes that are 117 regulated in a similar way by NuRD (i.e. up or down) tend to cluster with each other – see for example the 118 *Htral* gene, which is found in a region with other significantly downregulated genes (Extended Data Figure 119 2e).

Because changes in TADs and CTCF-Cohesin loops are thought to influence enhancer-promoter proximity^{45,48,49}, we used a modified version of the activity-by-contact algorithm⁵⁰ to determine changes in enhancer-promoter interactions in *Mbd3-ko* vs wild-type cells (**Extended Data Figures 3a,b**). Using active enhancers and promoters defined using H3K27ac and H3K4me3 ChIP-seq profiles¹⁸ (see Online Methods), this analysis revealed that enhancer-promoter interactions that only occur in the presence of the intact NuRD complex tend to link together genomic regions separated by longer distances than those that occur in the

- 126 absence of MBD3 (Figure 1d). In addition, when we considered enhancer-promoter interactions and
- 127 transcription levels in wild-type compared to *Mbd3-ko* cells³⁸, we found more contacts between intergenic
- 128 NuRD bound enhancers and promoters that are upregulated by NuRD, and fewer contacts between intergenic
- 129 enhancers and promoters that are downregulated by NuRD (Figure 1e).

130 Figure 1. MBD3-dependent assembly of the NuRD complex increases Mb-range genome interactions.

131 (a) Schematic representation of the NuRD complex interacting with chromatin in the presence and absence of 132 MBD3. (b) Log-log plots of contact probability as a function of genomic sequence separation (averaged 133 across the genome), derived from *in-nucleus* Hi-C experiments of *Mbd3*-ko (red) and wild-type (black) ES 134 cells, shows a significant (30%) increase in intermediate-range (~1 Mb) contacts in wild-type cells ($p<10^{-17}$, 135 Wilcoxon rank sum test, see Extended Data Figure 1b). (c) Representative plots showing part of the Hi-C 136 contact map for Chromosome 1 – the density of contacts is indicated by the colour intensity. Some TAD 137 boundaries are weakened resulting in an increased density of contacts between adjacent TADs, both within 138 and between A/B compartments (red and green arrows) - see also Extended Data Figures 1g-i for genome-139 wide comparisons. (d) Boxplots showing intra-chromosomal enhancer-promoter link lengths, determined 140 using a modified version of the activity-by-contact algorithm⁵⁰, present in both *Mbd3*-ko and wild-type ES 141 cells (orange), in only *Mbd3*-ko cells (red) or in only wild-type cells (green). The bars on the left of each plot 142 show all the data. (e) (Left) Fold-enrichment and (Right) chi-squared test when intergenic enhancer-promoter 143 interactions found in both wild-type and *Mbd3*-ko cells, as well as those found uniquely in either the *Mbd3*-ko 144 or the wild-type, are correlated with genes that are up-/down-regulated in the presence of intact NuRD. 145 Interactions where NuRD bound enhancers can be seen to contact the promoter, that are found in both the 146 *Mbd3*-ko and the wild-type, or uniquely in the wild-type, are enriched at up-regulated genes. In contrast, there 147 is a depletion in interactions between intergenic enhancers and the promoter at down-regulated genes. 148 Enriched and depleted interactions are coloured red and blue, respectively, and highlighted using solid and 149 dashed black boxes – see also Extended Data Figure 3c for an example of changes in enhancer-promoter 150 contacts. (f) Schematic interpretation of the results of the Hi-C experiments, showing that in the Mbd3 151 knockout there are more short range and fewer intermediate range chromatin interactions.



153



155 chromatin movement

156 To provide a mechanistic understanding of how assembly of the intact NuRD complex might increase mixing 157 of intermediate-range genome sequences, we carried out live-cell 3D single-molecule tracking of NuRD 158 complex subunits in the presence and absence of MBD3. We generated knock-in ES cell lines expressing the 159 endogenous Chd4, Mbd3 and Mta2 genes fused with C-terminal HaloTags and confirmed that the tags did not 160 prevent NuRD complex assembly, although subtle changes in expression were observed (Extended Data 161 Figure 4). We used a double-helix point spread function microscope²⁵ to generate 3D tracks of single NuRD-162 HaloTag-JF₅₄₉ complexes through a 4 µm slice of the nucleus, so that we could reliably determine biophysical 163 parameters (Figure 2a). (Tracking in 2D reduces the length of trajectories because molecules can move in/out of the imaging plane during the experiment.) We recorded trajectories at two distinct temporal regimes, 20 ms
and 500 ms. Recording at 20 ms time resolution allows the detection of both freely diffusing and chromatin
bound proteins²⁵, and can thus be used to extract the chromatin binding kinetics of NuRD complexes (Figure
2b). In contrast, at a 500 ms time resolution, 'motion blurring' substantially reduces the detection of freely
diffusing molecules, allowing us to focus on the slower sub-diffusive chromatin bound NuRD complexes¹
(Figure 2c).

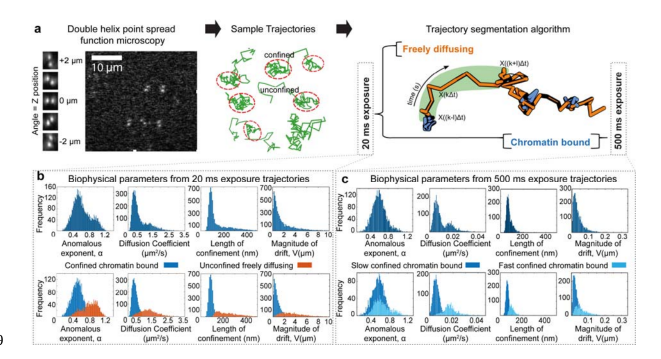
170 To extract dynamic biophysical parameters, we developed a machine learning method (a Gaussian 171 mixture model) to segment the single molecule trajectories into different classes by studying their behaviour 172 over a sliding window of 11 consecutive images (see Figure 2 and Extended Data Figure 5). To understand 173 how NuRD affects chromatin structure, we estimated from each sub-trajectory not just the apparent diffusion coefficient (as previously used for classifying sub-trajectories^{51,52}) but also the anomalous exponent α , the 174 175 localisation length Lc, and the drift magnitude V^{53} . The anomalous exponent α , defined as [mean squared 176 displacement $\propto t^{\alpha}$], is particularly informative. Diffusing proteins are characterised by an anomalous exponent close to 1 whereas chromatin bound proteins exhibit a lower anomalous exponent^{3,53,54}, whose value 177 is representative of the condensation state of the chromatin^{53,55}. In addition, values of α that are higher than 1 178 179 can represent energy-dependent directed motion. The localisation length Lc of chromatin bound proteins is 180 also informative as it reflects the spatial scale that the molecules explore within the nucleus. The localisation 181 length Lc is dependent on a range of parameters such as nucleosome density and CTCF/Cohesin-mediated 182 looping interactions: for example, Lc is larger at lower nucleosome densities when the linker length between 183 nucleosomes is longer^{55,56}. Finally, by computing the magnitude V_i (k) of the drift vector V_i in three 184 dimensions we can characterise directed movement of a molecule. (Further details of the approach – and of 185 the simulations we carried out to test the algorithm – are described in the Online Methods.)

Analysis of the 20 ms exposure trajectories of single CHD4 molecules using our approach revealed two diffusion states (**Figure 2b**). We identified a fast unconfined state that was freely diffusing with an α of 0.94 ± 0.12 and an apparent diffusion coefficient of 1.3 ± 0.3 μ m²/s (matching previous observations^{24,25}), as well as a confined chromatin bound state characterised by sub-diffusive motion with an α of 0.51 ± 0.02 and

190 an apparent diffusion coefficient of 0.43 \pm 0.03 μ m²/s. Similar results were obtained when segmenting the 191 trajectories of two other NuRD complex components, MBD3 and MTA2 (Extended Data Figure 6c). To 192 demonstrate that our approach can reliably determine relative differences in anomalous exponent for freely 193 diffusing and sub-diffusive chromatin bound molecules we imaged freely diffusing HaloTag. We found that only a small proportion of HaloTag molecules bind to chromatin (as observed previously^{57,58}), and that many 194 195 molecules have an anomalous exponent of around 1 [significantly higher than observed for fixed dye 196 molecules (α of 0.62 ± 0.01) (Extended Data Figure 6c)]. We conclude that we can use 20 ms exposure 197 trajectories to distinguish unconfined freely diffusing molecules from confined chromatin bound proteins. 198 [We note, however, that the apparent diffusion coefficient of chromatin bound NuRD molecules (0.43 ± 0.03) 199 μ m²/s) cannot be distinguished from immobile when imaging using 20 ms exposures: we found that immobile 200 dye molecules have a median precision of 60 nm and an apparent diffusion coefficient of $0.3 \pm 0.2 \ \mu m^2/s$ 201 (Extended Data Figure 6c).]

202

203 Figure 2. Live cell imaging to study NuRD complex binding kinetics and function. (a) (Left) Single JF_{549} -204 HaloTagged CHD4 molecules in the NuRD complex are tracked in 3D using a double-helix point spread 205 function microscope – two puncta are recorded for each fluorophore with their midpoint providing the lateral 206 x, y position and the angle between them representing the axial position in z relative to the nominal focal 207 plane. (Middle) Examples of extracted single particle trajectories from 20 ms exposure imaging show periods 208 of free unconfined and confined diffusion. (Right) Example trajectory of a single CHD4 molecule from 20 ms 209 exposure imaging showing a sliding window of 11 frames (shaded green) from which the biophysical 210 parameters are calculated. The resulting segmentation, into confined chromatin bound (blue) and unconfined 211 freely diffusing (orange) sub-trajectories, has been used to colour the trajectory. 500 ms exposure imaging, on 212 the other hand, can be used to extract slow and fast sub-diffusive motion of chromatin bound CHD4 213 molecules. (b, c) Distribution of the four biophysical parameters extracted from sliding windows within the 214 single-molecule trajectories of CHD4 taken with either (b) 20 ms or (c) 500 ms exposures - (top) before and 215 (bottom) after classification based on the anomalous exponent α , the apparent diffusion coefficient D, the 216 length of confinement Lc, and the drift magnitude, norm $\|V\|$ of the mean velocity. Each dataset was produced 217 by imaging 30 cells, and the parameters were extracted from 5,557 and 15,528 trajectories, respectively (see 218 Extended Data Figure 5 and the Online Methods for more detail).



220 Chromatin association of the deacetylase sub-complex requires assembly into the intact holo-NuRD

221 complex in ES cells

Having developed an approach to segment the 20 ms exposure trajectories of the NuRD complex into chromatin bound and freely diffusing molecules (**Figure 3a,b**), we investigated how the chromatin binding kinetics were affected by removal of MBD3, which disrupts the interaction between the HDAC- and CHD4containing NuRD sub-complexes^{18,24} (**Figure 1a**).

226 To explore whether the two sub-complexes are pre-assembled before binding to chromatin, we 227 imaged the CHD4 remodeller and the HDAC-containing sub-complex (using tagged MTA2 molecules) in 228 wild-type and Mbd3-ko ES cells. Single-molecule tracking of CHD4 in live ES cells revealed a small but 229 significant increase in the apparent diffusion coefficient of freely diffusing CHD4 in the absence of MBD3 230 (1.05-fold, p = 0.009), consistent with the formation of smaller CHD4 sub-complexes^{18,24,25} (Figure 3c). [A 231 larger increase might have been expected to arise from the disassembly of the holo-NuRD complex, but ES cells also contain CHD4 that is not present in NuRD (e.g. in the ChAHP complex with ADNP and HP1 β , γ^{59}). 232 233 When we imaged the HDAC-containing sub-complex using tagged MTA2 molecules, this revealed a much 234 more substantial increase in diffusion coefficient for freely diffusing MTA2 in the absence of MBD3 (1.7-235 fold, $p < 10^{-5}$; Figure 3c), demonstrating that the deacetylase subcomplex is normally associated with CHD4 236 in the intact holo-NuRD complex (Figure 1a). [As a control, we also imaged tagged MBD3 and showed that 237 both freely diffusing MBD3 and MTA2 molecules have similar diffusion coefficients, consistent with these 238 two proteins associating with each other in the deacetylase sub-complex^{18,24,25} (Figure 3c).] Finally, we 239 showed that MBD3 does indeed interact with CHD4 via GATAD2A, both in vitro using purified GATAD2A 240 in pull-down reconstitution experiments (**Extended Data Figure 6b**), and in ES cells where knock-down of 241 *Gatad2a* and *Gatad2b* increased the diffusion coefficient of CHD4 (1.05-fold, $p < 10^{-4}$) (Extended Data 242 Figures 6d,e).

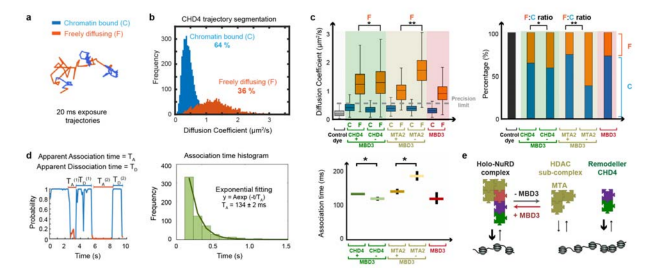
We then examined how the NuRD complex interacts with chromatin by comparing the percentage of freely diffusing versus chromatin bound CHD4 and MTA2 molecules in the presence and absence of MBD3. We observed a small decrease in the percentage of CHD4 molecules bound to chromatin in the absence of MBD3, but there was a much more significant decrease (2.4-fold, p < 0.01) in the percentage of chromatin

247 bound MTA2 molecules upon MBD3 depletion. This demonstrated that the chromatin bound deacetylase 248 subcomplex is also normally associated with CHD4 in the intact holo-NuRD complex. It also suggested that 249 CHD4 rather than the deacetylase sub-complex is primarily responsible for the association of NuRD with 250 chromatin (Figure 3c). This finding was supported by *in vitro* experiments which showed that, in comparison 251 to CHD4, the deacetylase subunit by itself does not bind strongly to nucleosomes (Extended Data Figure 252 **7a**). We conclude that NuRD normally exists as the intact complex *in vivo* and that, as with the removal of 253 MBD3, depletion of GATAD2A/B is sufficient to unlink CHD4 from the HDAC-containing sub-complex in 254 ES cells.

255 To investigate the chromatin binding kinetics of the CHD4 remodeller and the MTA2 deacetylase 256 sub-complex in the presence and absence of MBD3, we next determined association times from the time spent 257 freely diffusing between confined chromatin-bound states. We also attempted to determine dissociation times 258 from the time spent bound to chromatin between unconfined freely diffusing states (Figure 3d). The 259 distribution of association on times was well approximated by a single exponential, suggesting a Poissonian 260 process. Consistent with our finding that CHD4 is primarily responsible for recruitment of NuRD to 261 chromatin, we found no increase in the association time of CHD4 upon removal of MBD3. (Instead, we found 262 a small decrease consistent with the faster diffusion of the smaller CHD4 sub-complex.) However, we did find 263 a significant increase (1.3-fold, p < 0.01) in the association time of MTA2 upon MBD3 depletion (Figure 3d), 264 consistent with the finding that the deacetylase sub-complex does not strongly associate with chromatin in the 265 absence of CHD4 (Figure 3c and Extended Data Figure 7a). Although no changes in dissociation time were 266 observed (**Extended Data Figure 7b**), we reasoned that our trajectories may be truncated by photobleaching. 267 We therefore took advantage of 'motion blurring' when recording 500 ms trajectories to detect only chromatin bound proteins^{1,60}, and combined this with time-lapse imaging using different interval lengths in an attempt to 268 269 determine the dissociation time of the NuRD complex. To our surprise this showed that the dissociation times 270 were much longer than we had expected (greater than 100 seconds for MBD3, Extended Data Figure 7e), 271 such that it proved impossible to track individual molecules for long enough in order to determine reliable 272 dissociation rates. We conclude that, once bound to a target site the intact NuRD complex binds for 273 unexpectedly long times.

274

275 Figure 3. Live-cell single-molecule tracking reveals that the NuRD complex assembles before it binds to 276 chromatin. (a) Segmentation of an example 20 ms trajectory into confined chromatin bound (C) (blue) and 277 unconfined freely diffusing (F) states (orange). (b) Percentage of molecules and distribution of apparent 278 diffusion coefficients for chromatin bound and freely diffusing CHD4 molecules. (c) (Left) Box plot of 279 apparent diffusion coefficients for chromatin bound and freely diffusing CHD4 and MTA2 molecules in the 280 presence and absence of MBD3 [*p < 0.01, **p < 0.001 (Kolmogorov-Smirnov test)]. The data for MBD3 is 281 also shown as a control, and the grey dotted line indicates the upper bound of the precision limit calculated at 282 the 95 % confidence interval for immobilised JF_{549} dye molecules. (Right) Percentage of freely diffusing and 283 chromatin bound CHD4 and MTA2 molecules in the presence and absence of MBD3 [from Gaussian fitting, 284 *p < 0.01, **p < 0.001 (2-way ANOVA)]. [The number of cells/trajectories used in the analysis were: 285 30/5,557 (CHD4), 25/2,337 (CHD4-MBD3), 10/336 (MTA2), 10/652 (MTA2-MBD3) and 30/2,224 286 (MBD3).] (d) (Left) A plot of the confinement probability allows determination of the association T_A and 287 dissociation T_D times - defined respectively as the time a trajectory spends between periods of confined or 288 unconfined motion. (Middle) A single exponential curve of rate $lambda=1/T_A$ is then fit to the distribution of 289 association/dissociation times. (Right) The association times extracted for CHD4 and MTA2 were then 290 compared to those in the absence of MBD3, and to those for the MBD3 control [error bars show 95 % 291 confidence intervals, *p < 0.01 (2-way ANOVA)]. (e) Schematic representation of a model in which MBD3-292 dependent assembly of the NuRD complex increases the association rate of the deacetylase sub-complex.



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295 The Holo-NuRD complex modulates chromatin movement at enhancers and promoters

296 Because chromatin bound NuRD cannot be distinguished from immobile dye when imaging using 20 ms 297 exposures, we studied these slower moving molecules by imaging at a time resolution of 500 ms. (Here 298 motion blurring also prevents imaging of freely diffusing molecules.) The 500 ms imaging was facilitated by 299 the long residence times and trajectory lengths of the chromatin bound NuRD complex. Analysis of 500 ms 300 exposure trajectories (that last more than 5 seconds) initially revealed two states of chromatin bound CHD4. 301 Both the slow and fast states were primarily sub-diffusive (with an anomalous exponent α of around 0.5), and 302 were characterised by different confinement lengths of 62 ± 12 nm and 110 ± 30 nm and apparent diffusion 303 coefficients of 0.006 ± 0.002 and $0.018 \pm 0.006 \,\mu m^2/s$ (Figure 2c).

304 The apparent diffusion coefficient of the slow state was still below the precision limit even with 500 305 ms imaging: immobile dye molecules have a median precision of 34 nm and an apparent diffusion coefficient of $0.004 \pm 0.003 \,\mu\text{m}^2/\text{s}$. Interestingly, however, visualisation of the fast state trajectories revealed a proportion 306 307 of molecules exhibiting periods of directed motion, characterised by high anomalous exponents and high drift 308 (Figure 4a). To determine whether there is indeed a proportion of molecules undergoing directed motion, we 309 used Gaussian fitting to characterise the distributions in anomalous exponent for both the slow- and fast-states 310 of chromatin bound CHD4 (in separate calculations, see Extended Data Figures 8a-c). This revealed a single 311 slow state (S) with α_s of 0.59 ± 0.01 (67 % of sub-trajectories) and two fast states (F1 and F2) with different 312 anomalous exponents: α_{F1} of 0.60 ± 0.01 (26 %) and α_{F2} of 0.89 ± 0.02 (7 %) (Figure 4b). Molecules in the 313 fast F1 state have the same distribution of anomalous exponents as in the slow state and therefore explore the 314 same chromatin environment. However, they diffuse faster and have a larger length of confinement and thus 315 move further within the nucleus (Figure 4b). As can be seen in the example in Figure 4a, molecules in the 316 fast F2 state, however, have a higher anomalous exponent and they explore a larger area of the nucleus than 317 those in both the slow and the fast F1 states. Moreover, they have higher drift, indicative of movement in a 318 directed manner: this is also consistent with the higher anomalous exponent.

We next compared the dynamics of chromatin bound MBD3 to that of CHD4 and found that it too exhibited the one slow and two fast states. Both chromatin bound MBD3 and CHD4 molecules exhibited motion in the fast F1 and F2 states in around 22-26% and 7-8% of trajectories, respectively, confirming that these states are a property of the intact NuRD complex and not just of CHD4 (**Figures 4c,d**). Importantly, visualisation of the trajectories identified individual molecules that switch between the three states; S, F1 and F2 (**Figure 4a**). Thus, the three states are unlikely to either represent CHD4 forming different complexes or NuRD complex molecules bound in different regions of the nucleus.

326 We then compared the dynamics of chromatin bound CHD4 molecules in the *Mbd3*-ko and wild-type 327 cells. Surprisingly, we found that the anomalous exponent, length of confinement and apparent diffusion 328 coefficient of the fast F1 and F2 states were all higher in wild-type cells (Figure 4c). The increase in 329 anomalous exponent in wild-type cells unexpectedly suggests that in the presence of the intact NuRD complex 330 chromatin is less *condensed*, whilst the increased apparent diffusion coefficient and length of confinement 331 show that CHD4 molecules diffuse more rapidly and explore a larger nuclear volume. We had expected to 332 find that recruitment of the deacetylase by CHD4 would lead to less acetylated chromatin and greater condensation⁶¹⁻⁶⁵ – and that the bound CHD4 molecules in wild-type cells would thus explore a smaller 333 334 nuclear volume. In addition, in the presence of intact NuRD we also observed a significant decrease in the 335 proportion of CHD4 molecules in the fast F2 state exhibiting directed motion (7.4 % in wild-type cells vs 18% 336 in *Mbd3*-ko cells; see Figure 4d and Extended Data Figure 8d).

337 The fast F1 and F2 states of chromatin bound NuRD could result from movement on DNA due to 338 chromatin remodelling or, bearing in mind the long dissociation times we determined for CHD4 (see above), 339 from movement of NuRD-bound enhancers. To distinguish between these possibilities we targeted sites 340 nearby active enhancers with dCas9-GFP, either by transfecting a published and validated CARGO vector 341 expressing 36 different gRNAs targeting a Tbx3 enhancer³ or by transfecting a single gRNA that targets DNA 342 repeats near the *Nanog* gene (Extended Data Figures 9a,b). Targeting using our single gRNA was validated 343 by confirming co-localisation of GFP-tagged dCas9 (detected by immunofluorescence) with DNA FISH 344 probes (Extended Data Figure 9c). We carried out these experiments in cells that express an ER-MBD3-ER 345 (estrogen receptor-MBD3-estrogen receptor) fusion protein in which the nuclear localisation of MBD3 (and 346 thus assembly of the intact NuRD complex) is tamoxifen-inducible¹⁸. In this system the intact NuRD complex 347 assembles and remodels chromatin/transcription factor binding following induction: after 24 hours the 348 transcription factor landscape has been reset and transcriptional changes have occurred. This allowed us to

349 study the new chromatin environment created by NuRD shortly after it had become established (thus allowing 350 us to distinguish direct from downstream effects). We imaged cells showing bright undivided foci to avoid 351 using data from cells in the S or G2 phases of the cell cycle, which exhibit blurred foci or doublets (Extended 352 **Data Figure 10a**). Because the background fluorescence from freely diffusing dCas9-GFP prevented 3D 353 tracking of genomic loci on our DH-PSF microscope, we tracked the enhancer loci in 2D. Although this 354 meant that we could not directly compare the parameters obtained in the 2D (active enhancer) and 3D (NuRD 355 single molecule) tracking experiments, classification of the sub-trajectories once again revealed one slow and 356 two fast chromatin states. The enhancer locus trajectories (Figure 5a) were also much longer than those of 357 chromatin bound single CHD4 molecules (Figure 4a) allowing a better characterisation of switching between 358 the different fast states (Extended Data Figures 10b-d). We found that, as when tracking CHD4 single 359 molecules, addition of MBD3 (and thus intact NuRD assembly) significantly increased the apparent rate of 360 diffusion of both enhancers in the fast-diffusing F1 and F2 states (Figure 5b). As when tracking CHD4, an 361 increased proportion of sub-trajectories in the fast decondensed F2 state in wild-type cells was observed for 362 both the *Tbx3* and *Nanog* enhancers (Figure 5c).

363 To further confirm the effect of NuRD on enhancer dynamics we also imaged endogenous chromatin bound KLF4, a transcription factor which like NuRD binds to active enhancers in ES cells¹⁸ and which also 364 365 clusters with NuRD in 3D genome structures¹⁷. We carried out 3D single molecule tracking of KLF4-HaloTag 366 both before and 24 hours after inducing nuclear localisation of MBD3, and thus assembly of the intact NuRD 367 complex, using tamoxifen. Once again we observed a significant increase in the length of confinement and 368 apparent diffusion coefficient of both the slow and fast chromatin bound states of KLF4 (Figure 5d). The 369 observation of similar effects 24 hrs after reassembly of the NuRD complex (and also at shorter times of 6 hrs 370 after reassembly, data not shown), when either tracking active enhancers or chromatin bound KLF4, suggests 371 that we are looking at a direct effect of NuRD on enhancer dynamics. Finally, to strengthen our conclusion 372 that chromatin decompacts in the presence of the intact NuRD complex, leading to an increase in 373 intermediate-range chromatin interactions, we carried out two-colour enhancer-promoter DNA FISH studies 374 of three key pluripotency genes Bmp4, Sox2 and Tbx3. In the presence of NuRD we found a significant 375 increase (p<0.001) in the distance between their enhancers and promoters indicative of chromatin 376 decompaction (Extended Data Figures 9d.e).

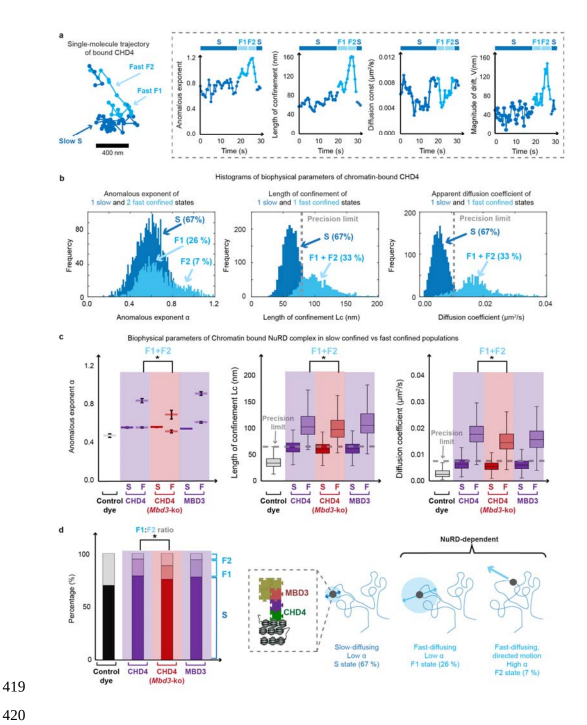
377 Relationship of NuRD mediated modulation of enhancer dynamics to previous studies

378 How might the fast sub-diffusive states of the chromatin bound NuRD complex (F1 and F2) relate to 379 the previous enhancer/promoter tracking experiments of Gu et al. (2018), and their transcription-dependent 380 enhancer 'stirring' model? The similarity in both the diffusion parameters and anomalous exponents of their 381 slow/fast and our S/F1 states, suggests that these correspond to either inactive (slow/S) or actively transcribed 382 (fast/F1) enhancers and promoters. In our experiments, however, we additionally observe a fast F2 state which 383 exhibits directed motion and we wondered whether this was due to chromatin movement resulting from 384 transcriptional elongation? We therefore tracked bound CHD4 molecules after adding DRB, a small molecule 385 inhibitor of transcriptional elongation⁶⁶. Consistent with the results of Gu et al. (2018), premature termination 386 by DRB led to a modest reduction in the proportion of bound CHD4 molecules exhibiting the fast F1 387 (transcription dependent) motion (from 26 % to 19 %). However, there was no reduction in the proportion of 388 molecules in the fast F2 state nor change in the chromatin environment (i.e. in the anomalous exponent or 389 length of confinement) in the presence of a block on transcriptional elongation (Extended Data Figures 390 **8b,d**). Finally, we tracked MBD3 molecules while blocking HDAC1/2 deacetylase activity with FK228⁶⁷. As 391 was the case when we prevented the association of the deacetylase sub-complex with CHD4 (see Figure 4c) 392 we again observed a decrease in the anomalous exponent (reflecting a more condensed state), but there was no 393 significant change in the proportion of molecules in the fast decondensed F2 state (Extended Data Figures 394 **8b,d**). We conclude that the fast F2 state is not dependent on transcriptional elongation or deacetylation 395 activity. Instead, the results suggest that the assembly of the intact holo-NuRD complex (from both the 396 chromatin remodelling and deacetylase subunits) modulates the dynamics of active enhancers, affecting their 397 conversion between the slow S, fast F1 and fast F2 states (Figure 5e).

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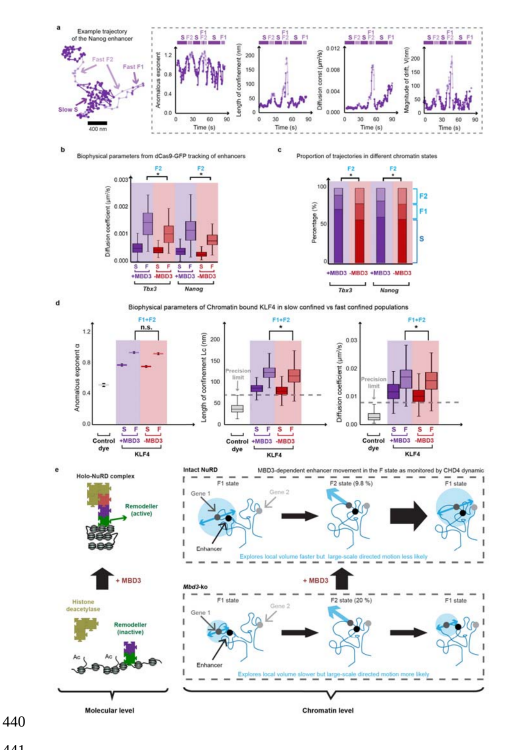
399 Figure 4. Assembly of the intact NuRD complex decondenses chromatin and reduces directed motion.

400 (a) (Left) Example trajectory of a chromatin bound CHD4 molecule showing periods of both slow (dark blue) 401 and fast (light blue) sub-diffusive motion. Two fast states (F1 and F2) are observed, with the F2 state showing 402 periods of directed motion. (Right) The four biophysical parameters calculated along this trajectory with the 403 fast F2 sub-trajectories showing a higher anomalous exponent, increased length of confinement and increased 404 drift. (b) (Left) Gaussian fitting to the distribution of anomalous exponents identified a single α for slow 405 confined NuRD complex molecules (dark blue), and two α values for molecules in the faster confined state 406 (light blue). The resulting distribution of the lengths of confinement (Middle) and apparent diffusion 407 coefficients (Right) are also shown. (c) Comparison of biophysical parameters for the CHD4 remodeller in the 408 presence and absence of MBD3, and for MBD3 itself [the number of cells/trajectories used in the analysis 409 were: 30/3,059 (CHD4), 15/2,111 (CHD4-MBD3), and 30/1,816 (MBD3)]. (Left) The anomalous exponents 410 resulting from Gaussian fitting (error bars show 95 % confidence intervals, *p < 0.01, 2-way ANOVA); 411 (Middle) Boxplots of the lengths of confinement; and (Right) the apparent diffusion coefficients are shown 412 with the precision limit of the single molecule imaging indicated by a grey dotted line (*p < 0.01, 413 Kolmogorov-Smirnov test). (We imaged JF₅₄₉ dye fixed on a coverslip and found that all three biophysical 414 parameters were significantly lower than the S, F1 and F2 states of NuRD components.) (d) (Left) Percentage 415 of molecules in the slow or fast chromatin bound states (from the Gaussian fitting, *p < 0.01, 2-way 416 ANOVA). (Right) Schematic representation of the three states of chromatin bound NuRD: a slow sub-417 diffusive S state with small exploration volume, one fast sub-diffusive F1 state with a larger length of 418 confinement (exploration volume) and another fast sub-diffusive F2 state exhibiting directed motion.



420

421 Figure 5. Assembly of the intact NuRD complex modulates the movement of active enhancers. (a) (Left) 422 Example trajectory of the Nanog enhancer segmented to show periods of slow and fast sub-diffusive motion 423 (dark and light blue respectively). (Right) The anomalous exponent, length of confinement, apparent diffusion 424 coefficient and the magnitude of drift of the locus extracted from the trajectory shown. (b) Boxplots of the 425 apparent diffusion coefficient calculated for 2D trajectories of loci nearby the Tbx3 and Nanog enhancers, in 426 the presence and absence of MBD3 (*p < 0.01, Kolmogorov-Smirnov test). Number of trajectories: 237 427 (Tbx3+MBD3) and 287 (Tbx3-MBD3); 546 (Nanog+MBD3) and 229 (Nanog-MBD3). (c) Gaussian fitting to 428 the distribution of the anomalous exponents identifies a single slow and two faster states for enhancer loci. 429 The percentage of sub-trajectories of enhancer loci exhibiting slow (S) and fast chromatin motions with either 430 a low (F1) or high (F2) anomalous exponent α is shown (*p < 0.01, 2-way ANOVA). (d) Comparison of 431 biophysical parameters for the KLF4 transcription factor in the presence and absence of MBD3. (Left) The 432 anomalous exponents (error bars show 95 % confidence intervals from Gaussian fitting, *p < 0.01, 2-way 433 ANOVA). (Middle and Right) Boxplots of the lengths of confinement and the apparent diffusion coefficients 434 are shown with the precision limit of the single molecule imaging indicated by grey dotted lines (*p < 0.01, 435 Kolmogorov-Smirnov test). Number of cells/trajectories used in the analysis were: 57/179 (KLF4+MBD3), 436 and 56/652 (KLF4-MBD3). (e) Schematic model depicting: (Left) a molecular level view of how assembly of the NuRD complex affects chromatin remodelling¹⁸; and (Right) a chromatin level view of how intact NuRD 437 438 increases the volume explored by an enhancer while at the same time reducing its likelihood of entering the 439 F2 state in which the directed motion of chromatin might re-organise enhancer-promoter proximity.



441

442 The NuRD complex modulates the dynamics of enhancers to control transcription

443 Whilst we cannot rule out that some of the fast motions we observe when imaging the chromatin bound NuRD 444 complex might result from chromatin remodelling, given that we observe similar dynamics for chromatin 445 bound NuRD, active enhancers and chromatin bound KLF4, the results suggest that single molecule tracking 446 of chromatin bound NuRD largely monitors the spatial dynamics of active enhancers. How might assembly of 447 the deacetylase sub-complex with CHD4 to form intact NuRD influence transcription? Histone acetylation is 448 thought to lead to a looser packing between nucleosomes and chromatin decompaction⁶¹⁻⁶⁵, but here, by 449 studying compaction in the immediate vicinity of the stably bound NuRD complex, we unexpectedly find that 450 assembly of the intact NuRD complex leads to a more open chromatin structure where enhancers can diffuse 451 faster and explore more of the nucleus (see Figure 5e). The chromatin decompaction and faster movement of 452 enhancers in the presence of intact NuRD provides a molecular explanation for the increase in intermediate-453 range chromatin interactions (Figure 1b), the blurring of TAD and A/B compartment boundaries (Figure 1c), 454 the increase in longer range enhancer-promoter interactions (Figure 1d), and the changes in enhancer 455 interactions with promoters that are regulated by NuRD (Figure 1e). Our experiments also show that 456 assembly of the intact NuRD complex reduces the time NuRD-bound enhancers spend undergoing directed 457 motion, and we propose that this may limit the re-organisation of enhancer-promoter interactions leading to 458 more stable, long-lived interactions. This could explain the observed increase in transcriptional noise, or low-459 level inappropriate transcription, observed in both human and mouse ES cells lacking functional NuRD^{68,69}. 460 CHD4 in the ChAHP complex has previously been shown to inhibit CTCF/Cohesin binding at repetitive DNA 461 sequences⁷⁰. Our results reveal that CHD4 in the NuRD complex plays a more general role by modulating 462 CTCF/Cohesin interactions and cross-linking in active chromatin to promote an environment where enhancers 463 and promoters can contact each other over longer distances, thereby reducing the fast F2 motions that our 464 imaging approach has uncovered.

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Online Methods

In-nucleus chromosome conformation capture (Hi-C)

Data acquisition

In-nucleus Hi-C was carried out as previously described¹⁷ on E14 wild-type and 7g9 *Mbd3-ko* (see Ref. 31 for construction and characterisation) ES cells. 50 bp paired-end sequencing was carried out on a HiSeq4000 instrument. Our own Hi-C analysis of *Mbd3-ko* ESCs cultured in 2i consisted of three experiments: SLX-18035, SLX-7676 and SLX-19611 (which was sequenced twice). Alongside these four datasets, we also collected wild-type 2i data (SLX-7672) and compared this with recently published Hi-C data for ESCs cultured in 2i conditions (https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-6591/) (Extended Data Figure 1a). We used 4 replicates of this published data (ERR239137, ERR239139, ERR239141, ERR239143).

In situ Hi-C							
Experiment	Source	Condition	Unique Contacts				
ERR2391437	E-MTAB-6591	WT	131,434,520				
ERR2391439	E-MTAB-6591	WT	133,825,748				
ERR2391441	E-MTAB-6591	WT	144,255,897				
ERR2391443	E-MTAB-6591	WT	132,181,206				
SLX-7672	This study	WT	55,178,175				
SLX-7676	This study	Mbd3-ko	64,883,460				
SLX-18035	This study	Mbd3-ko	124,812,397				
SLX-19611_s_1	This study	Mbd3-ko	69,008,264				
SLX-19611_s_2	This study	Mbd3-ko	68,483,385				

Pre-processing

For each experimental condition and replicate, FASTQ files were first processed and aligned against the mouse GRCm38.p6 reference genome using the nuc_processing package (<u>https://github.com/tjs23/nuc_processing</u>). The number of unique contacts observed varied significantly between replicates and conditions but overall a high number of reads were obtained in both conditions. The resulting raw contacts in NCC format were converted into hic and cooler format contact matrices using the

juicer Pre (<u>https://github.com/aidenlab/juicer/wiki/Pre</u>) and HiCExplorer hicConvertFormat tools for downstream analysis

(https://hicexplorer.readthedocs.io/en/latest/content/tools/hicConvertFormat.html)⁷¹⁻⁷³. To assess the correlation between replicates within and between the two experimental conditions, we made use of the HiCExplorer hicPlotdistvscounts tool. Clustering of replicates based on contact distance distributions identified clusters for the two experimental conditions showing good agreement between replicates. We therefore merged the obtained contacts resulting in 705,063,441 and 327,180,884 contacts for the wild-type and *Mbd3-ko* conditions, respectively. Due to the difference in read coverage, we down-sampled our merged wild-type dataset to have the same number of contacts as our merged *Mbd3-ko* dataset. Finally, we performed Knight-Ruiz normalisation of the merged datasets to ensure balanced matrices.

Compartment Analysis

To identify A/B chromatin compartments within our merged datasets we made use of the recently developed CscoreTool (https://github.com/scoutzxb/CscoreTool)⁷⁴. This tool assigns each genomic window *i* a score $\in [-1,1]$ with = 1 assigning a probability 1 that window *i* is in the A-compartment while = -1 assigns a probability 1 that window *i* is in the B-compartment. Importantly, unlike eigenvector analysis, c-scores of different samples can be directly compared since they represent probabilities. Binary compartments can also be assigned based on the sign of the c-score.

TAD Analysis

TAD calling was performed using Lavaburst, a recently published $tool^{41}$ which uses an insulation-style metric called the TAD-separation score to identify the degree of separation between the up and downstream region at each Hi-C matrix bin. Local minima of the TAD separation score are considered as putative TAD boundaries and assigned q-values for calculation of a false-discovery-rate (fdr). For this analysis we used parameters gamma = 10 and beta = 50,000. For both datasets, we identified TADs using a binning resolution of 25 kb.

Enhancer-Promoter Links

We wanted to investigate whether transcriptional mis-regulation in *Mbd3-ko* ESCs was significantly associated with a disruption of cis-regulatory interactions. To identify putative promoter-regulatory element (RE) interactions we made use of a recent study⁵⁰ which profiled RE-promoter interactions via thousands of separate CRISPR deletions. In particular, Fulco et al. (2019) found that a relatively simple activity-by-contact (ABC) model could be used to identify, with reasonable precision and recall, functional RE-promoter interactions. Crucially, the ABC model produced interactions with significantly higher accuracy than those identified using either linear distance along the genome or Hi-C contacts alone.

The ABC model considers the genome in 5 kb bins. For each promoter p and regulatory element r we define:

$$ABC_{p,r} = \frac{A_r \times C_{p,r}}{\sum_{s \in \mathcal{N}(p)} A_s \times C_{p,s}}$$

Where $\mathcal{N}(p)$ is the set of all regulatory elements within 5Mb of p, $C_{p,r}$ is the contact frequency between p and r, and A_r is the activity of r. For regions of the genome with poor read coverage, $C_{p,r}$ is estimated assuming a power-law decay of contact frequencies:

$$C_{p,r} \sim d_{p,r}^{-\gamma}$$

where $d_{p,r}$ is the genomic distance between the promoter and regulatory element while γ is inferred from the Hi-C contact maps. Fulco et al. (2019) define the activity of RE *r* as the geometric mean of read counts of DNaseI hypersensitive (DHS) and H3K27ac chromatin immunoprecipitation sequencing (ChIP-seq) at *r*. We did not have DHS tracks for our *Mbd3-ko* ES cells and therefore implemented our own version of the ABC model where we used the read counts in just the H3K27ac ChIP-seq data to score the activity of each regulatory element. Specifically, we identified putative regulatory elements using H3K4me3 and H3K27ac ChIP-seq data from wild-type (WT) cells and cells where MBD3 had been depleted¹⁸ (KO) as follows:

- promoter regions were assigned as those regions ±1kb of a TSS and overlapping with an H3K4me3 ChIP-seq peak,
- H3K27ac ChIP-seq peaks from that condition were considered as an initial putative list of condition specific RE's,

- 3. H3K27ac peaks closer than 500bp were merged,
- 4. Peaks with total length <500bp were discarded,
- 5. Peaks overlapping with promoter regions were discarded, and
- 6. The union of the non-promoter peaks and promoter regions was then treated as a master list of putative RE's which were each assigned a condition-specific score based on the mean H3K27ac peak strength across that RE.

In particular, REs could be assigned within a single condition as being either 'intergenic' or 'promoter' associated. Since REs were defined per condition, WT-unique, KO-unique and common overlapping peaks were assigned unique IDs for downstream analysis. The relevant code to perform this analysis and subsequent calculation of ABC scores can be found at: (https://github.com/dhall1995/Acitivity-By-Contact Enhancer-Promoter_Link_Prediction).

Using calculated ABC scores, WT and KO links were assigned unique link-IDs based on the promoter and enhancer pair in question as well as the promoter genomic position in the case of a gene with multiple possible promoter regions. The top 10% of all identified links (WT & KO) when ranked by ABC score were then selected as 'strong' links. In our data this corresponded to an ABC threshold of ~0.12. Based on this threshold, unique link IDs could be assigned as either WT-unique, KO-unique or common depending on the conditions in which we observed that link. These thresholds were chosen to maximise the precision of identified links while retaining a significant number of links to analyse although we acknowledge that maximising precision comes at the expense of recall using the ABC model. Finally, after identification of WT-unique, KO-unique and common links, enrichment analysis was performed by associating each link with a promoter and performing χ^2 analysis using the statsmodels python module.

CTCF and SMC3 Cut&Run

Cut&Run experiments were carried out according to Meers *et al.* $(2019)^{75}$ using a *Drosophila* genomic DNA spike-in and "input" controls consisting of samples which were processed in parallel with the Cut&Run samples but with unterhered MNase. Western blots of nuclear lysates were also carried out as previously described⁶⁸ to measure the relative levels of CTCF and SMC3 in wild-type and *Mbd3-ko* ES cells (**Extended Data Fig 4a**). The antibodies used were:

ANTIBODY	RAISED IN	COMPANY	CATALOGUE NUMBER	WESTERN BLOT	ChIP	Clonality
αCTCF	Rabbit	Millipore	07-729	1:1000	5µl	Polyclonal
aSMC3	Rabbit	Abcam	ab9263	1:1000	5µl (5µg)	Polyclonal

50 bp paired-end sequencing was carried out on a Novaseq instrument – three biological replicates per sample were obtained with 8-16 million mapped reads/replicate, respectively, whilst the input samples had 8-23 million mapped reads/replicate, respectively. All Cut&Run data was trimmed using trim_galore and then aligned using standard Bowtie parameters to the Mus musculus reference genome (mm10) (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/). Heatmaps of Cut&Run enrichment were made using Deeptools v2.5.0⁷⁶. Cut&Run bigwig tracks were calculated using BedTools and bedGraphToBigWig with standard parameters. The coverage was calculated with computeMatrix reference-point with options --binSize 10. The heatmap of standardised signal was then plotted using plotHeatmap. Peaks were called using MACS2 so as to give a False Discovery Rate of 1% and above 5-fold enrichment. All Venn Diagrams were plotted using the matplotlib-venn library in Python v3.6.

Mouse embryonic stem cell line generation

Mouse embryonic stem (ES) cell lines were cultured in 2iL conditions⁷⁷ – 50 % DMEM/F-12 medium (Gibco 21041025) and 50 % NeurobasalTM Medium (Gibco 12348017) supplemented with 1 x N2 to a final concentration of 2.5 µg/ml insulin (provided in-house by the Cambridge Stem Cell Institute), 0.5x B-27TM Supplement (Gibco 17504044), 1x minimum essential medium non-essential amino acids supplement (Sigma-Aldrich, M7145), 2mM L-glutamine (Life tech, 25030024), and 0.1 mM 2-mercaptoethanol (Life tech, cat. 21985023), 2i inhibitors (1 µM PD0325901, 3 µM CHIR99021) and 10 ng/ml mouse leukemia inhibitory factor (mLIF provided by the Biochemistry Department, University of Cambridge). Cells were passaged every two days by washing in PBS (Sigma-Aldrich, D8537), adding Trypsin-EDTA 0.25 % (Life tech, cat. 25200072) to detach the cells, and then washing in media before re-plating in fresh media. To help the cells attach to the surface, plates were incubated for 15 minutes at room temperature

(RT) in PBS containing 0.1 % gelatin (Sigma Aldrich, G1890). All cell lines were routinely screened for mycoplasma contamination at least twice yearly and tested negative.

ES cells expressing CHD4 tagged at the C-terminus with HaloTag were generated in the presence and absence of MBD3, as previously described^{17,24,25}. Briefly, this was achieved by CRISPR/Cas9 based knock-in of a cassette containing mEos3.2-HaloTag-FLAG and a puromycin selection gene into one *CHD4* allele. The puromycin cassette was then removed using Dre recombinase to generate the *CHD4* allele with a C-terminal HaloTag fusion. Since knockout of *CHD4* is lethal, we used cell viability assays and the ability to immunoprecipitate NuRD component proteins (**Extended Data Figure 4c**) to verify that the function of the tagged CHD4 was not significantly impaired. We similarly generated knock-in ES cells in an E14Tg2a (XY) background expressing MBD3 tagged at the C-terminus with HaloTag (**Extended Data Figure 4b**). MTA2-HaloTag knock-in cell lines were generated in MBD3-inducible ES cells¹⁸ (**Extended Data Figure 4d**), in which MBD3 is fused to the oestrogen receptor at both the N- and C-termini so that it initially localises at the cytoplasm but then translocates to the nucleus when induced with 4-hydroxytamoxifen added directly to the culture media to a final concentration of 0.4 nM. Western blots were carried out using nuclear lysates as previously described¹⁸, to confirm the expression and assembly of the NuRD complex (**Extended Data Figure 4a**). Immunoprecipitations were carried out using antibodies to CHD4 or MTA2, or Halo-Trap beads (ChromoTek) in the case of the CHD4-Halo line. The antibodies used were:

ANTIBODY	RAISED IN	COMPANY	CATALOGUE NUMBER	WESTERN BLOT	Clonality
αCHD4	Mouse	Abcam	ab70469	1:5000	Monoclonal [3F2/4]
αFlag	Mouse	Sigma	F1804	1:5000	Monoclonal M2
αGATAD2A	Rabbit	Abcam	ab87663	1:2000	Polyclonal
αHDAC1	Rabbit	Abcam	ab7028	1:2000	Polyclonal
aMBD3	Rabbit	Abcam	ab157464	1:5000	Monoclonal [EPR9913]
αMTA2	Mouse	Abcam	ab50209	1:5000	Monoclonal [MTA2- 276]
αΡCNA	Mouse	Santa cruz	Sc56	1:2000	Monoclonal [PC10]

Live-cell 3D single-molecule imaging

ES cells expressing HaloTag-tagged CHD4, MBD3 and MTA2 were passaged two days before imaging onto 35 mm glass bottom dishes No 1.0 (MatTek Corporation P35G-1.0-14-C Case) in serum/LIF imaging medium: FluorobriteTM DMEM (Thermo Fisher Scientific, A1896701) containing 100 mM 2-mercaptoethanol (Life tech, cat. 21985023), 1x minimum essential medium non-essential amino acids (Sigma-Aldrich, M7145), 2 mM L-glutamine (Life tech, cat. 25030024), 1 mM sodium pyruvate (Sigma-Aldrich, S8636-100ML), 10 % fetal bovine serum (HyClone FBS, Lot nr SZB20006, GE Healthcare Austria SV30180.03) and 10 ng/ml mLIF (provided by the Biochemistry Department, University of Cambridge). Glass bottom dishes had been prepared by incubation in 0.01% poly-L-ornithine (Sigma-Aldrich P4957) for 30 minutes, followed by two rinses in PBS and incubation in PBS containing 10 μ g/mL laminin (Sigma-Aldrich L2020) for at least 4 hours. Just before single-molecule imaging experiments, cells were labelled with 0.5 nM HaloTag[®]-JF₅₄₉ ligand for 15 minutes, followed by two washes in PBS and a 30 minute incubation at 37 °C in imaging medium, before imaging the cells in fresh serum/LIF imaging medium. Cells were under-labelled to prevent overlap of fluorophores during single-molecule tracking experiments. The HaloTag dyes were a kind gift from Luke D. Lavis (HHMI).

For the HaloTag-NLS control, the pEF-HaloTag-NLS vector was generated by replacing the HP1 sequence in a previously described HaloTag-HP1 expression vector⁷⁸ with a SV40 NLS sequence. 1µg of the pEF-HaloTag-NLS vector was transfected into ES cells using Lipofectamine 2000 Transfection Reagent (Thermo Fisher Scientific 11668019) during passaging onto 35 mm glass bottom dishes No 1.0 (MatTek Corporation P35G-1.0-14-C Case). Media was changed after 24 hours and samples were both labelled as above and imaged the following day.

Transcription elongation was inhibited using 100 μ M 5,6-dichloro-1- β -D-ribofuranosylbenzimidazole (DRB) and deacetylase activity using 10 nM FK228 (TOCRIS Bioscience, UK) both for two hours prior to imaging^{67,79}. qPCR experiments confirmed inhibition of transcription upon addition of DRB (data not shown).

A custom-made double-helix point spread function (DHPSF) microscope was then used for 3D single-molecule tracking as previously described²⁵. The setup incorporates an index-matched 1.2 NA water immersion objective lens (Plan Apo VC 60×, Nikon, Tokyo, Japan) to facilitate imaging above the coverslip surface. The DHPSF transformation was achieved by the use of a 580 nm optimized double-helix phase mask

(PM) (DoubleHelix, Boulder, CO, USA) placed in the Fourier domain of the emission path of a fluorescence microscope (Eclipse Ti-U, Nikon). The objective lens was mounted onto a scanning piezo stage (P-726 PIFOC, PI, Karlsruhe, Germany) to calibrate the rotation rate of the DHPSF. A 4f system of lenses placed at the image plane relayed the image onto an EMCCD detector (Evolve Delta 512, Photometrics, Tucson, AZ, USA). Excitation and activation illumination was provided by 561 nm (200 mW, Cobolt Jive 100, Cobolt, Solna, Sweden) and 405 nm (120 mW, iBeam smart-405-s, Toptica, Munich, Germany) lasers, respectively, that were circularly polarized, collimated, and focused to the back focal plane of the objective lens. Obliqueangle illumination imaging was achieved by aligning the laser off axis such that the emergent beam at the sample interface was near-collimated and incident at an angle less than the critical angle ($\theta_c \sim 67^\circ$) for a glass/water interface. The fluorescence signal was then separated from the excitation beams into the emission path by a quad-band dichroic mirror (Di01-R405/488/561/635-25x36, Semrock, Rochester, NY) before being focused into the image plane by a tube lens. Finally, long-pass and band-pass filters (BLP02-561R-25 and FF01-580/14-25, respectively; Semrock) placed immediately before the camera isolated the fluorescence emission. Using 561 nm excitation, fluorescence images were collected as movies of 60,000 frames at 20 ms or 4,000 frames at 500 ms exposure. A continuous 561 nm excitation beam at $\sim 1 \text{ kW/cm}^2$ was used for 20 ms exposure imaging and at $\sim 40 \text{ W/cm}^2$ for 500 ms exposure imaging. Each experiment was carried out with at least 3 biological replicates (3 fields of view, each containing around 3 cells).

Residence time analysis from time-lapse 500 ms exposure imaging

Since photobleaching is related to the number of exposures, and the residence time is related to the time a molecule spends bound to chromatin, it is possible to change the time-lapse between exposures and use the data to extract both the residence time¹ and photobleaching rate. However, when we imaged at time intervals of 0.5 s, 2.5 s, 8 s and 32 s, we discovered that at the longest time lapse (32 s) we could see no decrease in the mean number of frames imaged before photobleaching, implying that the residence time had no impact on the measurement, which was thus dominated by photobleaching (**Extended Data Figure 7e**). To estimate the residence time would likely require imaging at much longer time-lapses, but because chromosomes and the cell itself move significantly during periods longer than this, it becomes unreliable to track individual chromatin bound NuRD complex subunits.

3D single-molecule image processing, generation of trajectories and determination of experimental precision

Single molecules were localized from 3D movies using the easy-DHPSF software⁸⁰ with a relative localization threshold of 100 for all 6 angles for the 20 ms data and relative thresholds of 116, 127, 119, 99, 73 and 92 for the 500 ms data. The trajectories of individual molecules were then assembled using custom Python code for connecting localisations in subsequent frames if they were within 800 nm for 20 ms trajectories and within 500 nm for 500 ms trajectories (<u>https://github.com/wb104/trajectory-analysis</u>). This code also outputs average signal intensity per trajectory and trajectory lengths (OPTION - savePositionsFramesIntensities) and a summary of this data is reported in **Extended Data Figure 7d**.

Our precision values (measured for fixed dye molecules on the coverslip and calculated using the approach described by Endesfelder *et al.*⁸¹ were 60 nm and 34 nm for the 20 and 500 ms tracking experiments, respectively. The lower limits of the effective diffusion coefficient one can measure are dependent on the precision values. The effective diffusion coefficient (D_{eff}) is equal to the displacement squared/time. Thus, if the upper limit of the precision is 60 nm, then the lower limit of D_{eff} that we can measure for 20 ms imaging is: $0.06*0.06/0.02 = 0.18 \,\mu\text{m}^2 s^{-1}$. For 500 ms imaging on the other hand, the upper limit of our precision was 34 nm, corresponding to a lower limit of D_{eff} that we can measure of: $0.034*0.034/0.5 = 0.002 \,\mu\text{m}^2 s^{-1}$. Consistently, when we measured the apparent diffusion coefficients for dye molecules attached to a coverslip we determined values of $0.3 \pm 0.2 \,\mu\text{m}^2$ /s and $0.004 \pm 0.003 \,\mu\text{m}^2$ /s, respectively (see **Extended Data Figures 6c and 8b**). Any measured diffusion coefficients below these values do not have a biophysical interpretation.

Single-molecule trajectory analysis

We describe below (see Section 1) a switching model for a stochastic process between two or more states, characterized by different diffusion coefficients. We initially used this approach to decide whether we should develop a two- or three-state diffusion model to classify the distribution of displacements measured along single molecule trajectories. We then went on to develop an improved and novel algorithm to classify sub-

trajectories into confined and unconfined states based on four physical parameters using a Gaussian mixture model. This is described in Section 2. The algorithm was then tested on simulated trajectories which provided ground truth data. Finally, in Section 3, we describe how we applied this classification algorithm to single particle trajectories to estimate the diffusion coefficients in confined and unconfined states. However, the original classification based on diffusion coefficients provided a complementary analysis, and justified the results we obtained later, which is why it is included here.

1. Switching dynamics and displacement analysis

We describe the motion of a particle that switches between different states characterized by different diffusion coefficients^{82,83}. We initially used this description to identify the most appropriate number of switching states, based on the distribution of the length of the instantaneous displacements $\Delta X = X(t+\Delta t)-X(t)$. First, we describe the switching dynamics and fitting procedure, and then how we used the Bayesian Information Criterion (BIC) to select the optimal model.

1.1 Stochastic motion described by a diffusion model

The dynamics of a chromatin-binding protein complex such as NuRD, or some component of it, can be described by Langevin's equation^{84,85} in the classical overdamped limit, where the position X(t) satisfies:

$$\dot{\boldsymbol{X}} = \frac{\boldsymbol{f}(\boldsymbol{X})}{\gamma} + \sqrt{2D}\dot{\boldsymbol{\eta}}, \qquad (1)$$

where the force f depends on the position X and the second term corresponds to steady-state diffusion in a crowded medium, characterized by a constant diffusion coefficient D. Here η is a Gaussian variable with mean 0 and variance 1. When the protein binds to its molecular partners or chromatin, the diffusion coefficient could change, and the motion can become restricted, e.g. to that along the path of the DNA.

Experimental trajectories consist of a series of points ($X(k\Delta t)$ k = 1... and in the absence of any additional localization error noise, or an external force f = 0, the displacement dynamics at the sampling rate interval Δt , is given by:

$$\boldsymbol{X}(t + \Delta t) = \boldsymbol{X}(t) + \sqrt{2D\Delta t}\boldsymbol{\eta}, \qquad (2)$$

where η is a vector of Gaussian values of mean 0 and variance 1. In two dimensions, the distribution of displacements is:

$$Pr\{|\Delta \boldsymbol{X}| = u\} = \frac{u}{\sigma^2} \exp\left(-\frac{u^2}{2\sigma^2}\right) \qquad (3)$$

where $\sigma^2 = 2D\Delta t$ (see Eq. 5). For empirical displacements that contain a Gaussian localization error, characterized at the time resolution Δt by the amplitude σ_{le} , the formula should be modified⁸⁶:

$$Pr\left\{|\Delta \boldsymbol{X}|=u\right\} = \frac{u}{2D\Delta t + \sigma_{lc}} \exp\left(-\frac{u^2}{2(2D\Delta t + \sigma_{lc}^2)}\right)_{(4)}$$

We can thus use the approximated diffusion coefficient:

$$D_{app} = D + \frac{\sigma_{le}^2}{2\Delta t}.$$
(5)

1.2 Modelling stochastic switching behaviour

We next constructed a model for a chromatin-binding protein by using a classical Markov chain with two states 1 and 2, characterized by rate constants λ and μ for switching between the states, and by the diffusion coefficients D_1 and D_2 . The associated jump process is defined by:

$$X(t + \Delta t) = X(t) + \begin{cases} \sqrt{2D_1 \Delta t} \eta & \text{w.p.} 1 - \lambda \Delta t \\ 0 & \text{w.p.} \lambda \Delta t \end{cases}$$

$$X(t + \Delta t) = X(t) + \begin{cases} \sqrt{2D_2 \Delta t} \eta & \text{w.p.} 1 - \mu \Delta t \\ 0 & \text{w.p.} \mu \Delta t \end{cases}$$
(7)

where the transition rates from states 1 to 2 are described by the switching equation:

$$1 \underset{\lambda}{\stackrel{\mu}{\rightleftharpoons}} 2$$

The probability density function (pdf) p_d for the displacement of a molecule depends on the state of the process and can be computed using Bayes' law:

$$Pr_{d} \{ |\Delta X| = u \} = Pr\{ |\Delta X| = u | X(t) \text{ and } X(t + \Delta t) \text{ in state } 1 \} q_{1}(t)$$

$$+Pr\{|\Delta X| = u|X(t) \text{ and } X(t + \Delta t) \text{ in state } 2\}q_2(t), \tag{8}$$

where using Bayes relation, for state i = 1 or 2,

 $q_i(t) = Pr\{X(t) \text{ and } X(t + \Delta t) \text{ in state } i\} = Pr\{X(t + \Delta t) \text{ in state } i|X(t) \text{ in state } i\}$

 $\times Pr\{X(t) \text{ in state } i\}.$

Because $p_i(t) = Pr\{X(t) \text{ in state } i\}$ is the solution to the Master equation:

$$\frac{d}{dt}p_1 = -\lambda p_1 + \mu p_2 \qquad (9)$$
$$\frac{d}{dt}p_2 = \lambda p_1 - \mu p_2. \qquad (10)$$

and

$$p_1(t) = \kappa + Ce^{-(\lambda+\mu)t}$$

$$p_2(t) = 1 - \kappa - Ce^{-(\lambda+\mu)t},$$

where $\kappa = \frac{\lambda}{\lambda + \mu}$ and *C* depend on the initial distribution, we have:

 $Pr{X(t + \Delta t)$ in state 1|X(t) in state $1} = 1 - \lambda \Delta t$, and

 $Pr\{X(t + \Delta t) \text{ in state } 2 | X(t) \text{ in state } 2\} = 1 - \mu \Delta t.$

We conclude that in two-dimensions, for a long-time *t* and a short-time Δt , using Eqs. 8 and 3, the pdf can be written as:

$$Pr_{d}\left\{\left|\Delta \mathbf{X}\right| = u\right\} = \kappa \frac{u}{\sigma_{1}^{2}} \exp\left(-\frac{u^{2}}{2\sigma_{1}^{2}}\right) + (1-\kappa)\frac{u}{\sigma_{2}^{2}} \exp\left(-\frac{u^{2}}{2\sigma_{2}^{2}}\right)$$
(11),

where $\sigma_k = 2D_k \Delta t$ and k = 1,2, and κ is the ratio of the time spent in a confined state. In the case of a three-state model, the pdf for the displacement of a molecule is:

$$Pr_{\text{switch}} \left\{ \Delta \boldsymbol{X} = \boldsymbol{u} \right\} = \kappa_1 \frac{u}{\sigma_1^2} \exp\left(-\frac{u^2}{2\sigma_1^2}\right) + \kappa_2 \frac{u}{\sigma_2^2} \exp\left(-\frac{u^2}{2\sigma_2^2}\right) + (1 - \kappa_1 - \kappa_2) \frac{u}{\sigma_3^2} \exp\left(-\frac{u^2}{2\sigma_3^2}\right),$$
(12)

where $\sigma_k = 2D_k\Delta t$ with k = 1, 2, and 3 and D_1 , D_2 , and D_3 are the three diffusion coefficients. Here κ_1 (respectively κ_2) is the fraction of time spent in state 1 (respectively state 2).

To test the appropriateness of a one-, two- or three-state models, we extracted the diffusion coefficients from Eqs. 11-12 and the associated rate constants. We then estimated the quality of fit of the different models by using the Bayesian Information Criterion (BIC) defined as:

$$BIC = (p+1)n(\ln n) \left[\ln \left(\frac{2\pi RSS}{n} \right) + 1 \right]$$
(13)

where p is the number of parameters of the model from Eq. 12, n is the number of data points, and *RSS* is the residual sum of squares between the model and the data.

Analysis of the 20 ms trajectories revealed that the data could be characterized by either a two or a three state model, consisting of a confined and either one or two unconfined states (see Table 1, below). Similarly, we evaluated 1, 2 and 3 state models for analysis of the 500 ms data consisting of a confined and either one or two sub-diffusive states (see Table 2, below). Although this showed that a three-state model could best describe both the 20 and 500 ms data, the analysis also revealed that the BIC values for the three-state model were only slightly larger, suggesting that a two-state model would be sufficient to explain the data.

	D	BIC	Kc	Ku	D _c	$D_{\rm u}$	BIC	Kc	K _{u1}	K _{u2}	Dc	D_{u1}	D_{u2}	BIC	Ndisps
CHD4	0.206	-1718	0.58	0.42	0.118	0.616	-2932	0.23	0.56	0.21	0.068	0.222	1.126	-3268	237358
CHD4-MBD3	0.234	-1707	0.54	0.46	0.121	0.649	-2971	0.29	0.48	0.23	0.085	0.271	1.134	-3093	85046
CHD4-GATA	0.17	-1750	0.51	0.49	0.088	0.382	-2986	0.26	0.57	0.17	0.061	0.199	0.856	-3158	138121
MBD3	0.147	-1743	0.51	0.49	0.076	0.319	-2966	0.28	0.59	0.13	0.054	0.181	0.813	-3096	75809
MTA2	0.194	-1786	0.45	0.55	0.096	0.384	-2866	0.16	0.62	0.22	0.056	0.192	0.715	-2794	13767
MTA2-MBD3	0.383	-1628	0.5	0.5	0.174	1.2	-2805	0.16	0.46	0.38	0.065	0.279	1.495	-2908	24283

Table 1. Parameter estimation from trajectories with $\Delta t = 20$ ms.

	D	BIC	Kc	Ku	D _c	Du	BIC	K _c	K _{u1}	K _{u2}	D _c	D _{u1}	D_{u2}	BIC	Ndisps
CHD4	0.005	-1874	0.55	0.45	0.003	0.01	-3089	0.16	0.68	0.16	0.002	0.005	0.017	-3170	91470
CHD4+DRB	0.003	-1796	0.46	0.54	0.002	0.006	-3036	0.24	0.63	0.13	0.001	0.004	0.014	-3062	34164
CHD4-MBD3	0.004	-1855	0.52	0.48	0.002	0.008	-3145	0.2	0.63	0.17	0.002	0.004	0.014	-3218	60964
MBD3	0.004	-1827	0.55	0.45	0.003	0.009	-3094	0.22	0.62	0.16	0.002	0.005	0.017	-3205	53176
MBD3+FK228	0.004	-1803	0.56	0.44	0.002	0.008	-3151	0.36	0.51	0.13	0.002	0.005	0.017	-3181	64328

Table 2. Parameter estimation from trajectories with $\Delta t = 500$ ms. In both Tables 1 and 2, D_c and D_u are the estimated apparent diffusion coefficients (whilst K_c and K_u are their relative proportions). The blue boxes indicate our preferred model based on BIC analysis.

In summary, the displacement analysis revealed that a two-state diffusion model should be appropriate to analyse NuRD complex dynamics: in both cases there was only a small improvement in the BIC for the three-state diffusion model, suggesting that the 20 ms trajectories can be characterised by confined (low diffusion coefficient D) and unconfined (high diffusion coefficient D) states, whilst the 500 ms data can be characterised by slow and fast sub-diffusive chromatin bound states. However, this analysis supported our later identification of a slow and two fast states in the 500 ms data (see Section 5, below).

2. Gaussian mixture model for classification and segmentation of trajectories into two states using four biophysical parameters (4P)

The Gaussian classification described in Section 1 above, is limited as it considers each displacement as being independent and then pools them, destroying any possible causality present in the trajectories. Moreover, because parts of our trajectories could correspond to molecules that are confined (i.e. chromatin bound) and other parts to when they are unconfined (or diffusing more freely), our trajectories exhibit a variety of behaviours characterised not only by slow and fast diffusion coefficients but also by smaller/larger anomalous exponents, the exploration of smaller/larger regions of space, and less/more directional movement. We therefore needed an algorithm that took into account these types of behaviour and not just changes in diffusion coefficient. Based on recent work to characterise such biophysical parameters⁵⁵, we present here a new algorithm that uses these parameters to classify and segment a trajectory into two states – when analysing 20 ms data the resulting sub-trajectories are confined (C) and unconfined (U) in the associated four dimensional space, whilst for the 500 ms data, we observe two confined states, slow- and fast-diffusing (see **Figure 2** in the main text). The method is based on a generalized Gaussian mixture model⁸⁷, where the input data is an ensemble of trajectories and the output is two ensembles of sub-trajectories.

2.1 Input data

We obtain an ensemble of *N* trajectories $X_i(k\Delta t)$, i = 1,...N, $k = 0, 1,...,n_i$, with an acquisition time step Δt , such that each trajectory consists of n_i discrete points in three dimensions.

2.2 Statistical features are extracted in a sliding window along the trajectories

To classify sub-trajectories as confined C or unconfined U, we used four physical parameters^{53,88}, computed along single trajectories. For a trajectory given by the successive points $X_i(k\Delta t)$, we used a sliding window W_k containing 2l + 1 points, centered at $X(k\Delta t)$, and defined as:

$$W_k^{(d)}(m\Delta t) = \{X_i^{(d)}((k-l+m)\Delta t), m = 0, \dots, 2l, \quad d = 1, 2 \text{ and } 3\},$$
(14)

where *d* is the space dimension. The sliding window $W_k(m\Delta t)$ is applied for $k = 1,...,n_i$ along a trajectory containing n_i points.

We now briefly discuss the four parameters we used for the classification based on previous work which showed that they can be used individually for the classification of single-molecule trajectories:

Anomalous exponent α : The anomalous exponent characterizes the motion of a stochastic particle on a particular time scale involving several time steps Δt . It is computed from the Mean-Square-Displacement (MSD) $\langle |X(t + \Delta t) - X(t)|^2 \rangle$ that behaves like Δt^{α} where Δt is small compared to the time of the process. A value of $\alpha = 1$ reflects Brownian motion, and an $\alpha > 1$ is called super-diffusion, which may represent dynamics containing an element of deterministic (ballistic) directed motion. Finally, an $\alpha < 1$ is sub-diffusive motion and has been used previously to study changes in chromatin compaction^{53,89,90}. To estimate α for each point *X*, we first compute the MSD *S_k* over the sliding window *W_k(m\Delta t*) classically defined by:

$$S_k(m) = \langle ||W_k(m\Delta t) - W_k(0))||^2 \rangle, m = 0, \dots, 2l_s$$

Where $\langle \cdot \rangle$ denotes the average where we use the intermediate point from 1 to m. To estimate the exponent $\alpha_i(k)$ for point $X(k\Delta t)$, we fitted the function $S_k(m)$ computed by summing over each displacement contributing to Eq. (15) with:

$$f_k(t) = \beta_i(k) t^{\alpha_i(k)}$$
(16)

where $\beta_i(k) > 0$. For the fit, we constrain the variable *t* in the ensemble $[0, \Delta t, ..., (2l+1)\Delta t]$.

Effective Diffusion coefficient D: We estimate the effective diffusion coefficient^{88,91} by computing the second statistical moment along the trajectories. We use the empirical estimator to estimate for each sliding window $W_k(m\Delta t)^{53}$:

$$D_{i}(k) = \frac{1}{2d\Delta t} \sum_{m=0}^{2l} (W_{k}((m+1)\Delta t)) - W_{k}(m\Delta t))^{2}$$
(17)

Length of confinement Lc. The length of confinement estimates the size of a domain where a trajectory is confined. It is computed empirically in the window W_k by:

$$Lc_i(k) = \frac{1}{3} \sum_{d=1}^{3} \sqrt{\frac{1}{2l} \sum_{m=0}^{2l} (W_k^{(d)}(m\Delta t) - \langle W_k^{(d)} \rangle)^2} = STD(W_k),$$
(18)

where:

$$\langle W_k^{(d)} \rangle = \frac{1}{2l} \sum_{m=0}^{2l} W_k^{(d)}(m\Delta t), d = 1, 2 \text{ and } 3.$$
[9]

It is the standard deviation of the sub-trajectory W_k , where the average position is $(\langle W_k^{(1)} \rangle, \langle W_k^{(2)} \rangle, \langle W_k^{(3)} \rangle)$.

Magnitude of the drift vector $V_i(k)$. To characterize the displacement of a trajectory between the beginning and the end of the sliding window W_k , we compute the magnitude $V_i(k)$ of the drift vector V_i for each dimension d = 1, 2, 3, ⁹¹ using the formula:

$$V_i^{(d)}(k) = \frac{1}{2l\Delta t} \sum_{m=0}^{2l-1} W_k^{(d)}((m+1)\Delta t) - W_k^{(d)}(m\Delta t)$$
(20)

The norm of the drift $V_i(\mathbf{k})$ is:

$$\|V_i\| = \sqrt{\sum_{d=1}^{3} (V_i^{(d)}(k))^2}.$$
(21)

In summary, we compute four parameters: the anomalous exponent α , the apparent diffusion coefficient *D*, the length of confinement *Lc*, and the magnitude of the drift vector V_i (k). These are all

computed for each point along a trajectory using a sliding window. The sliding window of 11 points was chosen by trial and error. Below this value, the anomalous exponent distributions for confined and unconfined molecules tended to merge suggesting that long trajectories are essential for reliable estimation of this parameter. Above this value, the diffusion coefficient histograms for confined and unconfined molecules tended to merge suggesting that transitions between these populations occur leading to averaging of the diffusion coefficients. In the next section, we use these four parameters for classification.

2.3 Classification of a discrete time point in a trajectory into either a confined (C) or an unconfined (U) state

To classify each time point $X((k - 1)\Delta t)$ for $k = 1,...,n_i$ of all trajectories indexed by i = 1,...,N into C and U classes, we first collected all the values from Eqs. 16-21, computed from each sliding window W_k (Eq. 14). This led to a total of $n_i \times 4$ parameters, that we organized in a matrix R_i associated with trajectory *i*:

$$R_{i} = [\alpha_{i}^{T}, D_{i}^{T}, Lc_{i}^{T}, \|V\|_{i}^{T}] = \begin{bmatrix} r_{i}(\Delta t) \\ r_{i}(2\Delta t) \\ \vdots \\ \vdots \\ r_{i}(n_{i}\Delta t) \end{bmatrix}, \qquad (22)$$

where T is the transpose operator, and $r_i(k\Delta t)$ are four dimensional vectors of the parameters (Eqs. 16-18, 21).

We then concatenate all feature matrices R_i (Eq. 22), i = 1,...N into the $\left(\sum_{i=1}^{N} n_i\right) \times 4$ general matrices **R**, defined by:

$$\boldsymbol{R} = \begin{bmatrix} R_1 \\ R_2 \\ \cdot \\ \cdot \\ R_N \end{bmatrix}.$$
(23)

and we normalize each column (feature) in $R^{(j)}$, j = 1,...,4 by subtracting its mean and dividing by its standarddeviation.

To separate the histograms of the four parameters into two independent classes, we constructed an unsupervised binary classifier using a two-component Gaussian mixture model in a four-dimensional space, corresponding to the four parameters Eqs. 16-21. The Gaussian mixture distribution p is a weighted sum of two multivariate Gaussian densities, defined as:

$$p(r|w_{C,U}, \mu_{C,U}, \Sigma_{C,U}) = w_C g(r|\mu_C, \Sigma_C) + w_U g(r|\mu_U, \Sigma_U),$$
(24)

where $w_{\rm C}, w_{\rm U}$ are the mixing weights, such that $w_{\rm C} + w_{\rm U} = 1$, and $g(r|\mu, \Sigma)$ are the Gaussian densities

$$g(r|\mu, \Sigma) = \frac{1}{(2\pi)^2 |\Sigma|^{1/2}} \exp\left(-\frac{1}{2}(r-\mu)^T \Sigma^{-1}(r-\mu)\right), \quad (25)$$

and Σ_{C} , Σ_{U} and μ_{C} , μ_{U} are the four-dimensional covariance matrices and mean vectors for components C and U, respectively. We then find the values of the parameters μ_{C} , μ_{U} , Σ_{C} , Σ_{U} and w_{C} , w_{U} of Eq. 24 which best separate the data as the maximal likelihood estimators of the density (Eq. 25), given the observed statistics in *R* (Eq. 23), by using the Expectation-Maximization algorithm⁹².

For each point $X_i(k\Delta t)$, with its associated feature vector $r_i(k\Delta t)$ (Eq. 23), we assign a label $n \in \{C, U\}$, based on the posterior probability *P* of the density Eq. 25, given by:

$$P(n|r_i(k\Delta t), w_n, \mu_n, \Sigma_n) = \frac{w_n g(r_i(k\Delta t)|\mu_n, \Sigma_n)}{w_c g(r_i(k\Delta t)|\mu_c, \Sigma_c) + w_{\mathcal{U}} g(r_i(k\Delta t)|\mu_{\mathcal{U}}, \Sigma_{\mathcal{U}})}$$
(26)

such that for each point:

$$X_{i}(k\Delta t) \in \begin{cases} \mathcal{C}, & Pr(\mathcal{C}|r_{i}(k\Delta t), w_{\mathcal{C}}, \mu_{\mathcal{C}}, \Sigma_{\mathcal{C}}) > Pr(\mathcal{U}|r_{i}(k\Delta t), w_{\mathcal{U}}, \mu_{\mathcal{U}}, \Sigma_{\mathcal{U}}) \\ \mathcal{U}, & \text{else.} \end{cases}$$

$$(27)$$

yielding a segmented trajectory where each time point is assigned as confined (C) or unconfined (U). The resulting ensembles of sub-trajectories can then be analysed further, in particular to estimate the four biophysical parameters (Eqs. 16-18, 21). In practice, if a point $X_i(k\Delta t) \in C$ or U is isolated, e.g. C between two neighbouring U class points, then we relabel it as the class of its two immediate time neighbours $(k - 1)\Delta t$ and $(k + 1)\Delta t$.

3. Simulations to validate our 4P classification algorithm

In this section, we estimate the accuracy of the classification algorithm (subsections 2.2-2.3) using simulated (synthetic) trajectories that can switch between two states C and U. The dynamics were described by an Ornstein-Uhlenbeck equation⁸⁸ (Eq. 28), and the generated trajectories were considered as a ground-truth ensemble, which we used to estimate the accuracy of the classification procedure developed in the previous section.

3.1 Simulation of a ground-truth ensemble of synthetic trajectories

We generated *N* trajectories $X_i(t)$, $i = 1,...,n_i$, which can switch between confined C and unconfined U states at any time *t*. The stochastic switching process is defined by:

$$\frac{dX(t)}{dt} = \begin{cases} -\kappa (X(t) - X(\tau)) + \sqrt{2D_{\mathcal{C}}} \frac{d\omega}{dt}, & X(t) \in \mathcal{C}; \\ \sqrt{2D_{\mathcal{U}}} \frac{d\omega}{dt}, & X(t) \in \mathcal{U}, \end{cases}$$
(28)

where $D_{\rm C}$, and $D_{\rm U}$ are the diffusion coefficients for the confined C and unconfined U states respectively, $d\omega/dt$ are standard three-dimensional Brownian motions, with a mean of zero and a standard deviation of one, while κ is the strength of a potential well that attracts the trajectory $X(t) \in {\rm C}$ to a fixed-point $X(\tau)$, which is the last position, before time *t* where the trajectory was in state U prior to switching to C:

$$\tau = \sup\{s < t, X(s) \in U \mid X(t) \in C\}.$$
 (29)

The transition probability between states (C, U) is defined by a Markov chain: The transient probability matrix P_{ij} between state *j* at time *s* + Δs and state *i* at time *s*, is given by:

$$P_{ij} = \begin{cases} \lambda \Delta s, & i = \mathcal{C}, j = \mathcal{U}; \\ 1 - \lambda \Delta s, & i = \mathcal{C}, j = \mathcal{C}; \\ \mu \Delta s, & i = \mathcal{U}, j = \mathcal{C}; \\ 1 - \mu \Delta s, & i = \mathcal{U}, j = \mathcal{U}. \end{cases}$$
(30)

The probabilities $P_{\rm C}(s)$ and $(P_{\rm U}(s))$ satisfy⁹³:

$$\begin{bmatrix} \dot{P}_{\mathcal{C}}(s) \\ \dot{P}_{\mathcal{U}}(s) \end{bmatrix} = \begin{bmatrix} -\lambda & \mu \\ \lambda & -\mu \end{bmatrix} \begin{bmatrix} P_{\mathcal{C}}(s) \\ P_{\mathcal{U}}(s) \end{bmatrix}$$
(31)

and the solution of Eq. 31 is:

$$\begin{bmatrix} P_{\mathcal{C}}(s) \\ P_{\mathcal{U}}(s) \end{bmatrix} = \frac{1}{\lambda + \mu} \begin{bmatrix} \lambda e^{-(\lambda + \mu)s} + \mu & \mu(1 - e^{-(\lambda + \mu)s}) \\ \lambda(1 - e^{-(\lambda + \mu)s}) & \lambda + \mu e^{-(\lambda + \mu)s} \end{bmatrix} \begin{bmatrix} P_{\mathcal{C}}(0) \\ P_{\mathcal{U}}(0) \end{bmatrix}$$
(32)

The simulation procedure was as follows: we initialized $X_i(0)$ in one of the states C or U. We used Euler's scheme to discretize Eq. 28 at a time step Δt . For each consecutive step $k\Delta t$, $k = 1,...,n_i$, we determined the state of the trajectory from the probabilities $P_C(\Delta t)$, $P_U(\Delta t)$ (Eq. 30), using the previous state as the initial condition:

$$\begin{bmatrix} P_{\mathcal{C}}(0) \\ P_{\mathcal{U}}(0) \end{bmatrix} = \begin{cases} [1,0]^T, & X_i((k-1)\Delta t) \in \mathcal{C}; \\ [0,1]^T, & X_i((k-1)\Delta t) \in \mathcal{U}. \end{cases}$$
(33)

3.2 Evaluating the classifier using ground-truth trajectories

To measure the accuracy of our classification algorithm (Sections 2.2-2.3), we generated N = 100 trajectories $Y_i(k\Delta t)$, i = 1,...,N, $k = 1,...,n_i$ using Eq. 29, where the length of a trajectory n_i was chosen randomly from a Poisson distribution with average length n = 100 time points, $\Delta t = 0.02s$, $\kappa = 2$, different values of D_C and D_U (see below), and twenty equally spaced values of λ , $\mu \in [0,1]$ to obtain ground-truth states $S_i(k\Delta t) \in [C,U]$ associated with each trajectory $Y_i(k\Delta t)$. We used the four-parameter classifier (section 2.3) to classify the trajectories $Y_i(k\Delta t)$ and add a label $C_i(k\Delta t) \in [C, U]$. To evaluate the similarity between the true class set S_i and the output of the classification classes C_i , we used an indicator function $\delta_i(k\Delta t)$, defined by:

$$\delta_i(k) = \begin{cases} 1, & S_i(k\Delta t) = C_i(k\Delta t); \\ 0, & \text{else,} \end{cases}$$
(34)

and then used Eq. 34 to define a parameter M that measures the accuracy of the classification along all trajectories. It was defined by:

$$M = \frac{\sum_{i=1}^{N} \sum_{k=1}^{n_i} \delta_i(k)}{\sum_{i=1}^{N} n_i}$$
(35)

We evaluated the accuracy *M* (Eq. 35) for an ensemble of trajectories generated with twenty values of $D_{\rm C} \in [0.002, 0.1]$ and twenty values of $D_{\rm U} \in [0.2, 1] \, \mu m^2 / s$, where we set the switching rates to be $\lambda = 0.8, \, \mu = 0.9$): we found that the classifier achieved above 90% accuracy in the majority of the parameter space, dropping to 76% for confined and unconfined diffusion constants where we restricted $Dc \in [0.02, 0.1]$, and $D_{\rm U} \in [0.1, 0.2]$

[see **Figure 1b** (left), below]. In addition, to test the robustness of the classification algorithm to the switching rates, we evaluated M (Eq. 35) for twenty values of the switching rates λ , $\mu \in [0, 1]$ where $D_{\rm C}$ and $D_{\rm U}$ were similar (0.008 and 0.01 μ m²/s, respectively) and obtained a very good accuracy – above 85% for all values of λ , μ [see **Figure 1b** (right), below].

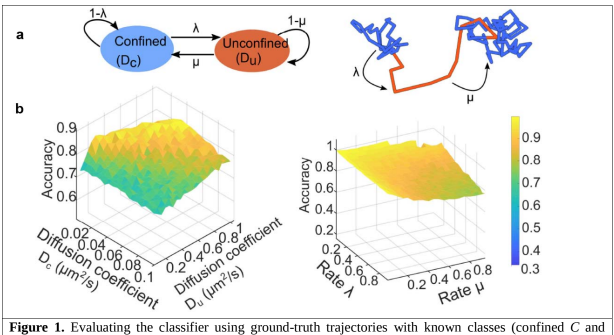


Figure 1. Evaluating the classifier using ground-truth trajectories with known classes (confined C and unconfined U). (a) The switching behaviour is described in Eq. 28, see above, where the class (C, U) is determined at each time point using a Markov chain (left) with switching rates λ , μ for the transitions between C and U and vice-versa, respectively (right). (b) After the ground truth set of trajectories was generated, we applied the classification procedure as described above, and computed the accuracy measure M (Eq. 35) defined as the fraction of correctly assigned classes by the segmentation algorithm according to the known classes in the ground-truth set. (Left) We computed M for trajectories generated using Eq. 29 with diffusion coefficients $D_c \in [0.002, 0.1] \, \mu m^2/s$ for class C and $D_U \in [0.2, 1] \, \mu m^2/s$ for U. We find that M > 0.76 for all tested values of D_C , D_U , while a well separated D_C , D_U resulted in M > 0.9. (Right) The accuracy M for switching rates λ , $\mu \in [0,1]$ and $D_C = 0.008$, $D_U = 0.01 \mu m^2/s$, M, resulted in M > 0.85 for all tested λ , μ .

In summary, when tested on simulated data, the classification algorithm was able to robustly classify

trajectories into confined and unconfined/sub-diffusive states.

4. Estimation of association and dissociation times from trajectories

After classifying the 20 ms data trajectories into two classes C and U (see section 2.3), we estimated the association and dissociation time constants as follows. The association time τ_A for trajectory X_i is defined as the time spent freely diffusing between confined chromatin-bound states:

$$t_{A} = \inf(t > 0; X_{i}(t) \in C \mid X_{i}(0) \in U).$$
(36)

The dissociation time τ_D is the time spent bound to chromatin between unconfined freely diffusing states:

$$t_{D} = \inf_{t} (t > 0; X_{i}(t) \in U \mid X_{i}(0) \in C).$$
(37)

To construct the histogram of these times, we reset the origin of time to zero after each association or dissociation event. In practice, for each partition of a trajectory X_i in a given ensemble, we collected the sequence τ_A^i of association times as the consecutive time points for which the trajectory spends in class U between being confined, as determined by the posterior probability *P* (Eq. 26). Equivalently, the dissociation times τ_D are those times in which the trajectory spends in confined state C between being unconfined. We fitted the histogram of association times and dissociation times with a single exponential⁹⁴:

$$f_{\rm A}(t) = \beta_{\rm D} \exp(-\mu t)$$
, and (38)
 $f_{\rm D}(t) = \beta_{\rm A} \exp(-\lambda t)$. (39)

The association and dissociation times are then obtained $\langle \tau_A \rangle = \frac{1}{\mu}$ by $\langle \tau_D \rangle = \frac{1}{\lambda}$ and.

5. Gaussian fitting of 500 ms exposure anomalous exponents

Although the algorithm we designed segments 500 ms exposure trajectories into slow-diffusing and fastdiffusing components, the distribution of the anomalous exponents in the fast-diffusing component showed that two peaks can clearly be identified by Gaussian fitting. Indeed, when we fitted 1, 2 or 3 Gaussians to the anomalous exponent distributions for chromatin bound NuRD complex subunits such as CHD4, MBD3 and MTA2 in wild-type ES cells, we found that 2 Gaussians were the best minimal model to account for the data – having the lowest BIC value (see Eq. 13 and **Extended Data Figure 8c**). This result was supported by the initial displacement analysis that we carried out (see Section 1), which also suggested that the fast components can be decomposed into two sub-populations. To ensure enough data had been collected to account for cell-to-cell heterogeneity, data was collated from 3 fields of view with around 6 cells in each field of view imaged, leading to a total of around 18 cells per condition. To assess the reproducibility of this result an additional 3 fields of view containing around 18 cells were collected for chromatin bound CHD4 on a different day and shown to have a similar anomalous exponent distribution (data not shown). Single molecule imaging of CHD4 in the *Mbd3*-ko was also shown to have a significantly different distribution when compared to both wild-type ES cell CHD4 datasets (p = 0.005 and p = 0.02 respectively).

6. Instructions for use of the code for classification and segmentation of trajectories into two diffusion states using four parameters (4P)

To use the classification code, click "Run" on MATLAB and then select all the relevant input files to include in the classification. The input format is a .csv file with the following columns:

Track number	frame	X	у	Z

The input parameters to define are as follows:

- 1. Exposure time for a frame e.g. 0.02 for 20 ms or 0.5 for 500 ms
 - 'dt',0.5,... % aquisition time interval [sec]
- Options to exclude short trajectories. This parameter also specifies the minimal length that a molecule must spend confined/unconfined for that sub trajectory to contribute to association/dissociation time calculations:
 - 'minNumPoints',2,... % (frames) threshold below which trajectories are discarded
- 3. Option to deal with misclassification errors for accurate final parameter calculation. To increase the accuracy of the calculation, we say that the period a molecule continuously spends confined/unconfined must be at least 7 frames long, i.e. at least 7 adjacent sliding windows must be assigned to the same class:

- 'minNumPointsLongTraj',7,... % minimal number of frames to be considered as non-confined long trajectory
- 4. Number of frames to include in the sliding window either side of a point on the trajectory
 - 'numMSDpoints',5,... % radius (number of points) to consider in the moving window around each point
 - For this study, we used optimised parameters of 5 frames before and after (11 frames in total)
 - Trade-off: longer sliding windows give better estimates of the parameters, but fail to detect

faster transitions

- 5. Option to classify by all 4 parameters or to choose a subset of them.
 - 'classifyUsingAlpha', true,... %use anomalous exponent in classification
 - 'classifyUsingDriftNorm', true, ... % use drift norm in classification
 - 'classifyUsingDiffusion',true,... % use diffusion constant in classification
 - 'classifyUsingLc', true,... % use Lc in classification
 - 'classifyUsingSpringConst', false,... % use Kc in classification

We currently do not classify by Kc, a spring constant useful for some analyses (see Shukron et al., Trends in

Genetics, 2019), but this is included in the code in case it is of use to others.

To generate outputs, set the following export function to true:

• 'exportResults', true,... % export results [true/false]

This will output the following:

1. .csv files for biophysical parameters calculated for sliding windows in each trajectory:

ID	frame	x	у	Z	confinementScore	confined	alpha	beta	diffusion	Lc	Кс	driftMagnitude
6	1778	27361	28318	-1700.2	7.29E-59	0	0.571	0.060	0.048	129.860	0.594	0.120
6	1779	27723	28205	-1648.9	2.41E-42	0	0.544	0.050	0.040	119.747	0.479	0.060
6	1780	27388	28286	-1383.2	1.48E-37	0	0.665	0.075	0.042	132.412	0.396	0.114
6	1781	27598	28117	-1419.6	3.65E-36	0	0.620	0.063	0.040	125.842	0.309	0.077
6	1782	27394	28360	-1747.8	1.14E-30	0	0.595	0.061	0.039	127.778	0.279	0.086
6	1783	27619	28289	-1548.6	1.27E-29	0	0.529	0.051	0.037	122.881	0.236	0.037
6	1784	27513	28323	-1602.4	1.13E-15	0	0.573	0.051	0.032	119.783	0.228	0.067
6	1785	27265	28367	-1316	5.54E-11	0	0.601	0.049	0.029	117.339	0.241	0.056
6	1786	27511	28317	-1431.7	2.19E-10	0	0.620	0.052	0.029	118.620	0.279	0.088
6	1787	27733	28230	-1649	1.81E-05	0	0.574	0.040	0.024	109.199	0.265	0.035
6	1788	27539	28282	-1666.7	0.002652771	0	0.625	0.041	0.022	107.263	0.283	0.075
6	1789	27412	28329	-1631.1	1.48E-08	0	0.627	0.046	0.026	114.530	0.334	0.042

- 2. .csv files for each biophysical parameter calculated for periods spent confined (note that this is the average parameter for a period spent confined, so if molecules spend a longer time confined, the estimate is based on values from more sliding windows)
- 3. .csv files for each biophysical parameter calculated for periods spent unconfined
- 4. .csv file of association times (time spent unconfined between periods spent confined) this excludes times where a molecule transitions for <10 frames as this may be a misclassification
- 5. .csv file for dissociation times (time spent confined between periods spent unconfined) this excludes times where a molecule transitions for <10 frames as this may be a misclassification

Additional outputs (set to 'true' by default) include:

6. Visualisation function in MATLAB to see all 3D trajectories but also all confined/unconfined trajectories outputted (sub-trajectory segments are coloured blue vs orange for confined vs unconfined segments).

- a. 'plotTrajectories',true,... % show all 3d trajectories [true/false] b. 'inspectTrajectories', true, ... % allows to inspect visually all, confined and unconfined trajectories
- 7. Options to plot the histograms of statistical parameters, association/dissociation times and trajectory duration times:
 - a. 'plotStatisticalParameters', true, ... % show the distribution of statistical parameters
 - b. 'plotTrajDurationHist', true, ...% plot trajectory duration histogram [true/false]
 - C. 'plotAssociationDissociationHist', true, ... % show the association dissociation histogram
- 8. Option to save figures of histograms outputted in Option 7
 - a. 'exportFigures', true, ... % export fig files

9. Export trajectories as PDB files to generate figures

- a. 'exportPDBtrajectories', true,... % export trajectories as pdb file
- 10. Changing the number of histogram fitting for association/dissociation times (default is 5)
 - a. 'numHistFitTrials',5,... % number histogram fitting trials
- 11. Option to classify by more than 2 Gaussian distributions (again not used in the paper but left as an option for other groups)

a. 'numberOfClasses',2,... % number of classes
 (confined/unconfined/others..)

In vitro biochemical assays of NuRD complex with and without nucleosomes

Drosophila PMMR and Human CHD4 were expressed in Sf21 cells and purified as described²⁴. Sf21 cells expressing Human GATAD2A-MBP were resuspended in 50 mM Tris-HCl pH 7.5, 1 M NaCl, 5 mM DTT, and $1 \times$ complete EDTA-free protease inhibitor cocktail (Roche), lysed by sonication and cleared by centrifugation at 50,000 *g* for 1 h. The supernatant was applied to amylose resin pre-equilibrated with lysis buffer and incubated for 2 h with rotation at 4 °C. The resin was washed with 20 X column volumes (CV) of lysis buffer and then eluted with 10 mM maltose in lysis buffer. Fractions containing hGATAD2A-MBP protein were concentrated and further purified by size exclusion chromatography using a Superose 6 Increase 3.2/300 column (GE Healthcare) equilibrated with 50 mM Tris-HCl pH 7.5, 150 mM NaCl, 5 % glycerol, and 1 mM DTT.

For pull-down experiments, purified protein was immobilised on MBP-Trap resin (ChromoTek) which was pre-equilibrated in pull-down buffer (50 mM HEPES pH 7.5, 300 mM NaCl, 1 mM DTT, and 5 % v/v glycerol) followed by incubation for 1 h with rotation at 4 °C. A sample of the 6% protein:bead mixture was retained as 'Input'. The resin was washed 3 times with pull-down buffer, then a washed 'beads' sample was retained for analysis on a 4-12% NuPAGE gel (Invitrogen).

Electrophoretic mobility shift assays (EMSAs) were performed with n3-Widom-78bp DNA or recombinant nucleosomes made with this template^{95,96} in 10 μ L of binding buffer (20 mM Hepes pH 7.5, 2 mM MgCl₂, 5% Glycerol, and 1 mM TCEP) with varying concentrations of the indicated proteins. The reaction mixtures were incubated at 30^oC for 30 min followed by centrifugation at 1000 x g. The resulting reaction mixtures were loaded onto 5% native polyacrylamide gels and run in 0.2 × TBE. Gels were stained with SYBR Gold (Invitrogen) and imaged using a Typhoon FLA 9000 (GE Healthcare).

dCas9-GFP imaging of enhancer loci

ES cells expressing dCas9 tagged with GFP were generated as described previously³. Briefly, MBD3inducible ES cells were transfected with the PB-TRE3G-dCas9-eGFP-WPRE-ubcp-rtTA-IRES-puroR vector containing a dual promoter backbone, with a TRE3G (Tet-on) promoter expressing GFP-tagged inactive dCas9 and the ubiquitin C promoter expressing the reverse tetracycline-controlled transactivator, rtTA, and a puromycin cassette via an IRES sequence. Puromycin-resistant ES cells were then selected for 7 days and doxycycline was added for 24 hours to induce expression of dCas9-GFP (through activation of the rtTA). Stable transfectants were then FACS sorted for low levels of GFP expression to select cells where only a few copies of the plasmid were integrated stably into the genome.

Prior to imaging, 1 µg/mL of doxycycline was added to ES cells for 24 hours to induce expression of low levels of dCas9-GFP. For imaging of *Tbx3* enhancer loci, three CARGO vectors in total expressing 36 gRNAs targeting the *Tbx3* enhancer were then transfected using lipofectamine 2000 (Invitrogen). The CARGO and dCas9-GFP expressing plasmids were kind gifts from the J. Wysocka lab. For imaging of *Nanog* enhancer loci, a custom designed gRNA was annealed with SygRNA® Cas9 Synthetic Modified tracrRNA (Sigma Aldrich TRACRRNA05N). The gRNA was designed such that a single gRNA sequence could be used to uniquely target a repetitive sequence close to the relevant enhancer (**Extended Data Figure 9b**). Cells were transfected with the tracr:crRNA complex using Lipofectamine 2000 (Invitrogen 11668019). In all cases, cells were transfected during passaging straight onto imaging dishes in Fluorobrite imaging medium as described above. After 24 hours, the media was replaced with fresh media and for +MBD3 samples, 4-hydrooxytamoxifen was added to a final concentration of 0.4 nM. All samples were then imaged after a further 24 hours.

2D tracking of genomic loci was carried out using oblique-angle illumination on a custom built 2D single-molecule tracking microscope as previously described⁹⁷. Briefly, an IX73 Olympus inverted microscope was used with circularly polarized laser beams aligned and focused at the back aperture of an Olympus 1.40 NA 100× oil objective (Universal Plan Super Apochromat, 100×, NA 1.40, UPLSAPO100XO/1.4). A 561 nm laser was used as a continuous wavelength diode laser light source. Oblique-angle illumination imaging was achieved by aligning the laser off axis such that the emergent beam at the sample interface was near-collimated and incident at an angle less than the critical angle ($\theta_c \sim 67^\circ$) for a glass/water interface. This generated a ~50 µm diameter excitation footprint. The power of the collimated 488 nm beam at the back aperture of the microscope was 100 W/cm². The lasers were reflected by dichroic mirrors which also separated the collected fluorescence emission from the TIR beam (Semrock, Di01-

R405/488/561/635). The fluorescence emission was collected through the same objective and then further filtered using a combination of long-pass and band-pass filters (BLP01-561R and FF01-587/35). The emission signal was projected onto an EMCCD (Photometrics, Evolve 512 Delta) with an electron multiplication gain of 250 ADU/photon operating in a frame transfer mode. The instrument was automated using the open-source software micro-manager (https://www.micro-manager.org) and the data displayed using the ImageJ software^{98,99}.

For image processing, PeakFit¹⁰⁰ was used to localise genomic loci from the images using the filter settings: "shiftFactor":1.0, "signalStrength":5.0, "minPhotons":30.0, "precisionThreshold":40.0, "minWidthFactor":0.5, "maxWidthFactor":0.5, "precisionMethod":"MORTENSEN". Trajectories were then tracked in 2D using custom python code for connecting foci in subsequent frames if they were within 500 nm (<u>https://github.com/wb104/trajectory-analysis</u>). Trajectories were classified as for single molecules using a Gaussian mixture model – see **Single molecule trajectory analysis**, above.

Enhancer-promoter DNA FISH (fluorescence in situ hybridisation)

FISH probes were prepared from mouse BAC library clones (Source Biosciences)¹⁰¹. BAC vector DNA was purified using the Qiagen Large Construct Kit. BAC DNA was labelled using Cy3 and Alexa Fluor 647 Nick Translation Labelling Kits (Jena Bioscience, PP-305S-CY3, PP-305S-AF647) and purified as previously described¹⁷. BAC probes were generated as follows:

Locus	BAC library code	Probe label
<i>Tbx3</i> promoter	BMQ-338M19	СуЗ
<i>Tbx3</i> enhancer	BMQ-309K22	Alexa Fluor 647
Sox2 promoter	BMQ-158G1	Cy3, Alexa Fluor 647
Sox2 enhancer	BMQ-53J20	Alexa Fluor 647
<i>Bmp4</i> promoter	BMQ-116K2	СуЗ
<i>Bmp4</i> enhancer	BMQ-249B4	Alexa Fluor 647

For two-colour DNA FISH, 2 x 10⁴ cells were seeded per well on microscope slides with removable 8 well silicone chambers (Ibidi 80841). Cells were fixed in 4% formaldehyde in PBS (Pierce[™] 28906, Thermo

Fisher Scientific) for 10 min at RT, followed by permeabilization in 0.3% Triton X-100 (Sigma-Aldrich X100) in PBS for 15 min at RT. After 3 washes in PBS, cells were incubated in pre-warmed 2 x saline-sodium citrate (SSC) with 100 ug/mL RNAse A (Qiagen 158922) for 1 hour at 37 °C. Chambers were removed and slides washed in 2 x SSC at RT. Slides were then dehydrated using 70% ethanol, 90% ethanol and 100% ethanol for 2 mins each and left to air dry. Cells were denatured in 70% deionised formamide (Sigma-Aldrich S4117) in 2 x SSC at 80 °C for 15 minutes. Slides were then again dehydrated quickly through ice-cold 70% ethanol, 90% ethanol RT, and 100% ethanol RT for 2 mins each and again left to air dry.

For each sample slide, 150 ng of Cy3-labelled BAC probe and 150 ng of AF647-labelled BAC probe were precipitated with 5 ug of Salmon Sperm DNA (Invitrogen 15632011) using 0.1 volumes of 3M sodium acetate (pH 5.2) and 2.5 volumes of 100% ethanol. Precipitated DNA was pelleted through centrifugation at 15000 x g for 20 minutes and resuspended in 50 uL hybridization buffer (50% formamide, 10% dextran sulfate (Sigma Aldrich 42867), 0.1% SDS, 2x SSC) through incubation for 1 hour at 37 °C. Probes were denatured at 80 °C for 10 mins and then transferred to 37 °C. Sample slides were overlaid with 50 μ L hybridization solution and covered with Parafilm, after which hybridization was allowed to occur at 37 °C overnight in a humidity chamber. The following day, the coverslip and hybridization solution were removed, and the slides washed four times in 2 x SSC at 40 °C for 3 mins each, then four times in 0.1 x SSC at 60 °C for 3 mins. Slides were cooled by washing in 4 x SSC at RT. After removing all the wash solution, cells were mounted in VECTASHIELD® Antifade Mounting Medium with DAPI (Vector Laboratories H-1200-10).

Sequential IF for dCas9-GFP and DNA FISH

ES cells expressing dCas9 tagged with GFP were transfected with either CARGO plasmids or a gRNA as described above (see "*dCas9-GFP imaging...*, above)". Fresh media was added after 24 hours and cells were fixed the following day in PBS containing 4% formaldehyde (PierceTM 28908, Thermo Fisher Scientific) for 10 min at RT. Cells were permeabilised in PBS containing 0.5% Triton X-100 (Sigma Aldrich X100) for 5 min, washed three times with PBS and then treated with blocking buffer (4% bovine serum albumin (Sigma-Aldrich, A9418) in 0.1% Triton-X 100 in PBS) for 30 min. Cells were incubated with GFP-Booster Alexa Fluor[®] 488 nanobody (ChromoTek, gb2AF488) in blocking buffer (1:1000) through incubation for 1 hour at RT. Samples were washed three times in PBS, each for 5 min.

Cells were post-fixed in PBS containing 3% formaldehyde (PierceTM, Thermo Fisher Scientific, 28908) for 10 min at RT, followed by re-permeabilization in 0.1 M HCl in 0.7% Triton X-100 in PBS for 10 min on ice. After 2 washes in 2 x SSC for 5 mins each, cells were incubated in pre-warmed 2 x SSC with 10 U/mL RNAse A (Qiagen 158922) for 1 hour at 37 °C. Slides were then equilibrated in 20% glycerol in PBS for 1 hour, followed by three consecutive freeze-thaw cycles using liquid nitrogen. After incubation for 1 hour in denaturing solution (50% formamide in 2 x SSC) at RT, slides were denatured at 70 °C for 5 mins and then washed several times in ice-cold 2 x SSC. Probes were denatured at 70 °C for 10 mins and then placed on ice to cool. Hybridisation solution was prepared with 50 ng of BAC probe and 10 µg Salmon Sperm DNA (Invitrogen 15632011) per 100 uL of hybridization buffer (50% formamide, 10% dextran sulfate, 1 mg/mL BSA, and 2 x SSC). Sample slides were overlaid with 25 uL hybridization solution per well and covered with Parafilm. After denaturation at 70 °C for 5 mins on a heat block, the slide was gradually cooled to 37 °C and hybridization solution were removed, and the slides washed three times in 2 x SSC at 40 °C for 5 mins each, then three times in 2 x SSC at RT for 5 mins. After removing all the wash solution, cells were mounted in VECTASHIELD® Antifade Mounting Medium with DAPI (Vector Laboratories).

IF and DNA-FISH image acquisition and analysis

Imaging was carried out using a Zeiss LSM Airy Scan 2 super-resolution microscope set for imaging of DAPI (405 nm laser, 0.8 %), Cy3 (514 nm laser, 10 %) and AF647 (639 nm laser, 90 %). Three stacks of horizontal plane images (3804×3804 pixels corresponding to 136.24×136.24 µm) with a z-step of 150 nm, were acquired for each field of view. CZI image files were then imported into IMARIS 9.6 (Bitplane) for three-dimensional modelling. Quantitative analysis of inter-probe distances within nuclei was carried out using the Surfaces and Spots modules of Imaris 9.6.

Software and code

1. Data collection

Microscope image acquisition

Micro-manager (https://www.micro-manager.org)

ImageJ software98,99

2. Data analysis

Hi-C analysis	NucProcess (https://github.com/tjs23/nuc_processing)						
	NucTools (<u>https://github.com/tjs23/nuc_tools</u>)						
	Juicer (https://github.com/aidenlab/juicer)						
	Cooler (https://cooler.readthedocs.io/en/latest/)						
	Lavaburst (https://github.com/nvictus/lavaburst)						
	Enhancer-promoter analysis (https://github.com/dhall1995/Acitivity-						
<u>By-Co</u>	ntact_Enhancer-Promoter_Link_Prediction						
Cut&Run analysis	R version 3.6.0						
	(https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/)						
	Read alignment ¹⁰² .						
	Deeptools v2.5.0 ⁷⁶ .						
2D single-molecule peak fitting	PeakFit ¹⁰⁰						
3D single-molecule peak fitting	easy-DHPSF software ⁸⁰						
Trajectory analysis	(https://github.com/wb104/trajectory-analysis)						
Gaussian mixture model classification	MATLAB v2016						
3D DNA FISH analysis	Imaris 9.6						

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Author contributions

SB, BDH, DHolcman and EDL designed the experiments. DL, SB, WB and TJS carried out the in-nucleus Hi-C data collection and processing. NR, WB, RR, XM and BDH carried out the Cut&Run data collection and processing. DHall, RR, XM and TJS analysed the preliminary ChIP-seq, Cut&Run and Hi-C data. LM, EB and LDC carried out preliminary ChIP-seq experiments. BDH, NR, JC, RF and SB made and characterised the cell lines used for the live-cell single-molecule imaging. ARC, AP, SFL and DK designed and built the 3D DH-PSF microscopes used for the live cell single molecule tracking experiments, and L-MN, SFL and DK designed and built the microscope for the 2D enhancer tracking. SB, DS and LHS carried out the live-cell 3D single-molecule imaging and analysed the data together with OS, PP, DK, BDH, EDL and DHolcman. SB, DS, GB, and KG carried out the live cell 2D tracking of enhancer movement. DS, SB, DL and MS carried out the immunofluorescence and DNA FISH experiments. WB developed the code to generate 3D single molecule tracks from the localisation data and dissociation time analysis, and OS, PP, AJ and DHolcman developed the code for segmentation of the 3D single-molecule trajectories. The *in vitro* biochemical experiments were carried out by WZ, JB, TD and TB, after purifying NuRD components and complexes expressed by AA, GC, IB and CS. SB, DHolcman and EDL wrote the manuscript with assistance from all the other authors.

Data Availability

The Hi-C and Cut&Run datasets reported in this study are available from the Gene Expression Omnibus (GEO) repository under accession code GSE147789. The *Chd4*, *Klf4*, *Mbd3* and *Mta2* Eos-Halo targeting constructs have been deposited with Addgene.

Code Availability

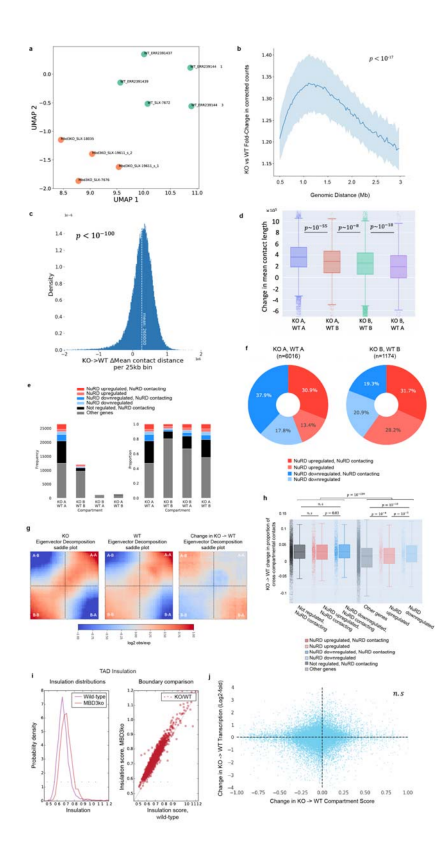
For the resubmission, we have made the data and analysis code available to the Editor and Reviewers using the following link to our group Dropbox:

https://www.dropbox.com/sh/t5dm7fqb3oxdq2b/AABwXmVhOo1QL5XyPn5Fec1_a?dl=0

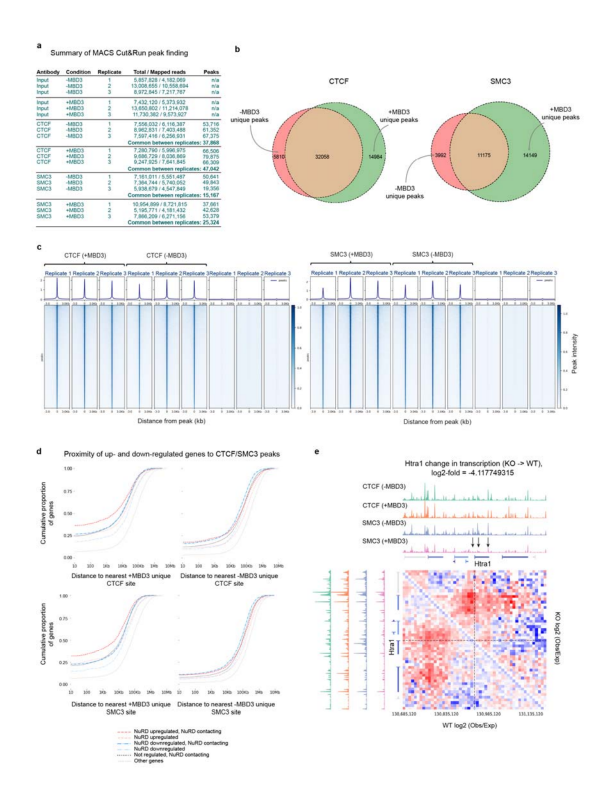
The password is: 5422801006. The code will be uploaded to Github upon acceptance of the paper and we will also make the source code openly available. We would also be happy to make the data available on request or to deposit it if a suitable site can be identified.

Extended Data

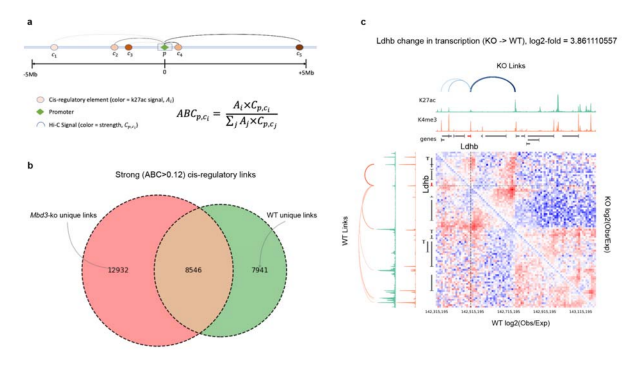
Extended Data Figure 1. MBD3-dependent assembly of the NuRD complex increases Mb-range chromosomal interactions. (a) UMAP projection plot showing the reproducibility between our own and previously published (E-MTAB-6591) Hi-C datasets for wild-type (blue) mouse ES cells grown in 2i/LIF conditions, and comparison with those from the Mbd3-ko (green). (b) The fold change in numbers of contacts at a range of genomic distances shows an increase in intermediate-range (~1 Mb) contacts in the presence of the intact NuRD complex $(p<10^{-17})$, Wilcoxon rank sum test). (c) Histogram showing the difference in mean contact distance for 25 kb genomic regions between the *Mbd3-ko* and wild-type cells. (d) Boxplot showing the changes in mean contact length for genomic regions that are in the A compartment in both wild-type and *Mbd3-ko* cells (blue), are in the B compartment in both conditions (green), that switch from A to B compartment in the presence of the intact NuRD complex (red), or that switch from B to A compartment in the presence of the intact NuRD complex (purple). (e) Bar and (f) pie charts showing the numbers of genes that remain or switch between compartments with the percentages of those genes that are up-/down-regulated. Saddle plots (g) show that there is an increase in inter-compartment contacts when going from the Mbd3-ko to wild-type cells, and thus A/B compartment mixing, and box plots (h) show that this is more noticeable in regions where NuRD-bound enhancers contact promoters. (i) Insulation scores derived from the contacts in the Hi-C data show a global decrease in TAD insulation in the presence of the intact NuRD complex. (j) A correlation plot shows that changes in A/B inter-compartment contacts do not correlate with changes in transcription.



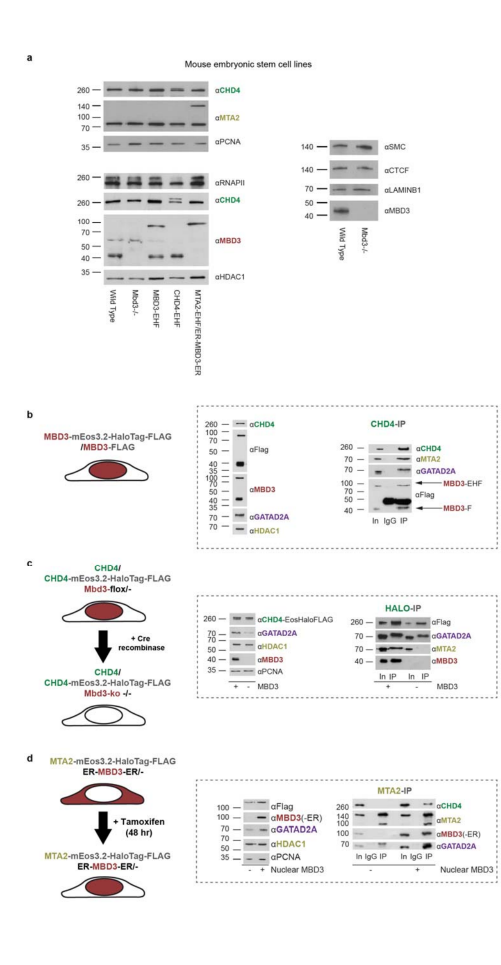
Extended Data Figure 2. Cut&Run experiments reveal that assembly of the intact NuRD complex leads to a redistribution of both CTCF and SMC3 (a Cohesin subunit) near NuRD-regulated genes. (a) Sequencing statistics and identification of peaks that are shared in the Cut&Run experiment replicates. (b) Venn diagrams showing the overlap in CTCF and SMC3 peaks in the absence and presence of MBD3. (c) (Top) Average peak profile and (Bottom) heatmap of CTCF and SMC3 signals +/-3 kb either side of identified peaks shows no significant changes in the overall levels of CTCF and Cohesin (SMC3) in the absence and presence of intact NuRD. (d) Cumulative probability plots of the distance from different categories of promoter to the nearest CTCF or Cohesin (SMC3) binding site found uniquely in either the presence (left) or absence (right) of MBD3. These plots are compared to genes with no transcriptional change (dotted orange lines) and they all have p-values of $< 1 \times 10^{-30}$ (Mann-Whitney U test). (e) Comparison of the CTCF and Cohesin (SMC3) Cut&Run data, plotted together with the observed/expected contact frequencies for a given genomic distance around the Htra1 gene. Positions where Cohesin is lost within the body of the Htral gene and upstream of its promoter are indicated with black arrows. The contact maps for the Mbd3-ko and wild-type cells are shown above/below the diagonal, respectively, and they are coloured according to their log-fold-change in contact frequency (red = increased, blue = decreased, intensity = absolute log fold change). The genes are coloured according to their log-fold-change in levels of expression (red = upregulated, blue = downregulated, intensity = absolute log fold change) and the black dotted lines mark the position of the *Htra1* promoter.



Extended Data Figure 3. Activity-by-contact model analysis of enhancer-promoter interaction strength in *Mbd3* knockout and wild-type ES cells. (a) Schematic of the activity-by-contact (ABC) analysis illustrating how the enhancer-promoter (E-P) interaction strength (ABC score) is calculated from the H3K4me3 signal at promoters and H3K27ac signal at nearby enhancers (defined using our previously published ChIP-seq data in wild-type and *Mbd3*-depleted ES cells)¹⁸ as well as the Hi-C signal strength. (b) Venn diagram showing the overlap between strong enhancer-promoter contacts identified in *Mbd3* knockout and wild-type ES cells. (c) Contact maps for a region around the *Ldhb* gene with changes in enhancerpromoter interactions and loops/TADs indicated. The maps for the *Mbd3-ko* and wild-type cells are shown above/below the diagonal, respectively, and they are coloured according to their log-fold-change in contact frequency relative to that expected theoretically at that particular distance (red = increased, blue = decreased, intensity = absolute log fold change). The black dotted lines mark the position of the *Ldhb* promoter.

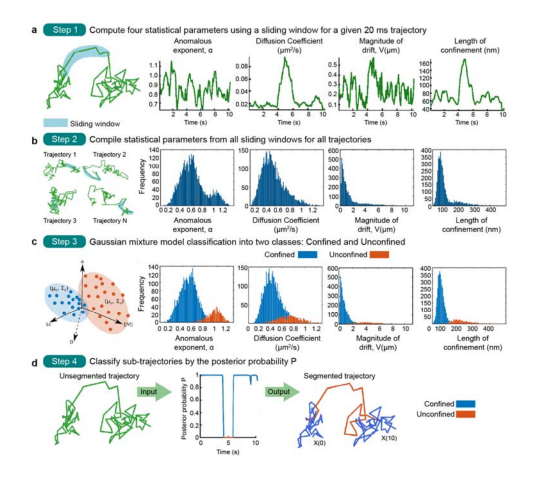


Extended Data Figure 4. Mouse embryonic stem cell lines expressing mEos3.2-HaloTag-FLAG tagged NuRD complex subunits. (a) Western blot comparison of expression of (Left) NuRD components, and (Right) CTCF and Cohesin in the cell lines used. Detailed schematic of the (b) *Mbd3*, (c) *Chd4* and (d) *Mta2* cell lines generated. CHD4 was tagged as previously described²⁴. MTA2 was tagged in ES cells expressing the ER-MBD3-ER (estrogen receptor-MBD3-estrogen receptor) fusion protein so that nuclear localisation of MBD3 is tamoxifen-inducible¹⁸. (Left) Expression of NuRD complex subunits was confirmed by western blot. Note that the stability of MTA2 and GATAD2A are both dependent upon MBD3, but that of CHD4 is not⁶⁸. (Right) Immunoprecipitation of either CHD4 or MTA2 confirms that the Eos-Halo-FLAG tags do not prevent association with other NuRD components, and that NuRD complex integrity is dependent upon the presence of MBD3.

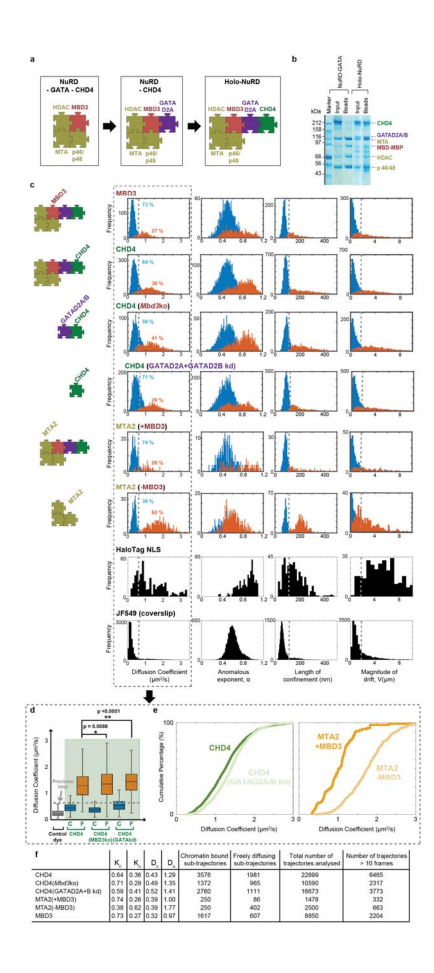


Extended Data Figure 5. Segmentation of single-molecule trajectories of CHD4-HaloTag-JF₅₄₉ (a) (Left)

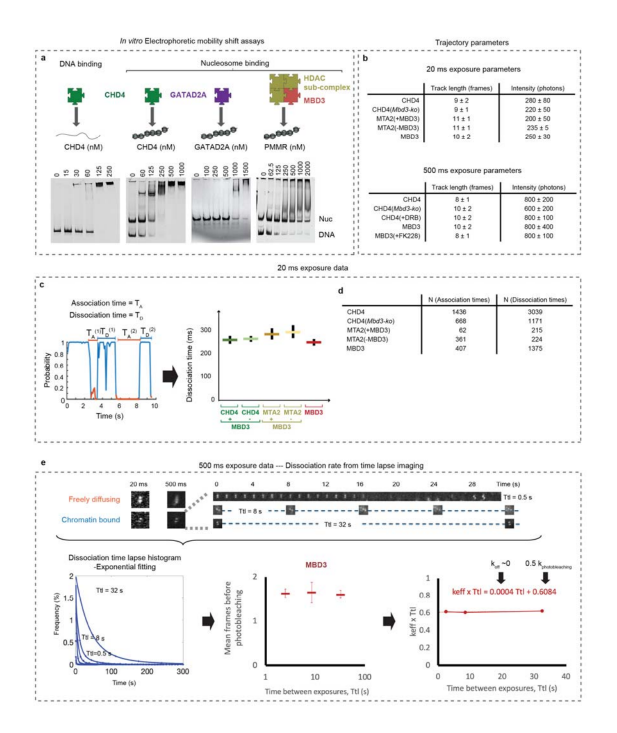
A single molecule trajectory, where we show an example sliding window (blue). (Right) Four biophysical parameters are extracted from sliding windows within this trajectory: the anomalous exponent α , the effective diffusion coefficient D, the norm||V|| of the mean velocity, and the length of confinement Lc, were all estimated from a sliding window of size w=10 Delta t (see Online Methods). (b) (Left) Several trajectories with example sliding windows (blue). (Right) Histograms of the biophysical parameters extracted in (a) from individual sub-trajectories are computed over the ensemble of all the recorded trajectories. (c) (Left) Scheme of the Gaussian mixture model (GMM) used to separate the histograms of the four parameters (four-dimensional feature space) into confined (C) and unconfined (U) populations (for 20 ms trajectories). The mean vectors μ C, μ U and covariance matrices Σ C, Σ U are estimated from the maximum likelihood. (Right) Separation of histograms in (b) into confined (blue) and unconfined (orange) populations. (d) Classification procedure applied to each trajectory, resulting in confined (blue) and unconfined (orange) sub-trajectories. The posterior probability P of the GMM (panel c) is computed on the four parameters for each point [Xi(k Δ t) \in C with P(k Δ t) > 1 – P(k Δ t) (blue); otherwise X(k Δ t) \in U (orange)] (see Online Methods). The result is a segmented trajectory where each time point is assigned as confined (C) or unconfined (U).



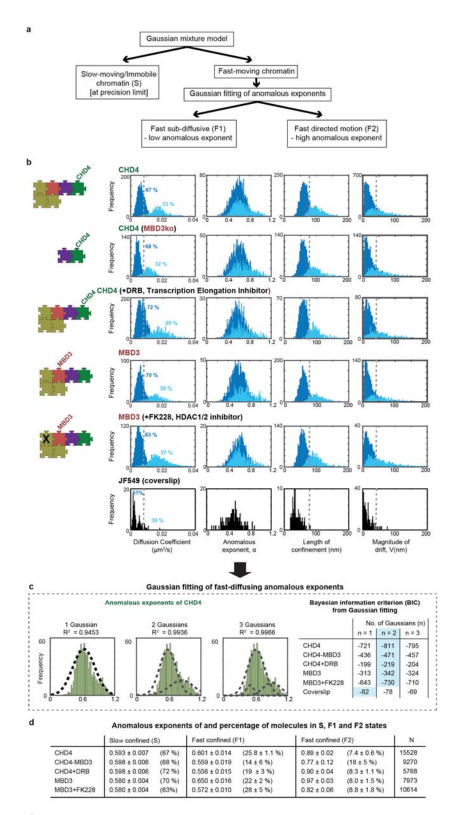
Extended Data Figure 6. In vitro and live cell single-molecule imaging experiments delineate holo-NuRD complex assembly. (a) Schematic of holo-NuRD complex assembly with GATAD2A linking MBD3 to the CHD4 remodeller. (b) Pull-down experiments of MBP-tagged MBD with and without GATAD2A confirm that GATAD2A is required for CHD4 to interact with the deacetylase sub-complex. (c) Distribution of the four biophysical parameters described in Figure 2 for 20ms exposure tracking of MBD3 and CHD4 in wild-type ES cells, as well as CHD4 in the absence of either MBD3 or GATAD2A/B. The data for MTA2 in the presence and absence of nuclear localised MBD3 are also shown. The grey dotted lines indicate the upper bound of the precision limit calculated at the 95 % confidence interval for an immobilised JF₅₄₉ dye control sample. HaloTag with a nuclear localisation sequence (HaloTag-NLS) is also shown as a control for a (mostly) freely diffusing molecule^{57,58}. (d) Box plot of apparent diffusion coefficients extracted from chromatin bound (C) and freely diffusing (F) CHD4 molecules in wild-type, Mbd3 knockout and GATAD2A/B knock-down ES cells (*p < 0.01, **p < 0.001, Kolmogorov-Smirnov test). The grey dotted line indicates the upper bound of the precision limit calculated at the 95 % confidence interval for an immobilised JF₅₄₉ dye control sample. (e) Cumulative distribution functions showing a higher diffusion coefficient for freely diffusing unconfined CHD4 upon removal of GATAD2A/B, and for freely diffusing MTA2 molecules upon removal of MBD3 from the nucleus. (f) Table showing the proportions (K) and estimated values of the apparent diffusion coefficients (D), as well as the number of chromatin bound and freely diffusing subtrajectories obtained from the total number of trajectories analysed. (NB - many trajectories were discarded as they were either too short for analysis or because they had a low probability of being classified as confined or unconfined.)



Extended Data Figure 7. The NuRD complex interacts with both DNA and nucleosomes, but only forms tight interactions through the CHD4 remodeller. (a) In vitro electrophoretic mobility shift assays confirm that CHD4 binds to both DNA and nucleosome core particles (NCPs) to form large complexes that only just enter the gel. GATAD2A alone shows low affinity binding to NCPs whilst the deacetylase complex interacts with DNA, but does not bind stably to NCPs. (b) Table showing the mean track length in frames, and mean photons detected per image frame, in 20 ms (Top) and 500 ms (Bottom) exposure trajectories. (c) (Left) Confinement probability allows collection of the association T_A or dissociation T_D times - defined respectively as the time a trajectory spends between periods of confined or unconfined motion. (Right) Dissociation times calculated using transitioning trajectories as periods of confined motion between two periods of unconfined motion (see also Figure 3). Error bars show 95 % confidence intervals. (d) Table with the number of single molecule tracks that were used to determine the association and dissociation times. (e) (Top) Example images demonstrating how long 500 ms exposures motion blur freely diffusing molecules, but allow detection and tracking of those that are chromatin bound. Images of single chromatin bound CHD4 molecules during time-lapse imaging with various dark times. (Bottom) Exponential fitting of time-lapse residence time histograms can be used to extract the photobleaching rate k_b and the effective dissociation rate k_{eff} . However, examination of the mean number of frames before photobleaching for MBD3, where the time between exposures is varied, shows that the results are completely dominated by photobleaching.



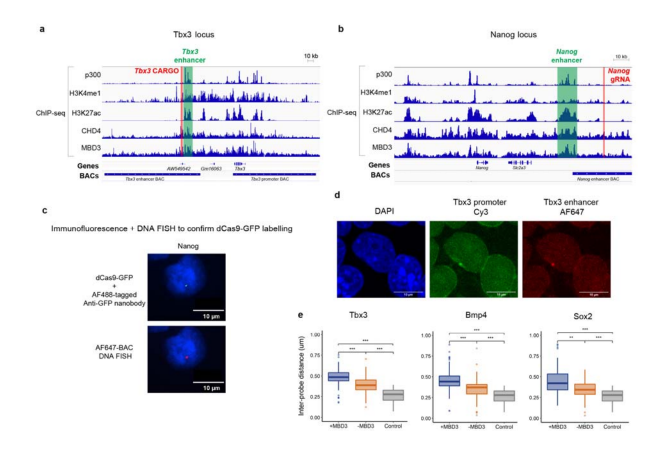
Extended Data Figure 8. Live cell single-molecule imaging experiments to study the chromatin-bound NuRD complex. (a) Schematic of the approach used for analysing the 500 ms exposure trajectories of chromatin bound NuRD complex subunits. First, a Gaussian mixture model is used to distinguish slowmoving/immobile and fast-moving chromatin bound sub-trajectories based on the 4 biophysical parameters (see also Figure 2 and Extended Data Figure 5). Then, Gaussian fitting of the anomalous exponent distributions of the fast-moving chromatin bound sub-trajectories revealed two fast-moving populations (F1 and F2) characterised by distinct anomalous exponents. (b) Distribution of the four biophysical parameters for 500 ms exposure tracking of: (i) chromatin bound CHD4 in wild-type ES cells, in the absence of MBD3, and in the presence of DRB (an inhibitor of transcriptional elongation); (ii) chromatin bound MBD3 in wild-type ES cells, and in the presence of the HDAC1/2-specific inhibitor FK228; and (iii) JF549 dye bound to the coverslip. The grey dotted lines indicate the upper bound of the precision limit calculated at the 95 % confidence interval for the immobilised JF_{549} dye control sample. (c) (Left) Fitting of 1, 2 or 3 Gaussians to the anomalous exponent distributions for fast moving chromatin bound CHD4 in wild-type ES cells – the R^2 values above indicate the goodness of fit. (Right) The Bayesian information criterion (BIC) was calculated for all the datasets shown in (b) to determine that 2 Gaussians are the best minimal model to account for the data - i.e. that has the lowest BIC value (light blue box). (d) Table summarising the changes in anomalous exponent of the slow and fast chromatin bound NuRD complex subunits in the presence and absence of MBD3, or in the presence of specific inhibitors. Errors given are for 95 % confidence intervals. (e) Table showing the proportions (K) and estimated values of the apparent diffusion coefficients (D), as well as the number of slow (S) and fast (F1+F2) diffusing sub-trajectories obtained from the total number of trajectories analysed. (NB – many trajectories were discarded as they were either too short for analysis or because they had a low probability of being classified as slow or fast.)



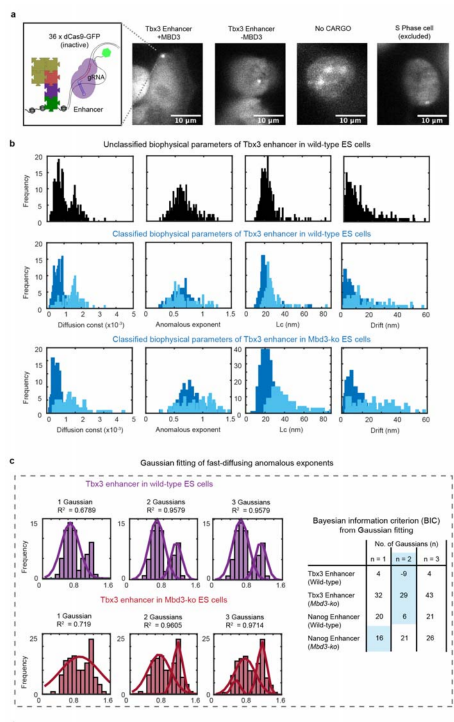
e	K.	к,	D,	D,	No. of slow (S) sub-trajectories	No. of fast (F1+F2) sub-trajectories	Total number of sub-trajectories analysed	Number of trajectories > 10 frames
CHD4	0.67	0.33	0.006	0.018	2045	1014	15528	3592
CHD4+DRB	0.72	0.28	0.007	0.018	838	384	5768	1401
CHD4(Mbd3ko)	0.68	0.32	0.006	0.015	1431	680	9270	2206
MBD3	0.7	0.3	0.006	0.016	1271	545	7973	1935
MBD3+FK228	0.63	0.37	0.006	0.018	1415	837	10614	2583

Extended Data Figure 9. DNA FISH validation to confirm NuRD-dependent changes in enhancer-

promoter compaction. Genomic locations of the (a) Tbx3 and (b) Nanog genes annotated with the locations to which dCas9-GFP was targeted using either CARGO vectors³ or a single gRNA that targets nearby genomic repeats (red lines). Locations of bacterial artificial chromosome (BAC) DNA FISH probes for the Tbx3 and Nanog enhancers are also indicated as are the targeted enhancers themselves (green). The corresponding ChIP-seq profiles¹⁷ indicate the binding of the NuRD complex subunits CHD4 and MBD3 as well as the location of active enhancers (determined from the ChIP-seq profiles for H3K27ac, H3K4me1 and p300). (c) Representative confocal images showing co-localisation of dCas9-GFP (labelled using AF488tagged anti-GFP nanobody) and AF647-BAC DNA FISH probes targeting the Nanog enhancer. (d) Representative confocal images of Cy3-labelled BAC DNA FISH probes targeting the Tbx3 promoter and AF647-labelled BAC DNA FISH probes targeting the Tbx3 enhancer. (e) Boxplots showing the enhancerpromoter distances in MBD3-inducible ESCs with and without tamoxifen: +MBD3 (blue) and -MBD3 ESCs (orange) respectively. There is a significant increase in enhancer-promoter distance in the presence of intact NuRD for *Tbx3* (+MBD3, n = 70; -MBD3, n = 101), *Bmp4* (+MBD3, n = 172; -MBD3, n = 71) and *Sox2* (+MBD3, n = 42; -MBD3, n = 50) (**p<0.01, ***p < 10^{-5} , Kolmogorov-Smirnov test). To estimate the precision limit of the experiment, control samples were generated in which the distance was measured for the Sox2 enhancer labelled with both Cy3 and AF647 (grey, n = 32).



Extended Data Figure 10. 2D dCas9-GFP tracking of active enhancers. (a) Representative images of 36 gRNAs targeted to the *Tbx3* enhancer in the presence or absence of MBD3 with a negative control expressing no gRNAs. (Right) An example of a cell with a doublet indicating that it is in S phase (that was excluded from the analysis) is also shown. (b) Distribution of the four biophysical parameters extracted from sliding windows within the 2D single-molecule trajectories of dCas9-GFP bound at the *Tbx3* enhancer in wild-type ES cells imaged using 500 ms exposures – (Top) before and (Middle) after classification based on the anomalous exponent α , the apparent diffusion coefficient D, the length of confinement Lc, and the drift magnitude, norm $\|V\|$ of the mean velocity. (Bottom) Distribution of the four biophysical parameters after classification for the *Tbx3* enhancer in *Mbd3-ko* cells. (c) (Left) Fitting of 1, 2 or 3 Gaussians to the fast-moving anomalous exponent distributions of the *Tbx3* enhancer tracked in either wild-type ES (Top) or *Mbd3-ko* cells (Bottom) – the R² values above indicate the goodness of fit. (Right) The Bayesian information criterion (BIC) was calculated for all the datasets to determine which number of Gaussians best modelled the data – that with the lowest BIC value (light blue box). (d) Table showing the Gaussian fitted anomalous exponent values for slow- and the fast-moving chromatin bound dCas9-GFP at both enhancers tracked in the presence and absence of MBD3.





Anomalous exponents of and percentage of molecules in S, F1 and F2 states

	Slow confined (S)	Fast confined (F1)	Fast confined (F2)	N
Tbx3 Enhancer Wild-type	0.675 ± 0.008 (71.73%)	0.635 ± 0.001 (18.47%)	1.071 ± 0.001 (9.80%)	237
Tbx3 Enhancer Mbd3-ko	0.754 ± 0.004 (57.49%)	0.621 ± 0.007 (21.39%)	1.075 ± 0.007 (21.12%)	287
Nanog Enhancer Wild-type	0.720± 0.027 (61.17%)	0.677 ± 0.025 (21.48%)	1.087 ± 0.013 (17.35%)	546
Nanog Enhancer Mbd3-ko	0.773 ± 0.029 (58.08%)	0.712 ± 0.028 (20.38%)	1.128 ± 0.018 (21.54%)	229

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