#### Title

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- Full title: Coupling chromatin structure and dynamics by live super-resolution imaging
- Short title: Coupling chromatin structure and dynamics

## 5 Authors

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## 19 Abstract

Chromatin conformation regulates gene expression and thus constant remodeling of chromatin 20 structure is essential to guarantee proper cell function. To gain insight into the spatio-temporal 21 organization of the genome, we employ high-density photo-activated localization microscopy and 22 23 deep learning to obtain temporally resolved super-resolution images of chromatin in vivo. In combination with high-resolution dense motion reconstruction, we confirm the existence of 24 elongated ~ 45 to 90 nm wide chromatin 'blobs', which appear to be dynamically associating 25 chromatin fragments in close physical and genomic proximity and adopt TAD-like interactions in 26 the time-average limit. We found the chromatin structure exhibits a spatio-temporal correlation 27 extending ~ 4  $\mu$ m in space and tens of seconds in time, while chromatin dynamics are correlated 28 over  $\sim 6 \ \mu m$  and outlast 40 s. Notably, chromatin structure and dynamics are closely interrelated. 29 which may constitute a mechanism to grant access to regions with high local chromatin 30 31 concentration.

#### 32 Introduction

The three-dimensional organization of the eukaryotic genome plays a central role in gene regulation 33 (1-3). Its spatial organization has been prominently characterized by molecular and cellular 34 approaches including high-throughput chromosome conformation capture (Hi-C) (4) and 35 fluorescent in situ hybridization (FISH) (5). Topologically associated domains (TADs), genomic 36 regions that display a high degree of interaction, were revealed and found to be a key architectural 37 feature (6). Direct 3D localization microscopy of the chromatin fiber at the nanoscale (7) confirmed 38 the presence of TADs in single cells but also, among others, revealed great structural variation of 39 chromatin architecture (8, 9). To comprehensively resolve the spatial heterogeneity of chromatin, 40 super-resolution microscopy must be employed. Previous work showed that nucleosomes are 41 distributed as segregated, nanometer-sized accumulations throughout the nucleus (10-13) and that 42 43 the epigenetic state of a locus has a large impact on its folding (14, 15). However, to resolve the fine structure of chromatin, high labeling densities, long acquisition times and, often, cell fixation 44 are required. This precludes capturing dynamic processes of chromatin in single live cells, yet 45 chromatin moves at different spatial and temporal scales. 46

The first efforts to relate chromatin organization and its dynamics were made using a combination 47 of Photo-activated Localization Microscopy (PALM) and tracking of single nucleosomes (16). It 48 could be shown that nucleosomes mostly move coherently with their underlying domains, in 49 accordance with conventional microscopy data (17); however, a quantitative link between the 50 51 observed dynamics and the surrounding chromatin structure could not yet be established in realtime. Although it is becoming increasingly clear that chromatin motion and long-range interactions 52 are key to genome organization and gene regulation (18), tools to detect and to define bulk 53 chromatin motion simultaneously at divergent spatio-temporal scales and high resolution are still 54 missing. 55

Here we apply deep learning-based photo-activated localization microscopy (Deep-PALM) for 56 temporally resolved super-resolution imaging of chromatin in vivo. Deep-PALM acquires a single 57 resolved image in a few hundred milliseconds with a spatial resolution of  $\sim 60$  nm. We observed 58 elongated ~ 45 to 90 nm wide chromatin domain 'blobs'. Employing a computational chromosome 59 model, we inferred that blobs are highly dynamic entities, which dynamically assemble and 60 disassemble. Consisting of chromatin in close physical and genomic proximity, blobs nevertheless 61 adopt TAD-like interaction patterns when chromatin confirmations are averaged over time. Using 62 a combination of Deep-PALM and high-resolution dense motion reconstruction (17), we 63 simultaneously analyzed both structural and dynamic properties of chromatin. Our analysis 64 emphasizes the presence of spatio-temporal cross-correlations between chromatin structure and 65 dynamics, extending several micrometers in space and tens of seconds in time. Furthermore, 66 extraction and statistical mapping of multiple observables from the dynamic behavior of chromatin 67 blobs shows that local chromatin density regulates local chromatin dynamics. 68

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#### 70 **Results**

#### 71 **Deep-PALM reveals dynamical chromatin remodeling in living cells.**

Super-resolution imaging of complex and compact macromolecules such as chromatin requires 72 dense labeling of the chromatin fiber in order to resolve fine features. We employ Deep-STORM, 73 a method which uses a deep convolutional neural network (CNN) to predict super-resolution images 74 from stochastically blinking emitters (19) (Figure 1A; Materials and Methods). The CNN was 75 trained to specific labeling densities for live-cell chromatin imaging using a photo-activated 76 fluorophore (PATagRFP); we therefore refer to the method as Deep-PALM. We chose three 77 labeling densities 4, 6 and 9 emitters per  $\mu m^2$  per frame in the ON-state to test, based on the 78 comparison of simulated and experimental wide field images (Supplementary Figure 1A). The 79 CNN trained with 9 emitters per  $\mu$ m<sup>2</sup> performed significantly worse than the other CNNs and was 80

81 thus excluded from further analysis (Supplementary Figure 1B; Materials and Methods). We applied Deep-PALM to reconstruct an image set of labeled histone protein (H2B-PATagRFP) in 82 human bone osteosarcoma (U2OS) cells using the networks trained on 4 and 6 emitters per  $\mu m^2$  per 83 frame (Materials and Methods). A varying number of predictions by the CNN of each individual 84 frame of the input series were summed to reconstruct a temporal series of super resolved images 85 (Supplementary Figure 1C). The predictions made by the CNN trained with 4 emitters per  $um^2$ 86 show large spaces devoid of signal intensity, especially at the nuclear periphery, making this CNN 87 inadequate for live-cell super-resolution imaging of chromatin. While collecting photons from long 88 89 acquisitions for super-resolution imaging is desirable in fixed cells, Deep-PALM is a live imaging approach. Summing over many individual predictions leads to considerable motion blur and thus 90 loss in resolution. Quantitatively, the Nyquist criterion states that the image resolution  $R = 2/\sqrt{\tau\rho}$ 91 depends on  $\rho$ , the localization density per second and the time resolution  $\tau$  (20). In contrast, motion 92 blur strictly depends on the diffusion constant D of the underlying structure  $R = \sqrt{4D\tau}$ . There is 93 thus an optimum resolution due to the tradeoff between increased emitter sampling and the 94 95 avoidance of motion blur, which was at a time resolution of 360 ms for our experiments (Figure 1B; Supplementary Figure 1D). Super-resolution imaging of H2B-PATagRFP in live cells at this 96 temporal resolution shows a pronounced nuclear periphery while fluorescent signals in the interior 97 98 vary in intensity (Figure 1C). This likely corresponds to chromatin-rich and chromatin-poor regions 99 (13). These regions rearrange over time, reflecting the dynamic behavior of bulk chromatin. 100 Chromatin-rich and chromatin-poor regions were visible not only at the scale of the whole-nucleus, 101 but also at the resolution of a few hundred nanometers (Figure 1D). Within chromatin-rich regions, the intensity distribution was not uniform but exhibited spatially segregated accumulations of 102 labeled histones of variable shape and size, reminiscent of nucleosome clutches (10), nanodomains 103 (14, 16) or TADs (21). At the nuclear periphery, prominent structures arise. Certain chromatin 104 structures could be observed for  $\sim 1$  s, which underwent conformational changes during this period 105

(Figure 1E). The spatial resolution at which structural elements can be observed (Materials and Methods) in time-resolved super-resolution data of chromatin was  $63 \pm 2$  nm (Figure 1E), slightly more optimistic than the theoretical prediction (Figure 1B) (22). Thus, Deep-PALM identifies spatially heterogeneous coverage of chromatin as previously reported (10, 13, 14, 16, 21). We further monitor chromatin temporally at nanometer scale in living cells.

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# 112 Chromatin appears in elongated nanometer-sized blobs with a non-random spatial 113 distribution.

To quantitatively assess the spatial distribution of H2B, we developed an image segmentation 114 scheme (Materials and Methods, Supplementary Figure 2) which allowed us to segment spatially 115 separated accumulations of H2B signal with high fidelity (Supplementary Note 1, Supplementary 116 Figure 3, Supplementary Figure 4). Applying our segmentation scheme,  $\sim 10,000$  separable 117 elements, blob-like structures were observed for each super-resolved image (166 resolved images 118 per movie; Figure 2A). To elucidate their origin and formation, we used a transferable 119 computational model introduced by Qi et al. (23), which is based on one-dimensional genomics 120 and epigenomics data, including histone modification profiles and CTCF binding sites. Super-121 resolution images were generated from the modeled chromosomes. Within these images we 122 identified and characterized 'chromatin blobs' analogously as for experimental data (Materials and 123 Methods; Figure 2B). 124

For both imaged and modeled chromatin, we first computed the k<sup>th</sup> nearest neighbor distance (NND; centroid-to-centroid) distributions, taking into account the nearest 1<sup>st</sup> to 40<sup>th</sup> neighbors (Figure 2C, blue to red). Centroids of nearest neighbors are  $(95 \pm 30)$  nm (mean  $\pm$  standard deviation) apart, consistent with previous super-resolution images of chromatin in fixed cells (*14*) and slightly further than what was found for clutches of nucleosomes (*10*). The envelope of all NND distributions (Figure 2C, black line) shows several weak maxima at ~ 95 nm, 235 nm, 335 nm, and

450 nm, which roughly coincide with the peaks of the 1<sup>st</sup>, 7<sup>th</sup>, 14<sup>th</sup> and 25<sup>th</sup> nearest neighbors respectively (Figure 2C, red dots). In contrast, simulated data exhibit a prominent first nearest neighbor peak at slightly smaller distance and higher-order NND distribution decay quickly and appear washed out (Figure 2D). This hints towards greater levels of spatial organization of chromatin *in vivo*, which is not readily recapitulated in the employed state-of-the-art chromosome model.

Next, we were interested in the typical size of chromatin blobs. Their area distribution (Figure 2E) 137 fit a log-normal distribution with parameters  $(3.3 + 2.8) \cdot 10^{-3} \,\mu\text{m}^2$  (mean + standard deviation), 138 which is in line with the area distribution derived from modeled chromosomes. Notably, blob areas 139 vary considerably as indicated by the high standard deviation and the prominent tail of the area 140 distribution towards large values. Following this we calculated the eccentricity of each blob to 141 resolve their shape (Figure 2F). The eccentricity is a measure of the elongation of a region reflecting 142 the ratio of the longest chord of the shape and the shortest chord perpendicular to it (Figure 2F, 143 illustrated shapes at selected eccentricity values). The distribution of eccentricity values shows an 144 accumulation of values close to 1, with a peak value of  $\sim 0.9$ , which shows that most blobs have an 145 elongated, fiber-like shape and are not circular. In particular, the eccentricity value of 0.9 146 corresponds to a ratio between the short and long axis of the ellipse of  $\sim \frac{1}{2}$  (Materials and Methods), 147 which results, considering the typical area of blobs, in roughly 92 nm long and 46 nm wide blobs 148 149 on average. The length coincides with the value found for the typical nearest-neighbor distance (Figure 2C;  $(95 \pm 30)$  nm). However, due to the segregation of chromatin into blobs, their 150 elongated shape and their random orientation (Figure 2A), the blobs cannot be closely packed 151 throughout the nucleus. We find that chromatin has a spatially heterogeneous density, occupying 5 152 - 60% of the nuclear area (Supplementary Figure 5A, B), which is supported by a previous electron 153 microscopy study (24). The blob dimensions also fall within previously determined size ranges (10, 154 14), confirming the existence of spatially segregated chromatin structures in the sub-100 nm range. 155

#### 156 Chromatin blobs identified by Deep-PALM are coherent with sub-TADs

Due to the projection of the nuclear volume to the imaging plane, the observed blobs could simply 157 be overlays of distant, along the one-dimensional genome, non-interacting genomic loci. To 158 exclude this possibility, we analyzed the gap length along the simulated chromosome between 159 beads belonging to the same blob. The analysis showed that the blobs are mostly made of 160 consecutive beads along the genome, thus implying an underlying domain-like structure, similar to 161 TADs (Figure 3A). Using the affiliation of each bead to an intrinsic chromatin state of the model 162 (Figure 3B), it became apparent that blobs along the simulated chromosome consisting mostly of 163 active chromatin are significantly larger than those formed by inactive and repressive chromatin 164 (Figure 3C). These findings are in line with experimental results (15) and results from the 165 simulations directly (23), thereby validating the projection and segmentation process. 166

Since chromatin is dynamic in vivo and in computer simulations, each bead can diffuse in and 167 out of the imaging volume from frame to frame. We estimated that, on average, each bead spent 168 approximately 1.5 s continuously within a slab of 200 nm thickness (Figure 3D). Furthermore, a 169 bead is on average found only  $0.55 \pm 0.33$  s continuously within a blob, which corresponds to 1 - 1170 2 experimental super-resolved images (Figure 3D). These results suggest that chromatin blobs are 171 highly dynamic entities, which usually form and dissemble within less than one second. We thus 172 constructed a time-averaged association map for the modeled chromosomes, quantifying the 173 frequency at which each locus is found with any other locus within one blob. The association map 174 is comparable to interaction maps derived from Hi-C (Figure 3E). Strikingly, inter-loci association 175 and Hi-C maps are strongly correlated, and the association map shows similar patterns as those 176 identified as TADs in Hi-C maps, even for relatively distant genomic loci (> 1 Mbp). A similar 177 TAD-like organization is also apparent when the average inverse distance between loci is 178 considered (Figure 3F, upper panel), suggesting that blobs could be identified in super-resolved 179 images due to the proximity of loci within blobs in physical space. Using the computational 180

chromosome model, we conclude that chromatin blobs identified by Deep-PALM are mostly made of continuous regions along the genome and cannot be attributed to artifacts originating from the projection of the three-dimensional genome structure to the imaging plane. The blobs associate and dissociate within less than one second, but loci within blobs are likely to belong to the same TAD. Their average genomic content is 75 kb, only a fraction of typical TAD lengths in mammalian cells (average size 880 kb) (*6*), suggesting that blobs likely correspond to sub-TADs or TAD nanocompartments (*21*).

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#### 189 Super-resolution chromatin dynamics using Deep-PALM

To quantify the experimentally observed chromatin dynamics at the nanoscale, down to the size of 190 one pixel (13.5 nm), we used a dense reconstruction of flow fields, Optical Flow (OF; Figure 4A, 191 Materials and Methods), which was previously used to analyze images taken on confocal (25, 26), 192 and Structured Illumination Microscopes (13). We examined the suitability of OF for super-193 resolution based on single molecule localization images using simulations. We find that the 194 accuracy of OF is slightly enhanced on super-resolved images, compared to conventional 195 fluorescence microscopy images (Supplementary Note 2, Supplementary Figure 6). Experimental 196 197 super-resolution flow fields are illustrated on the basis of two subsequent images, between which the dynamics of structural features are apparent to the eve (Figure 4B-C). On the nuclear periphery, 198 connected regions spanning up to ~ 500 nm can be observed (Figure 4B (i-ii), marked by arrows). 199 200 These structures are stable for at least 360 ms but move from frame to frame. The flow field is shown on top of an overlay of the two super-resolve images and color-coded (Figure 4B (iii), the 201 intensity in frame 1 is shown in green, the intensity in frame one is shown in purple, co-localization 202 of both is white). Displacement vectors closely follow the redistribution of intensity from frame to 203 204 frame (roughly from green to purple). Similarly, structures within the nuclear interior (Figure 4C)

can be followed by eye, thus further validating and justifying the use of a dense motion
 reconstruction as a quantification tool of super-resolved chromatin motion.

Using Optical Flow fields, we linked the spatial appearance of chromatin to their dynamics. Effectively, the blobs were characterized with two structural parameters (NND and area) and their flow magnitude (Figure 4D). Supplementary Movie 1 shows the time evolution of those parameters for an exemplary nucleus. Blobs at the nuclear periphery showed a distinct behavior from those in the nuclear interior. In particular, the periphery exhibits a lower density of blobs, but those appear slightly larger and are less mobile than in the nuclear interior (Supplementary Figure 7), in line with previous findings using conventional microscopy (*26*).

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#### 215 Chromatin structure and dynamics are linked

To further elucidate the relationship between chromatin structure and dynamics, we analyzed the 216 correlations between each pair of parameters in space and time. Therefore, we computed the auto-217 and cross-correlation of parameter maps with a given time lag across the entire nucleus (in space) 218 (Figure 5A). In general, a positive correlation denotes a low-low or a high-high relationship (a 219 variable de-/increases when another variable de-/increases) while analogously a negative 220 correlation denotes a high-low relationship. The autocorrelation of NND maps (Figure 5A(i)) shows 221 a positive correlation, thus regions exist spanning 2 - 4  $\mu$ m, in which chromatin is either closely 222 packed (low-low) or widely dispersed (high-high). Likewise, blobs of similar size tend to be in 223 spatial proximity (Figure 5A(iii)). These regions are not stable over time but rearrange 224 continuously, an observation bolstered by the fact that the autocorrelation diminishes with 225 increasing time lag. The cross-correlation between NND and area (Figure 5A(ii)) shows a negative 226 correlation for short time lags, suggesting that large blobs appear with a high local density while 227 small ones are more isolated. Interestingly, the correlation becomes slightly positive for time lags 228  $\geq 20$  s, indicating that big blobs are present in regions which were sparsely populated before and 229

small blobs tend to accumulate in previously densely populated regions. This is in line with dynamic

reorganization and reshaping of chromatin domains on a global scale as observed in snapshots of
the Deep-PALM image series (Figure 1A).

The flow magnitude is positively correlated for all time lags, while the correlation displays a slight 233 increase for time lags  $\leq 20$  s (Figure 5A(vi)), which has also been observed previously (13, 25, 27). 234 The spatial autocorrelation of dynamic and structural properties of chromatin are in stark contrast. 235 While structural parameters are highly correlated at short, but not at long time scales, chromatin 236 motion is still correlated at a time scale exceeding 30 s. At very short time scales (< 100 ms), 237 238 stochastic fluctuations determine the local motion of the chromatin fiber, while coherent motion becomes apparent at longer times (27). However, there exists a strong cross-correlation between 239 structural and dynamic parameters: the cross-correlation between the NND and flow magnitude 240 shows striking negative correlation at all time lags (Figure 5A(iv)), strongly suggesting that sparsely 241 distributed blobs appear less mobile than densely packed ones. The area seems to play a negligible 242 role for short time lags, but there is a modest tendency that regions with large blobs tend to exhibit 243 increased dynamics at later time points ( $\geq 10$  s; Figure 5A(v)), likely due to the strong relationship 244 between area and NND. 245

In general, parameter pairs involving chromatin dynamics exhibit an extended spatial auto- or cross-246 correlation (up to  $\sim 6 \,\mu\text{m}$ ; the lower row of Figure 5A), compared to correlation curves including 247 solely structural parameters (up to 3 - 4 µm). Furthermore, the cross-correlation of flow magnitude 248 249 and NND does not considerably change for increasing time lag, suggesting that the coupling between those parameters is characterized by a surprisingly resilient memory, lasting for at least 250 tens of seconds (28). Concomitantly, the spatial correlation of time-averaged NND maps and maps 251 of the local diffusion constant of chromatin for the entire acquisition time enforce their negative 252 correlation at the time scale of  $\sim 1 \min$  (Supplementary Figure 8). Such resilient memory was also 253 proposed by a computational study that observed that interphase nuclei behave like concentrated 254

solutions of unentangled ring polymers (29). Our data support the view that chromatin is mostly
unentangled since entanglement would influence the anomalous exponent of genomic loci in
regions of varying chromatin density (29, 30). However, our data do not reveal a correlation
between the anomalous exponent and the time-averaged chromatin density (Supplementary Figure
8), in line with our previous results using conventional microscopy (26).

Overall, the spatial cross-correlation between chromatin structure and dynamics indicates that the NND between blobs and their mobility stand in a strong mutual, negative, relationship. This relationship, however, concerns chromatin density variations at the nanoscale, but not global spatial density variations such as in eu- or heterochromatin (*26*). These results support a model in which regions with high local chromatin density, larger blobs are more prevalent and are mobile, while small blobs are sparsely distributed and less mobile (Figure 5B). Blob density and dynamics in the long-time limit are to a surprisingly large extent influenced by preceding chromatin conformations.

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#### 268 The local chromatin density is a key regulator of instantaneous chromatin dynamics

The spatial correlations above were only evaluated pairwise, while the behavior of every blob is 269 likely determined by a multitude of factors in the complex energy landscape of chromatin (23, 27). 270 Here, we aim to take a wider range of available information into account in order to reveal the 271 principle parameters, driving the observed chromatin structure and dynamics. Employing a 272 microscopy-based approach, we have access to a total of six relevant structural, dynamic and global 273 274 parameters, which potentially shape the chromatin landscape in space and time (Figure 6A). In addition to the parameters used above, we included the confinement level as a relative measure, 275 allowing the quantification of transient confinement (Materials and Methods). We further included 276 the bare signal intensity of super-resolved images and, as the only static parameter, the distance 277 from the periphery since it was shown that dynamic and structural parameters show some 278 dependence on this parameter (Supplementary Figure 7). We then employed t-Distributed 279

Stochastic Neighbor Embedding (31) (t-SNE), a state of the art dimensionality reduction technique, 280 to map the six-dimensional chromatin 'features' (the six input parameters) into two dimensions. 281 (Figure 6A, see Supplementary Note 3). The t-SNE algorithm projects data points such that 282 neighbors in high-dimensional space likely stay neighbors in two-dimensional space (31). Visually 283 apparent grouping of points (Figure 6B) implies that grouped points exhibit great similarity with 284 respect to all input features and it is of interest to reveal which subset of the input features can 285 explain the similarity among chromatin blobs best. It is likely that points appear grouped because 286 their value of a certain input feature is considerably higher or lower than the corresponding value 287 of other data points. We hence labeled points in t-SNE maps which are smaller than the first quartile 288 point or larger than the third quartile point. Data points falling in either of the low/high partition of 289 one input feature are colored accordingly for visualization (Figure 6D; blue/red points respectively). 290 We then assigned a rank to each of the input features according to their nearest-neighbor fraction 291 (n-n fraction): Since the t-SNE algorithm conserves nearest neighbors, we described the extent of 292 grouping in t-SNE maps by the fraction of nearest neighbors which fall in either one of the 293 subpopulations of low or high points (illustrated in Supplementary Figure 9). A high nearest 294 neighbor fraction (n-n fraction; Figure 6C) therefore indicates that many points marked as low/high 295 are indeed grouped by t-SNE and are therefore similar. The ranking (from low to high n-n fraction) 296 reflects the potency of a given parameter to induce similar behavior between chromatin blobs with 297 respect to all input features. 298

The relative frequency at which each parameter ranked first provides an intuitive feeling for the most 'influential' parameters in the dataset (Figure 6E). The signal intensity plays a negligible role, suggesting that our data is free of potential artifacts related to the bare signal intensity. Furthermore, the blob area and the distance from the periphery likewise do not considerably shape chromatin blobs. In contrast, the NND between blobs was found to be the main factor inducing the observed characteristics in 67 % of all-time frames across all nuclei. The flow magnitude and confinement

305 level together rank 1<sup>st</sup> in 26 % of all cases (11 % and 17 %, respectively). These numbers suggest that the local chromatin density is a universal key regulator of instantaneous chromatin dynamics. 306 Note that no temporal dependency is included in the t-SNE analysis and thus the feature extraction 307 concerns only short-term ( $\leq$  360 ms) relationships. The characteristics of roughly one-fourth of all 308 blobs at each time point are mainly determined by similar dynamical features. Mapping chromatin 309 blobs as marked in Figure 6C, D back to their respective positions inside the nucleus (Figure 6F) 310 shows that blobs with low/high flow magnitude or confinement level markedly also grouped in 311 physical space, which is highly reminiscent of coherent motion of chromatin (25). In contrast, blobs 312 313 with extraordinary low or high NND were found interspersed throughout the nucleus, in line with spatial correlation analysis between structural and dynamic features (Figure 5). Our results point 314 towards a large influence of the local chromatin density on the dynamics of chromatin at the scale 315 of a few hundred nanometers and within a few hundred milliseconds. At longer time and length 316 scales, however, previous results suggest that this relationship is lost (26) 317

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#### 319 **Discussion**

With Deep-PALM we present temporally resolved super-resolution images of chromatin in living 320 cells. Our technique identified chromatin nanodomains, called "blobs", which mostly have an 321 elongated shape, consistent with the curvilinear arrangement of chromatin as revealed by Structured 322 Illumination Microscopy (13) with typical axes length of 45 - 90 nm. A previous study reported 323  $\sim$ 30 nm wide 'clutches of nucleosomes' in fixed mammalian cells using STORM nanoscopy (10), 324 while the larger value obtained using Deep-PALM may be attributed to the motion blurring effect 325 in live-cell imaging. However, histone acetylation and methylation marks were shown to form 326 nanodomains of diameter 60 - 140 nm, respectively (14), which includes the computed dimensions 327 for histone H2B using Deep-PALM. 328

Using an established chromosome model, chromatin blobs were shown to consist of continuous genomic regions with an average length of 75 kb. While assembling and disassembling dynamically within less than one second. Monomers within blobs display a distinct TAD-like association pattern in the long-time limit, suggesting that the identified blobs represent sub-TADs. Transient formation is consistent with recent findings that TADs are not stable structural elements, but exhibit extensive heterogeneity and dynamics (7, 9).

We found that structural and dynamic parameters exhibit extended spatial and temporal (cross-) 335 correlations. Structural parameters such as the local chromatin density (expressed as the NND 336 between blobs) and area lose their correlation after  $3 - 4 \mu m$  and roughly 40 s in the spatial and 337 temporal dimension, respectively. In contrast, chromatin mobility correlations extend over  $\sim 6 \,\mu m$ 338 and persist during the whole acquisition period ( $\geq 40$  s). Extensive spatio-temporal correlation of 339 chromatin dynamics have been presented previously, both experimentally (25) and in simulations 340 (27), but was not linked to the spatio-temporal behavior of the underlying chromatin structure until 341 now. We found that the chromatin dynamics are closely linked to the instantaneous, but also to past 342 local structural characterization of chromatin. In other words, the instantaneous local chromatin 343 density influences chromatin dynamics in the future and vice versa. Based on these findings, we 344 suggest that chromatin dynamics exhibit an extraordinary long memory. This strong temporal 345 relationship might be established by the fact that stress propagation is affected by the folded 346 chromosome organization (32). Fiber displacements cause structural reconfiguration, ultimately 347 leading to a local amplification of chromatin motion in local high-density environments. This 348 observation is also supported by the fact that increased nucleosome mobility grants chromatin 349 accessibility even within regions of high nucleosome density (33). 350

Given the persistence at which correlations of chromatin structure and, foremost, dynamics occur in a spatio-temporal manner, we speculate that the interplay of chromatin structure and dynamics could involve a functional relationship (*34*): transcriptional activity is closely linked to chromatin

accessibility and the epigenomic state (35). Because chromatin structure and dynamics are related, 354 dynamics could also correlate with transcriptional activity (26, 36). However, it is currently 355 unknown if the structure-dynamics relationship revealed here is strictly mutual or if it may be 356 causal. Simulations hint that chromatin dynamics follows from structure (27, 28), this question will 357 be exciting to answer experimentally and in the light of active chromatin remodelers in order to 358 elucidate a potential functional relationship to transcription. Chromatin regions which are switched 359 from inactive to actively transcribing, for instance, undergo a structural reorganization 360 accompanied by epigenetic modifications (37, 38). The mechanisms driving recruitment of 361 enzymes inducing histone modifications such as histone acetyltransferases, deacetylases or 362 methyltransferases is largely unknown, but often involves the association to proteins (39). Their 363 accessibility to the chromatin fiber is inter alia determined by local dynamics (33). Such a structure-364 dynamics feedback loop would constitute a quick and flexible way to transiently alter gene 365 expression patterns upon reaction to external stimuli or to co-regulate distant genes (40, 41). Future 366 work will study how structure-dynamics correlations differ in regions of different transcriptional 367 activity and/or epigenomic states. Furthermore, to probe the interactions between key 368 transcriptional machines such as RNA polymerases with the local chromatin structure and to record 369 their (possibly collective) dynamics could shed light into the target search and binding mechanisms 370 of RNA polymerases with respect to the local chromatin structure. Deep-PALM in combination 371 with Optical Flow paves the way to answer these questions by enabling the analysis of time-372 373 resolved super-resolution images of chromatin in living cells.

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#### 379 Materials and Methods

#### 380 Cell Culture.

Human osteosarcoma U2OS expressing H2B-PATagRFP cells were a gift from Sébastien Huet 381 (CNRS, UMR 6290, Institut Génétique et Développement de Rennes, Rennes, France); the histone 382 H2B was cloned as described previously (42). U2OS cells were cultured in DMEM (with 4.5 g/l 383 glucose) supplemented with 10% fetal bovine serum (FBS), 2 mM glutamine, 100 ug/ml penicillin, 384 and 100 U/ml streptomycin in 5% CO<sub>2</sub> at 37°C. Cells were plated 24 hours before imaging on 35 385 mm Petri dishes with a #1.5 coverslip like bottom (ibidi, Biovalley) with a density of  $2 \times 10^5$ 386 cells/dish. Just before imaging, the growth medium was replaced by Leibovitz's L-15 medium (Life 387 Technologies) supplemented with 20% FBS, 2 mM glutamine, 100 µg/ml penicillin, and 100 U/ml 388 streptomycin. 389

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#### 391 PALM Imaging.

Imaging of H2B-PAtagRFP in live U2OS cells was carried out on a fully automated Nikon TI-E/B 392 PALM (Nikon Instruments) microscope. The microscope is equipped with a full incubator 393 enclosure with gas regulation to maintain a temperature of ~37°C for normal cell growth during 394 live-cell imaging. A laser line of 561 nm (~50-60 W/cm<sup>2</sup> at the sample) was applied for PATagRFP 395 excitation and the 405 nm laser used for photo-activation (~2-2.5 W/cm<sup>2</sup> at the sample). Laser beam 396 powers were controlled by acoustic optic-modulators (AA Opto-Electronics). Both wavelengths 397 were united into an oil immersion 1.49 NA TIRF objective (100x; Nikon). An oblique illumination 398 was applied to acquire image series with high signal to noise ratio. The fluorescence emission signal 399 was collected by using the same objective and spectrally filtered by a Quad-Band beam splitter 400 (ZT405/488/561/647rpc-UF2, Chroma Technology) with Ouad-Band emission filter 401 (ZET405/488/561/647m-TRF, Chroma). The signal was recorded on an EMCCD camera (Andor 402 iXon X3 DU-897, Andor Technologies) with a pixel size of 108 nm. For axial correction, Perfect 403

404 Focus System was applied to correct for defocusing. NIS-Elements software was used for acquiring

405 the images at 30 ms per frame.

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**Deep-PALM analysis.** The convolutional neural network (CNN) was trained using simulated data following Nehme *et al.* (*19*) for three labeling densities (4, 6 and 9 emitters per  $\mu$ m<sup>2</sup> per frame). Raw imaging data were checked for drift as previously described (*25*). The detected drift in raw images is in the range < 10 nm and therefore negligible. The accuracy of the trained net was evaluated by constructing ground truth images from the simulated emitter positions. The Structural Similarity Index is computed to assess the similarity between reconstructed and ground truth images (*43*):

$$SSIM = \sum_{x,y} \frac{(2\mu_x\mu_x + C_1)(2\sigma_{xy} + C_2)}{(\mu_x^2 + \mu_y^2 + C_1)(\sigma_x^2 + \sigma_y^2 + C_2)},$$
(1)

where x and y are windows of the predicted and ground truth images, respectively,  $\mu$  and  $\sigma$  denote their local means and standard deviation, respectively and  $\sigma_{xy}$  their cross-variance.  $C_1 = (0.01L)^2$ and  $C_2 = (0.03L)^2$  are regularization constants, where *L* is the dynamic range of the input images. The second quantity to assess CNN accuracy is the Root Mean Square Error between the ground truth G and reconstructed image R:

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$$RMSE = \sqrt{\frac{1}{N} \sum_{N} (R - G)^2},$$
(2)

420

Where *N* is the number of pixels in the images. After training, experimental data were supplied to the trained network and predictions of single Deep-PALM images were summed to a final superresolved image. An up-sampling factor of 8 was used, resulting in an effective pixel size of 108 nm/8 = 13.5 nm. The image quality assessment in order to determine the optimal number of predictions to be summed, we use a blind/referenceless image spatial quality evaluator (BRISQUE)

426 (44). For visualization, super-resolved images were convolved with a Gaussian kernel ( $\sigma = 1$ 

427 pixel) and represented using a false RGB colormap.

428

Fourier Ring Correlation analysis. Fourier Ring Correlation (FRC) is an unbiased method to 429 estimate the spatial resolution in microscopy images. We follow an approach similar to the one 430 described in Nieuwenhuizen et al. (45). For localization-based super-resolution techniques, the set 431 of localizations is divided into two statistically independent subsets and two images from these 432 subsets are generated. The FRC is computed as the statistical correlation of the Fourier transforms 433 of both sub-images over the perimeter of circles of constant frequency in the frequency domain. 434 Deep-PALM, however, does not result in a list of localizations, but in predicted images directly. 435 The set of 12 predictions from Deep-PALM were thus split into two statistically independent 436 subsets and the method described in Nieuwenhuizen et al. (45) was applied. 437

438

Chromatin blob identification. The super-resolved images displayed isolated regions of 439 accumulated emitter density. To quantitatively assess the structural information implied by these 440 accumulation of emitters in the focal plane, we developed a segmentation scheme which aims to 441 identify individual blobs (Supplementary Figure 2). A marker-assisted watershed segmentation was 442 adapted in order to accurately determine blob boundaries. For this purpose, we use the raw 443 predictions from the deep convolutional neural network without convolution (Supplementary 444 Figure 2A). The foreground in this image is marked by regional maxima and pixels with very high 445 density (i.e. those with  $I > 0.99 I_{max}$ , Supplementary Figure 2B). Since blobs are characterized by 446 447 surrounding pixels of considerably less density, the Euclidian distance transform is computed on the binary foreground markers. Background pixels (i.e. those pixels not belonging to any blobs) are 448 expected to lie far away from any blob center and thus, a good estimate for background markers are 449 those pixels being furthest from any foreground pixel. We hence compute the watershed transform 450

on the distance transform of foreground markers and the resulting watershed lines depict 451 background pixels (Supplementary Figure 2C). Equipped with fore- and background markers 452 (Supplementary Figure 2D), we apply a marker-controlled watershed transform on the gradient of 453 the input image (Supplementary Figure 2E). The marker-controlled watershed imposes minima on 454 marker pixels preventing the formation of watershed lines across marker pixels. Therefore, the 455 marker-controlled watershed accurately detects boundaries as well as blobs which might not have 456 been previously marked as foreground (Supplementary Figure 2F). Finally, spurious blobs whose 457 median- or mean intensity is below 10% of the maximum intensity are discarded and each blob is 458 assigned a unique label for further correspondence (Supplementary Figure 2G). The area and 459 centroid position are computed for each identified blob for further analysis. This automated 460 segmentation scheme performs considerably better than other state-of-the-art algorithms for image 461 segmentation due to the reliable identification of fore- and background markers accompanied by 462 the watershed transform (Supplementary Note 1). 463

464

465 **Chromatin blob properties.** Centroid position, area, and eccentricity were computed. The 466 eccentricity is computed by describing the blobs as an ellipse:

$$E = \sqrt{1 - a^2/b^2} \tag{3}$$

467 where *a* and *b* are the short and long axes of the ellipse, respectively.

468

Chromatin blob identification from a computational chromatin model. We chose to employ a computational chromatin model, recently introduced by Qi *et al.* (*23*), in order to elucidate the origin of experimentally determined chromatin blobs. Each bead of the model covers a sequence length of 5 kb and is assigned one of 15 chromatin states to distinguish promoters, enhancers, quiescent chromatin, etc. Starting from the simulated polymer configurations, we consider monomers within a 200 nm thick slab through the center of the simulated chromosome. In order to

generate super-resolved images as those from Deep-PALM analysis, fluorescence intensity is 475 ascribed to each monomer. Monomer positions are subsequently discretized on a grid with 13.5 nm 476 spacing and convolved with a narrow point-spread function, which results in images closely 477 resembling experimental Deep-PALM images of chromatin. Chromatin blobs were then be 478 identified and characterized as on experimental data (Figure 2A, B). Mapping back the association 479 of each bead to a blob (if any) allows us to analyze principles of blob formation and maintenance 480 using the distance and the association strength between each pair of monomers, averaged over all 481 20,000 simulated polymer configurations. 482

483

**Radial distribution function.** The radial distribution function g(r) (RDF, also pair correlation function) is calculated (in two dimensions) by counting the number of blobs in an annulus of radius *r* and thickness *dr*. The result is normalized by the bulk density  $\rho = n/A$ , with the total number of blobs *n* and *A* the area of the nucleus, and the area of the annulus,  $2\pi r \, dr$ :

$$dn(r) = \rho \cdot g(r) \cdot 2\pi r \, dr \tag{4}$$

Quantification of chromatin dynamics. Super-resolved images of chromatin showed spatially 488 distributed blobs of varying size, but the resolved structure is too dense for state-of-the-art single 489 particle tracking methods to track. Furthermore, are highly dynamic structures, assembling and 490 dissembling within 1 - 2 super-resolved frames (Figure 3D), which makes a Single Particle 491 Tracking approach unsuitable. Instead, we used a method for dynamics reconstruction of bulk 492 macromolecules with dense labeling, Optical Flow. Optical Flow builds upon the computation of 493 flow fields between two successive frames of an image series. The integration of these flow fields 494 from super-resolution images results in trajectories displaying the local motion of bulk chromatin 495 with temporal and high spatial resolution. Further, the trajectories are classified into various 496 diffusion models and parameters describing the underlying motion are computed (26). Here, we use 497 the effective diffusion coefficient D (in units of  $m^2/s^{\alpha}$ ), which reflects the magnitude of 498 displacements between successive frames (the velocity of particles or monomers in the continuous 499

limit) and the anomalous exponent  $\alpha$  (26). The anomalous exponent reflects if the diffusion is free 500  $(\alpha = 1, e.g.$  for non-interacting particles in solution), directed ( $\alpha > 1$ , e.g. as the result from active 501 processes) or hindered ( $\alpha < 1$ , e.g. due to obstacles or an effective back-driving force). 502 Furthermore, we compute the length of constraint  $L_c$  which is defined as the standard deviation of 503 the trajectory positions with respect to its time-averaged position. Denoting  $R(t; R_0)$  the trajectory 504 at time t originating from  $R_0$ , the expression reads (46)  $L_c(R_0) = var(R(t; R_0))^{1/2}$ , where var 505 denotes the variance. The length of constraint is a measure of the length scale explored of the 506 monomer during the observation period. A complementary measure is the confinement level (47), 507 which computes the inverse of the variance of displacements within a sliding window of length  $\omega$ : 508  $C \propto \omega / \operatorname{var}(\mathbf{R}(t; \mathbf{R}_0))$ , where the sliding window length is set to 4 frames (1.44 s). Larger values 509 of *C* denote a more confined state than small ones. 510

511

Spatial correlation for temporally varying parameters. The nearest-neighbor distance and the 512 513 area, as well as the flow magnitude, were calculated and assigned to the blobs' centroid position. In order to calculate the spatial correlation between parameters, the parameters were interpolated 514 from the scattered centroid positions onto a regular grid spanning the entire nucleus. Because not 515 every pixel in the original super-resolved images is assigned a parameter value, we chose an 516 effective grid spacing of 5 pixels (67.5 nm) for the interpolated parameter maps. After interpolation, 517 the spatial correlation was computed between parameter pairs: Let  $\mathbf{r} = (x, y)^T$  denote a position 518 on a regular two-dimensional grid and  $f(\mathbf{r},t)$  and  $g(\mathbf{r},t)$  two scalar fields with mean zero and 519 variance one, at time t on that grid. The time series of parameter fields consist of N time points. 520 The spatial cross-correlation between the fields f and g, which lie a lag time  $\tau$  apart, is then 521 522 calculated as

$$C(\boldsymbol{\rho},\tau) = \frac{1}{N} \sum_{t} \frac{\sum_{x,y} f(\boldsymbol{r},t) g(\boldsymbol{r}+\boldsymbol{\rho},t+\tau)}{\sum_{x,y} f(\boldsymbol{r},t) \cdot g(\boldsymbol{r},t+\tau)}$$
(5)

where the space lag  $\rho$  is a two-dimensional vector  $\rho = (\Delta x, \Delta y)^T$ . The sums in the numerator and denominator are taken over the spatial dimensions, the first sum is taken over time. The average is thus taken over all time points which are compliant with time lag  $\tau$ . Subsequently, the radial average in space is taken over the correlation, thus effectively calculating the spatial correlation  $C(\rho, \tau)$ over the space lag  $\rho = \sqrt{\Delta x^2 + \Delta y^2}$ . If f = g, the spatial autocorrelation is computed.

528

**Spatial correlation for static parameters.** We denote global parameters as those, which reflect the structural and dynamic behavior of chromatin spatially resolved, but in a time-averaged manner. Examples involve the diffusion constant, the anomalous exponent, the length of constraint, but also time-averaged nearest-neighbor distance maps, etc. (Supplementary Figure 8). Those parameters are useful to determine time-universal characteristics. The spatial correlation between those parameters is equivalent to the expression given for temporally varying parameters when the temporal dimension is omitted, effectively resulting in a correlation curve  $C(\rho)$ .

536

t-Distributed Stochastic Neighbor Embedding (t-SNE). The distance from the periphery, 537 intensity, their nearest-neighbor distance, area, flow magnitude and confinement level of each 538 identified blob form the six-dimensional input feature space for t-SNE analysis. The parameters for 539 540 each blob (n = 3.260,232, divided into subsets of approximately 10,000) were z-transformed prior to the t-SNE analysis. The t-SNE analysis was performed using MATLAB and the Statistics and 541 Machine Learning Toolbox (Release 2017b, The MathWorks, Inc., Natick, Massachusetts, United 542 States) with the Barnes-Hut approximation. The algorithm was tested using different distance 543 metrics and perplexity values and showed robust results within the examined ranges 544 (Supplementary Note 3, Supplementary Figure 10). 545

#### 547 H2: Supplementary Materials

- 548
- 549 Note S1. Performance of the segmentation scheme employed for this study in comparison to other
- 550 state-of-the-art algorithms for general purpose and comparison of blob segmentation on
- 551 experimental images and on images containing randomly distributed emitters.
- 552 Note S2. Suitability of Optical Flow for super-resolution images of chromatin.
- 553 Note S3. t-SNE and its robustness with respect to distance metrics and perplexity values.
- 554 Figure S1. CNN training and time-resolution determination.
- 555 Figure S2. Chromatin blob identification pipeline.
- 556 Figure S3. Performance of segmentation algorithms on super-resolved images of chromatin *in vivo*.
- 557 Figure S4. Segmentation on images of randomly distributed emitters.
- 558 Figure S5. Chromatin area fraction.
- 559 Figure S6. Performance of Optical Flow on conventional and super-resolved images.
- 560 Figure S7. Structural and dynamic parameters are dependent on the proximity to the nuclear
- 561 periphery.
- 562 Figure S8. Global spatial correlation of structural and dynamic parameters.
- 563 Figure S9. Clustering illustration of points within a subset based on nearest-neighbors in t-SNE
- 564 maps.
- 565 Figure S10. t-SNE for different distance metrics and perplexity values.
- 566 Movie S1. Time series of super-resolved chromatin structure and dynamics.
- 567

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| 722 | Author contributions: H.A.S. designed and supervised the project; R.B. designed the data analysis   |
|-----|---|
| 723 | and wrote the code, H.A.S carried out experimental work; R.B. carried out the data analysis; H.A.S. |
| 724 | and R. B interpreted results: H.A.S, R.B. and K.B. wrote the manuscript.                            |
| 725 |   |
| 726 | Competing interests: The authors declare no competing financial interest.                           |
| 727 |   |
|     |   |

728 Data and materials availability: All data needed to evaluate the conclusions in the paper are 729 present in the paper and/or the Supplementary Materials. Additional data related to this paper may 730 be requested from the authors.

731

#### 732 Figures and Tables

Figure 1: Temporally resolved super-resolution images of chromatin in U2OS nuclei. A) 733 Widefield images of U2OS nuclei expressing H2B-PATagRFP are input to a trained convolutional 734 neural network (CNN) and predictions from multiple input frames are summed to construct a super-735 resolved image of chromatin *in vivo*. **B)** The resolution tradeoff between the prolonged acquisition 736 of emitter localizations (green line) and motion blur due to diffusion of the underlying diffusion 737 processes (purple line). For our experimental data, the localization density per second is  $\rho =$ 738  $(2.4 \pm 0.1) \mu m^{-2} s^{-1}$ , the diffusion constant is  $D = (3.4 \pm 0.8) \cdot 10^{-3} \mu m^2 s^{-1}$ (see 739 740 Supplementary Figure 8B) and the acquisition time per frame is  $\tau = 30 \text{ ms}$ . The spatial resolution assumes a minimum ( $69 \pm 5$  nm) at a time resolution of 360 ms. C) Super-resolution images of a 741 single nucleus at time intervals of about 10 seconds. Scale bar is 2 µm. D) Magnification of 742 segregated accumulations of H2B within a chromatin-rich region. Scale bar is 200 nm. E) 743 Magnification of a stable, but dynamic structure (arrows) over three consecutive images. Scale bar 744 is 500 nm. F) Fourier Ring Correlation (FRC) for super-resolved images resulting in a spatial 745 resolution of  $63 \pm 2$  nm. 746

747

| 748 | Figure 2: Chromatin blob identification and characterization of imaged and modeled  |
|-----|---|
| 749 | chromatin. A) Super-resolved images show blobs of chromatin (left). These blobs are segmented                             |
| 750 | (Materials and Methods, Supplementary Note 1) and individually labeled by random color (right).                           |
| 751 | Magnifications of the boxed regions are shown. Scale bars: whole nucleus 2 $\mu$ m, magnifications                        |
| 752 | 200 nm. B) Generation of super-resolution images and blob identification and characterization for                         |
| 753 | a 25 Mbp segment of chromosome 1 from GM12878 cells as simulated in Qi et al. (23). Beads (5                              |
| 754 | kb genomic length) of a simulated polymer configuration within a 200 nm thick slab are projected                          |
| 755 | to the imaging plane, resembling experimental super-resolved images of live chromatin. Blobs are                          |
| 756 | identified as on experimental data. C) From the centroid positions, the nearest-neighbor distance                         |
| 757 | (NND) distributions are computed for up to 40 nearest neighbors (blue to red). The envelope of the                        |
| 758 | k-NND distributions (black line) shows peaks at approximately 95 nm, 235 nm, 335 nm and 450                               |
| 759 | nm (red dots). D) k-NND distributions as in B) for simulated data. E) Area distribution of                                |
| 760 | experimental and simulated blobs. The distribution is in both cases well described by a lognormal                         |
| 761 | distribution with parameters $(3.3 \pm 2.8) \cdot 10^{-3} \mu\text{m}^2$ for experimental blobs and $(3.1 \pm 3.2) \cdot$ |
| 762 | $10^{-3} \mu\text{m}^2$ for simulated blobs (mean $\pm$ standard deviation). F) Eccentricity distribution for             |
| 763 | experimental and simulated chromatin blobs. Selected eccentricity values are illustrated by ellipses                      |
| 764 | with the corresponding eccentricity. Eccentricity values range from 0, describing a circle, to 1,                         |
| 765 | describing a line. Prominent peaks arise due to the discretization of chromatin blobs in pixels.                          |

**Figure 3: Chromatin blobs on modeled chromosomes consist of continuous loci along the genome and exhibit a TAD-like time-averaged conformation. A)** Gap length between beads belonging to the same blob. An exemplary blob with small gap length is shown. The blob is mostly made of consecutive beads being in close spatial proximity. **B)** A representative polymer configuration is colored according to chromatin states (red: active, green: inactive, blue: repressive). **C)** The cumulative distribution function of clusters within active, inactive and

772 repressive chromatin. Inset: Mean area of clusters within the three types of chromatin. The distributions are all significantly different from each other as determined by a two-sample 773 Kolmogorov-Smirnov test ( $p < 10^{-50}$ ). **D**) Distribution of the continuous residence time of any 774 monomer within a cluster (0.5 + 0.3 s; mean + standard deviation). Inset: Continuous residence 775 time of any monomer within a slab of 200 nm thickness  $(1.5 \pm 1.6 \text{ s}; \text{mean} \pm \text{standard deviation})$ . 776 E) The blob association strength between any two beads is measured as the frequency at which any 777 two beads are found in one blob. The association map is averaged over all simulated configurations 778 (upper triangular matrix; from simulations) and experimental Hi-C counts are shown for the same 779 chromosome segment (lower triangular matrix; from Rao et al. (48)). The association and Hi-C 780 maps are strongly correlated (Pearson's correlation coefficient PCC = 0.76) F) Close-up views 781 around the diagonal of Hi-C-like matrices. The association strength is shown together with the 782 inverse distance between beads (upper panel; PCC = 0.85) and with experimental Hi-C counts 783 (lower panel; as in E)). 784

785

Figure 4: Super-resolution chromatin dynamics. A) A time series of super-resolution images 786 (left panel) is subject to Optical Flow (right panel). B) (i-ii) Two subsequent images of chromatin. 787 Magnifications show prominent mobile blobs on the nuclear periphery in both images (colored 788 arrows). (iii) The flow field and corresponding magnification are shown on top of a fused image of 789 both super-resolved images in (i) and (ii) (green and purple respectively, co-localization are white). 790 The flow field is colored according to the direction of vectors (see color wheel) C) As for B) in the 791 nuclear interior. Scale bars: whole nucleus 2 µm, magnifications 500 nm. Flow vectors are not 792 drawn to scale and down-sampled 8-fold for clarity. D) Blobs of a representative nucleus (see 793 794 Supplementary Movie 1) are labeled by their NND (left), area (middle) and flow magnitude (right). Colors denote the corresponding parameter magnitude. 795

#### 797 Figure 5: Spatio-temporal correlations between structural and dynamic parameters. A) The

spatial auto- and cross-correlation between parameters were computed for different time lags. The graphs depict the correlation over space lag for each parameter pair and different colors denote the time lag (increasing from blue to red). **B**) Illustration of the instantaneous relationship between local chromatin density and dynamics. The blob density is shown in blue; the magnitude of chromatin dynamics is shown by red arrows. The consistent negative correlation between NND and flow magnitude is expressed by increased dynamics in regions of high local blob density.

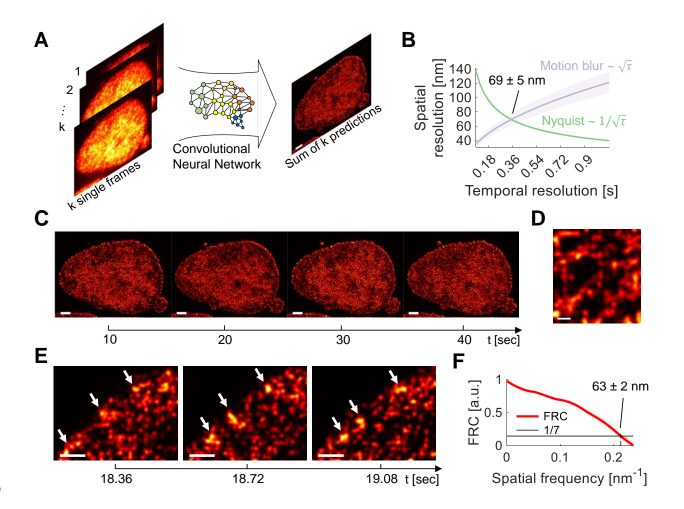
Figure 6: Chromatin feature extraction. A) The six-dimensional parameter space is input to the 804 t-SNE algorithm and projected to two dimensions. B) The 2D embedding of an exemplary data set 805 is shown and colored according to the magnitude of each input feature (blue to red, the parameter 806 average is shown in beige) C) Points below the first (blue) and above the third (red) quartile points 807 of the corresponding parameter are marked and the parameters are ranked according to the fraction 808 of nearest neighbors which fall in one of the marked regions. D) Data points marked below the first 809 or above the third quartile points are labeled according to the feature in which they were marked. 810 Priority is given to the feature with the higher nearest-neighbor fraction if necessary. E) t-SNE 811 analysis is carried out for each nucleus over the whole time series and it is counted how often a 812 parameter ranked 1<sup>st</sup>. The results are visualized as a pie chart. The NND predominantly ranks 1<sup>st</sup> in 813 about 2/3 of all cases. F) Marked points in C-D) are mapped back onto the corresponding nuclei 814 and the cumulative distribution function (CDF) over space is shown. 815

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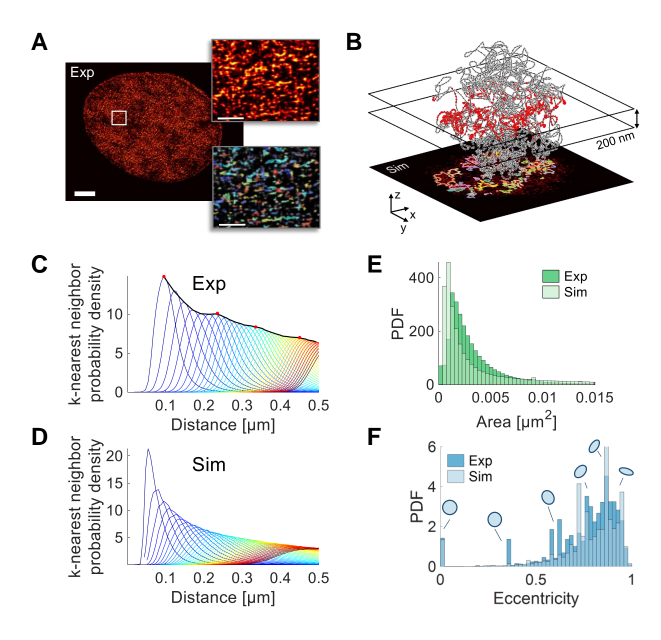
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818

## 819 **Figure 1**

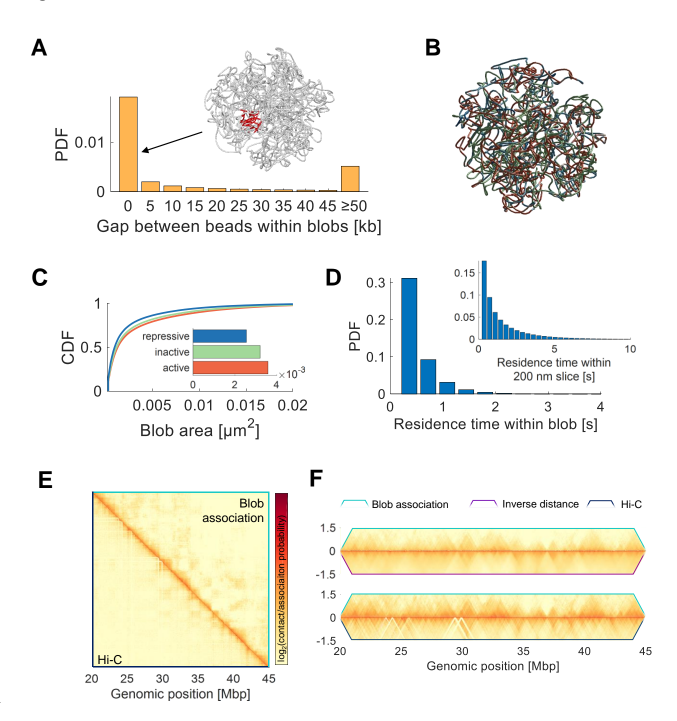


## 821 Figure 2



822

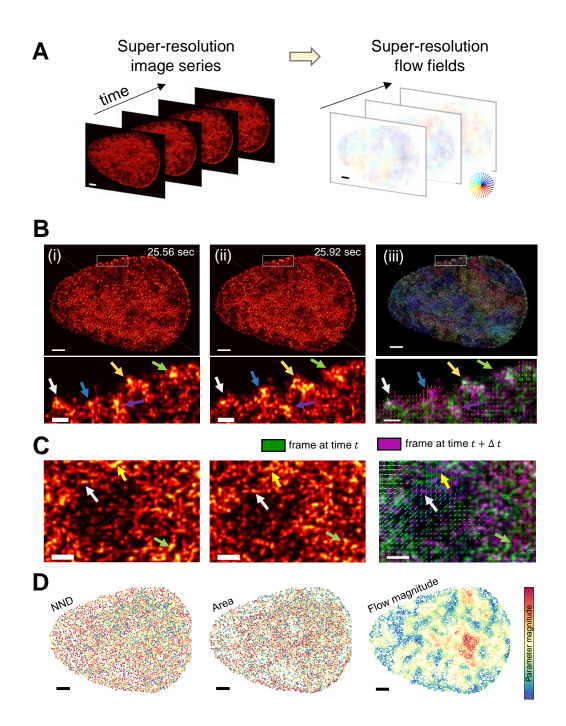
## 823 Figure 3



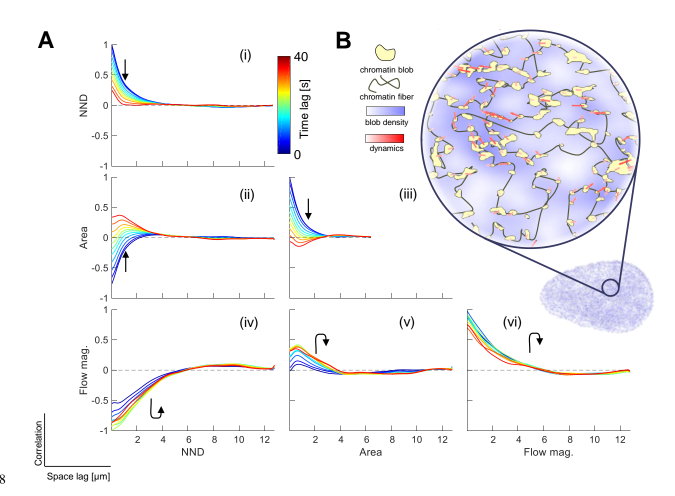
824

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### 825 Figure 4

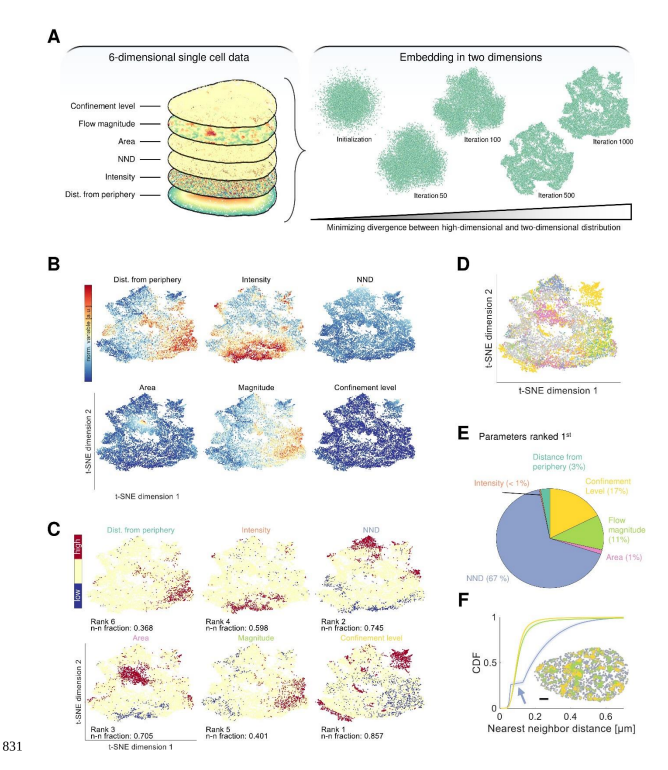


### 827 Figure 5



828

#### Figure 6 830



# SUPPLEMENTARY MATERIAL

# <sup>834</sup> Coupling chromatin structure and dynamics by live

# super-resolution imaging

| 836               | R. Barth <sup>1, 2,</sup> K. Bystricky <sup>1</sup> and H. A. Shaban <sup>,1, 3, †, *</sup>  |
|-------------------|--|
| 837<br>838        | 1: Laboratoire de Biologie Moléculaire Eucaryote (LBME), Centre de Biologie Intégrative (CBI),<br>CNRS; University of Toulouse, UPS; 31062 Toulouse; France  |
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|                   |  |

846 <u>hshaban@fas.harvard.edu</u>

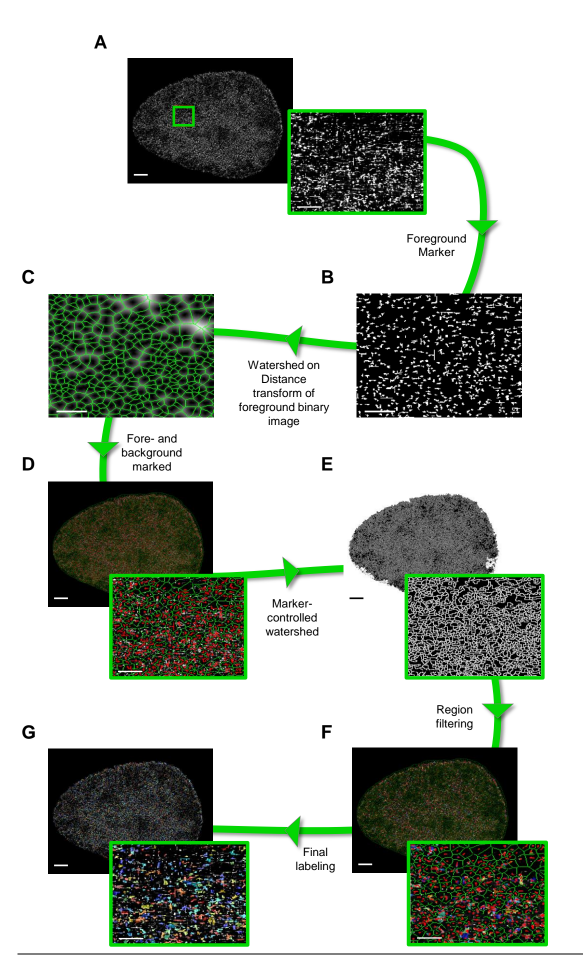
| Α |  | <i>.</i> .   |                           | , C                        | С        | NN 4                      | CNN 6   |
|---|--|--|---------------------------|----------------------------|----------|---------------------------|---|
|   | Experimen  |  | 4 emitters /<br>Predicted | µm²<br><sub>*</sub> ×      | 180 ms   |                           |   |
|   |  |  | ***<br>                   |                            | 240 ms   |                           |   |
|   | 6 emitters   | / um <sup>2</sup>  | 9 emitters /              | nulated<br>um <sup>2</sup> | 300 ms   |                           |   |
|   | Predicted  | ×  | Predicted                 | ×.                         | 360 ms   |                           |   |
|   |  |  |                           |                            | 420 ms   |                           |   |
| В | huller   | mulated  |                           | nulated.                   | 480 ms   |                           |   |
|   | y Index<br>96.0  | <u>n. s.</u>   | ***                       |                            | 540 ms   |                           |   |
|   | Structural<br>6.0<br>58.0<br>58.0<br>58.0  |  |                           |                            | 600 ms   |                           |   |
|   |  | сли <sup>н А</sup> сли <sup>к</sup><br>×10 <sup>-3</sup> | CHN9                      |                            | 660 ms   |                           |   |
|   | Root Mean<br>Square Error [A.U.]<br>50<br>1 51<br>52<br>53<br>53<br>54<br>54<br>54<br>54<br>55<br>55<br>55<br>55<br>55<br>55<br>55<br>55<br>55 | ×10 <sup></sup>  | ***                       | D<br>III                   |          |                           |   |
|   | Root<br>Square E   |  |                           | BRISQUE<br>score           | 50<br>45 |                           |   |
|   | 07   | CMMA CM  | NO CMNO                   |                            | 0.       | ې مې<br>Time<br>esolutior | , <sup>()</sup> |
|   |  |  |                           |                            |          | Solution                  | . []  |

#### Supplementary Figure 1: CNN training and time-resolution determination. A) Experimental 848 and simulated widefield images with varying labeling density. Predictions by the trained CNNs are 849 shown as false-color images and the ground truth emitter positions are overlaid as green crosses. 850 Scale bar for experimental widefield image is 2 $\mu$ m, simulated images are the same size. Scale bar 851 on predictions are 200 nm. B) The accuracy of the trained CNNs was evaluated using the Structural 852 Similarity Index and the Root Mean Square Error (Materials and Methods). The CNN trained with 853 9 emitters per $\mu m^2$ (CNN 9) performs significantly worse than the other two CNNs. Statistical 854 significance assessed by a two-sample t-test (\*\*\* p < 0.001). C) Predictions from single acquired 855

images are summed over different times from 360 ms to 1020 ms. A subset of reconstructions up

to a time resolution of 660 ms are shown for CNN 4 and CNN 6. Scale bar is  $2 \mu m$ . **D**) The structural image quality as quantified by BRISQUE (44) for CNN 6 at varying temporal resolution. The optimal (maximum) value is found at a time resolution of 360 ms.

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Supplementary Figure 2: Chromatin blob identification pipeline. A) A grayscale image as 863 output from the Deep-PALM algorithm is to be segmented. Magnified views correspond to the red 864 rectangle. B) Foreground pixels are marked by finding the regional maxima. Additionally, very 865 bright pixels (i.e. those with  $I > 0.99 \cdot I_{max}$ ) are marked as foreground. Only the magnification is 866 shown. C) The watershed algorithm (without markers) is applied to the distance transform of the 867 foreground mask. The resulting watershed lines represent lines in between pixels marked as 868 foreground and therefore represent pixels of (local) low intensity, i.e. background markers. The 869 870 background mask is corrected for pixels which nevertheless exert high intensity (i.e. those with I > $0.8 \cdot I_{max}$ ). Only the magnification is shown. D) Input image with the foreground (red) and 871 background (green) markers superimposed. E) The watershed algorithm is applied to the gradient 872 of the input image with minima imposed on pixels belonging to fore- or background. F) The regions 873 defined by watershed lines (the basins) are post-processed in order to keep only those whose mean 874 and median intensity exceed a threshold of  $0.1 \cdot I_{max}$ . G) Resulting segmentation of the input 875 image. Scale bars correspond to 2 µm and 0.5 µm in the full and magnified regions respectively. 876

# 878 SUPPLEMENTARY NOTE 1

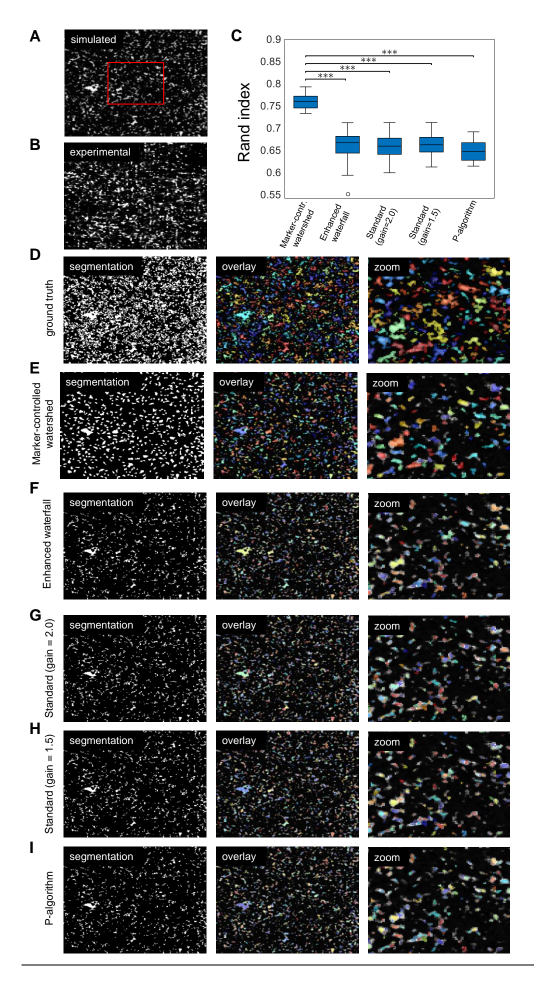
#### 879 Segmentation performance

The marker-assisted watershed segmentation pipeline developed to segment chromatin blobs in 880 super-resolved images is tested against other state-of-the-art segmentation algorithms in order to 881 validate its performance. We evaluated our approach against three fully automated improvements 882 of the widely spread watershed algorithm: the enhanced waterfall algorithm, the 'standard' 883 algorithm, and the P-algorithm (49). To this end, we simulated images which closely resemble 884 experimental super-resolved images of histone-labeled chromatin. We generated random shapes by 885 thresholding randomly generated 1/f noise and gave intensity to the shapes according to 886 experimental images. An exemplary simulated and experimental image is shown in Supplementary 887 Figure 3A, B. The segmentation result of an exemplary simulated image is shown in Supplementary 888 Figure 3D, I. We used the Rand index as a measure of the similarity of two segmentation algorithms. 889 In particular, the Rand index is computed between the ground truth and the segmentation result of 890 each algorithm as an estimate of the likelihood of a correctly classified element. An image 891 segmentation problem can be formulated as a partitioning of the image pixels into several subsets. 892 Let P denote the set of n pixels to be segmented,  $G = \{G_1, ..., G_n\}$  denotes the ground truth 893 segmentation of n pixels into g subsets and  $R = \{R_1, ..., R_r\}$  denotes the resulting segmentation 894 into r subsets from an algorithm to test. The number of agreements between G and R consists of (i) 895 the true positives (TP), i.e. the pairs of elements in P that are in the same subset in G and R and (ii) 896 the true negatives (TN), i.e. the pairs of elements in P that are not in the same subset in G and not 897 in R. Likewise, the number of disagreements consists of (iii) the false negatives (FN), i.e. elements 898 899 in P that are in the same subset in G but not in R and (iv) the false positives (FP), i.e. elements in P that are not in the same subset in G but that are in R. Taken together, the Rand index expresses 900 901 the number of agreements over the total number of pairs, i.e. agreements and disagreements (50):

Rand Index = 
$$\frac{TP + \bar{T}N}{TP + TN + FN + FP} = \frac{\bar{T}P + TN}{\binom{n}{2}}$$
 (6)

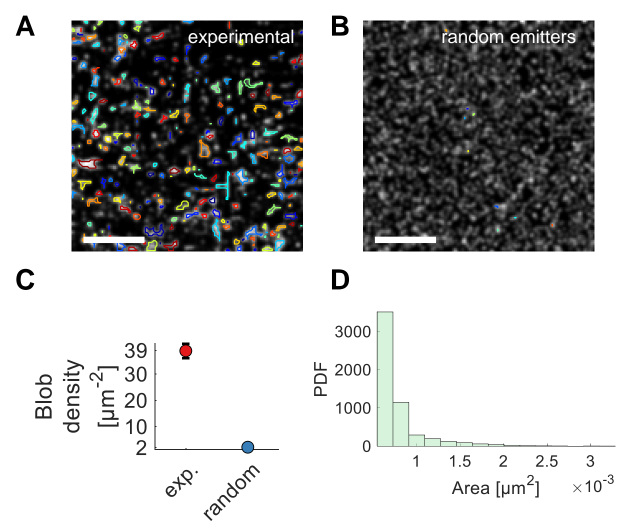
The Rand index ranges from 0 to 1 and gives the fraction of matching pairs of pixels in the ground truth and computed segmentation. Our custom marker-controlled watershed algorithm performs significantly better than other tested algorithms, with a Rand index of about 75%.

To further validate the segmentation algorithm, the blob segmentation on experimental images of chromatin (Supplementary Figure 4A) was compared to images, in which emitters were randomly distributed (Supplementary Figure 4B). On random images, the blob density was ~ 19-fold reduced compared to segmentation on experimental images (Supplementary Figure 4C) and the blob area was about one order of magnitude smaller (Supplementary Figure 4D). These results show that blobs were identified due to the appearance of chromatin as blobs and not due to apparent grouping of randomly distributed emitters.



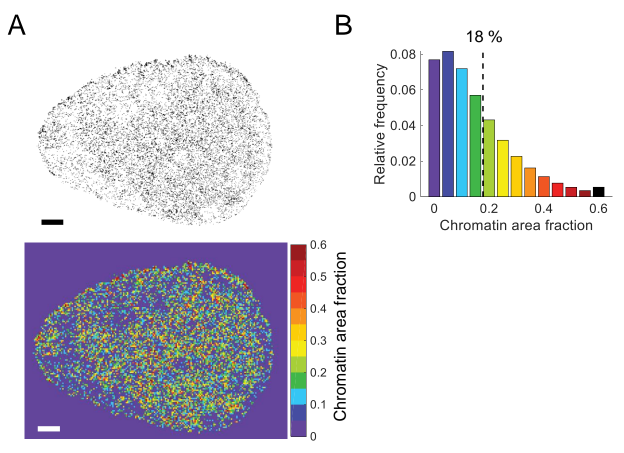
#### Supplementary Figure 3: Performance of segmentation algorithms on super-resolved images 914 of chromatin in vivo. A) Exemplary simulated image for which the ground truth segmentation is 915 known, used to compare different segmentation algorithms. The section in the red triangle is 916 917 magnified in D-I). B) Exemplary experimental image to be compared to the simulated image. C) The Rand index reveals that our custom maker-controlled watershed algorithm significantly 918 919 outperforms all other tested algorithms. Statistical significance assessed from 20 independent segmentation runs by a two-sample t-test (\*\*\* p < 0.001). **D**) Binary image (right) displaying the 920 ground truth segmentation of the simulated image. Different segments are randomly colored for 921 clarity and the magnified region marked in A) is shown. E-I) Exemplary segmentation results from 922 the tested algorithms. 923

- 924 925
- 923





Supplementary Figure 4: Segmentation on images of randomly distributed emitters. A) An 927 exemplary super-resolved image of chromatin in vivo, the identified blobs are overlaid and 928 929 randomly colored. B) Emitters were randomly distributed, and the segmentation algorithm was applied to those images. The number of emitter was matched to the number of beads of the modeled 930 chromosomes within the imaging volume (compare Figure 2B). Scale bar is 0.5 µm. C) The blob 931 932 density for blobs identified on experimental images and images containing randomly distributed emitters. Experimental images contained  $\sim$ 20-fold more blobs than could be identified using 933 random images. D) Area distribution for blobs identified on images containing randomly distributed 934 emitters (mean  $\pm$  std.: (0. 4  $\pm$  0. 2)  $\cdot$  10<sup>-3</sup>  $\mu$ m<sup>2</sup>). 935



936

Supplementary Figure 5: Chromatin area fraction. A) Segmented images are divided into boxes with dimensions 120 nm x 120 nm and the chromatin area fraction is computed for each box. Exemplary, a map of chromatin area fractions is shown color-coded from low to high chromatin density (purple to red). Scale bar is 2  $\mu$ m. B) Histogram of the observed chromatin area fractions. The black line denotes 18 ± 14 % (mean ± std).

# 942 SUPPLEMENTARY NOTE 2

### 943 Suitability of Optical Flow for super-resolution images of chromatin

Optical Flow is used to compute a flow field between two subsequent images in scenarios in which dynamic information cannot be retrieved from single-particle tracking approaches, for example, due to high labeling densities (26). Optical Flow algorithms are generally evaluated with respect to the angular error (AE), a measure for the error in the direction between a computed and a ground truth vector. Likewise, the endpoint error (EE) is a measure for the error in the magnitude. For two computed vectors **a** and **b**, the AE and EE are computed as (25)

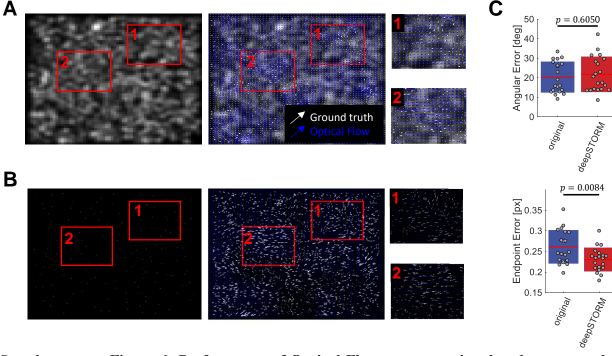
$$AE = \cos^{-1}\left(\frac{ab}{|\mathbf{a}||\mathbf{b}|}\right) \tag{7}$$

$$EE = |\mathbf{a} - \mathbf{b}|$$

Current Optical Flow algorithms achieve sub-pixel EE and AE of around 20° for bulk chromatin 951 imaging. Here, we prove that the Optical Flow algorithm used previously for conventional 952 microscopes (25, 26) results in comparable AE and EE values using super-resolved time series of 953 chromatin. To this end, ground truth data is simulated as described previously (26). A density of 6 954 emitters per  $\mu m^2$  and an acquisition time of 30 ms was used to resemble experimental data 955 (Supplementary Figure 1). The images were either summed up directly in sets of 12 in order to 956 achieve experimental time resolution of 360 ms or first processed by Deep-PALM and then 957 summed. Optical Flow was computed for both sets and the AE and EE were computed. Exemplary 958 simulated images and ground truth vectors, as well as computed flow fields, are superimposed in 959 Supplementary Figure 6A-B. The resulting AE and EE from 20 independent runs are summarized 960 in Supplementary Figure 6C. Optical Flow on super-resolved images did not significantly change 961 the accuracy in direction, however the endpoint error is slightly smaller for flow fields computed 962 963 on super-resolved images. These results thus validate Optical Flow for the use on super-resolution time series of chromatin. 964

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967

968 Supplementary Figure 6: Performance of Optical Flow on conventional and super-resolved
 969 images. A) Exemplary simulated conventional fluorescence microscopy images (left), ground truth
 970 and estimated flow field (middle). Magnified regions as indicated by the red boxes (right). B) As

(8)

A) for images analyzed with Deep-PALM. For visualization, only every eighth vector is shown (Deep-PALM images are up-sampled 8-fold compared to the input images). C) Angular and endpoint error over 20 independent sets of simulated images. Statistical significance was determined by a two-sided t-test.

975 976

# 977 Supplementary Movie 1: Time series of super-resolved chromatin structure and dynamics.

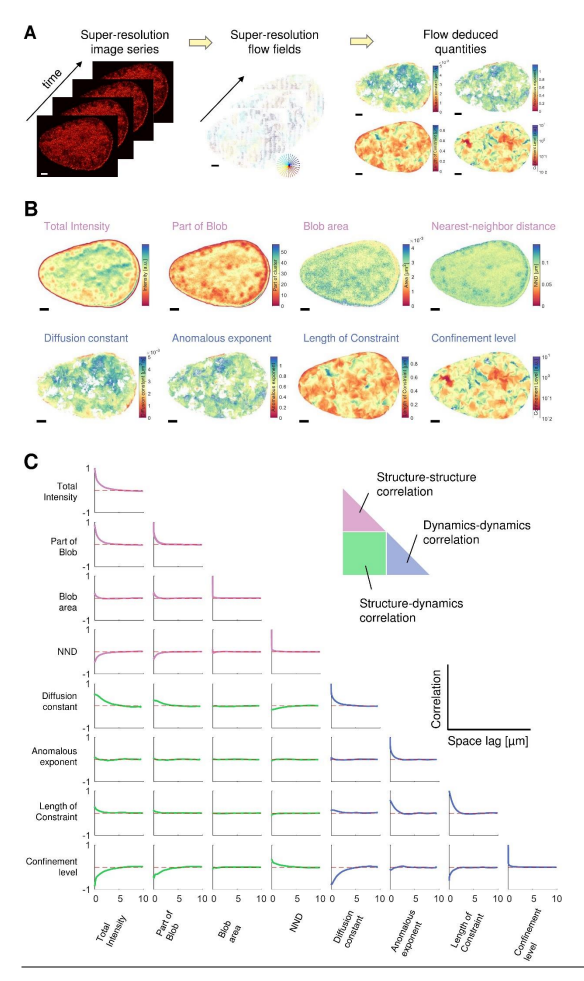
The centroid positions of each identified blob are mapped onto the nucleus and colored according to their nearest-neighbor distance, area, mean displacement direction, and magnitude. Colors are given such that the respective maximum parameter value over the whole image series is red, the minimum parameter value is blue. Colors thus indicate parameter values relative to the parameter range. Displacement direction is color-coded according to the color-wheel shown.



С Α  $\times 10^{-5}$ ×10<sup>-3</sup> 3.5 3 3 5.2 μm<sup>2</sup>] 3 2.5 0 0.2 0.6 0 0.2 0.4 0.6 0.8 0.4 0.8 1 Norm. distance from periphery Norm. distance from periphery В Flow magnitude [µm] 0.09 0.12 NND [µm] 0.08 0.11 0.07 0.1 0.06 0.05 0.09 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Norm. distance from periphery Norm. distance from periphery



986 Supplementary Figure 7: Structural and dynamic parameters are dependent on the proximity 987 to the nuclear periphery. A) The average area, B) NND, C) density and D) flow magnitude versus 988 the normalized distance from the nuclear periphery (0 is on the periphery and 1 is at the center of 989 the nucleus). Line and shaded area denote the mean ± standard error.



#### 992 Supplementary Figure 8: Global spatial correlation of structural and dynamic parameters.

A) Illustration of the Hi-D workflow. A time series of super-resolution images (left panel) is input 993 to the Optical Flow algorithm resulting in flow fields with a pixel size of 13.5 nm (middle panel). 994 995 By trajectory reconstruction and motion classification, quantities describing the underlying bulk motion are computed. B) Eight parameters characterizing the global chromatin structure and 996 dynamics during the whole time series are shown. The structural parameters (upper row) are the 997 total intensity of super-resolved images, the counts how often each pixel was identified as part of a 998 chromatin blob, the average blob area per pixel and the average nearest-neighbor distance for each 999 pixel. Dynamic parameters are the diffusion constant and anomalous exponent, which was 1000 computed by regression of mean squared displacement curves (Materials and Methods), the length 1001 of constraint and the average confinement level. Scale bar is 3 µm. C) The spatial correlation 1002 between all combinations of structural and dynamic parameters over space lag is shown. 1003

1004

## 1006 SUPPLEMENTARY NOTE 3

### 1007 t-SNE and its robustness with respect to distance metrics and perplexity

- 1008 High-Dimensional parameter space is input to the t-SNE algorithm. The underlying principle is that 1009 data points, which are similar with respect to a number of factors (dimensions) lie close in high-
- dimensional space (employing a certain distance metric). The mapping into lower dimensionality
- 1011 (for instance in 2D) by t-SNE is initialized by assigning each point a random position in 2D (Figure
- 1012 6A). Illustratively, a set of springs between all data points exert a repelling or attractive force on
- 1013 each other depending on if the current distances between data points in 2D represent the distances
- between the data points in high-dimensional space. The 2D positions are iteratively refined in order
- 1015 to minimize the divergence between the high-dimensional and two-dimensional distributions.
- 1016 More specifically, a high-dimensional pairwise distance measure can be defined between all points. 1017 The similarity of data points  $x_i$  and  $x_i$  is expressed as the conditional probability,  $p_{ij}$ , that  $x_i$  would
- pick  $\mathbf{x}_i$  as its neighbor under the assumption that neighbors are picked in proportion to their
- 1019 probability density under a Gaussian centered at  $\mathbf{x}_i(31)$ :

$$p_{j|i} = \frac{\exp\left(-\|\mathbf{x}_{i} - \mathbf{x}_{j}\|^{2}/2\sigma_{i}^{2}\right)}{\sum_{k \neq i} \exp(-\|\mathbf{x}_{i} - \mathbf{x}_{k}\|^{2}/2\sigma_{i}^{2})'}$$
(9)

and as the symmetrized conditional probabilities

$$p_{ij} = \frac{p_{j|i} + p_{i|j}}{2n},\tag{10}$$

1021 where n is the number of data points. The variance of the Gaussian,  $\sigma_i^2$ , is determined by a binary 1022 search in order to obtain a user-specified value for the *Perplexity*, which is defined as

$$Perplexity(P_i) = 2^{H(P_i)}, (11)$$

- 1023
- 1024 where  $H(P_i)$  is Shannon's entropy

$$H(P_i) = -\sum_j p_{j|i} \log_2 p_{j|i}$$
 (12)

- and  $P_i$  is the conditional probability distribution over all other data points given  $x_i$ . The perplexity is, loosely speaking, controlling the number of close neighbors of each point and can have a complex non-linear influence of the resulting distribution of points.
- 1028 The conditional probability  $q_{j|i}$  of two points  $y_i$  and  $y_j$  in the two-dimensional space is modelled by 1029 a t-distribution:

$$q_{ij} = \frac{\left(1 + \|\mathbf{y}_i - \mathbf{y}_j\|^2\right)^{-1}}{\sum_{k \neq l} (1 + \|\mathbf{y}_k - \mathbf{y}_l\|^2)^{-1}}$$
(13)

1030

The algorithm first randomly assigns a position to each data point in two-dimensional space and then iteratively refines the position of data points such as to minimize the Kullback-Leibler (KL) divergence, a natural measure for the mismatch between the joint probability distributions in the high-dimensional space, P, and in the low-dimensional space, Q. Thus, the cost function at every iteration is the KL divergence between P and Q,

$$KL(P||Q) = \sum_{i} \sum_{j} p_{ij} \log\left(\frac{p_{ij}}{q_{ij}}\right),$$
(14)

which is minimized using a gradient descent method. In other words, the two-dimensional position y<sub>i</sub> is modified such as to minimize the KL divergence between P and Q. This minimization scheme depends critically on the conditional probabilities  $p_{ij}$  and  $q_{ij}$  and therefore on the distance  $\|\cdot\|$ between points and the perplexity. We tested the influence of different distance metrics and perplexity values on our data set to exclude artifacts arising through an improper choice of parameters. The following distance metrics were tested:

1042 1) Euclidian distance

$$|x_i - x_j|| = \sqrt{(x_i - x_j)^T (x_i - x_j)}$$
 (15)

1043 2) Mahalanobis distance

$$\|x_{i} - x_{j}\| = \sqrt{(x_{i} - x_{j})^{T} S^{-1} (x_{i} - x_{j})}$$
(16)

1044 where S is the covariance matrix. The Mahalanobis distance reduces to the Euclidian 1045 distance if S is the identity matrix.

1046 3) Correlation distance

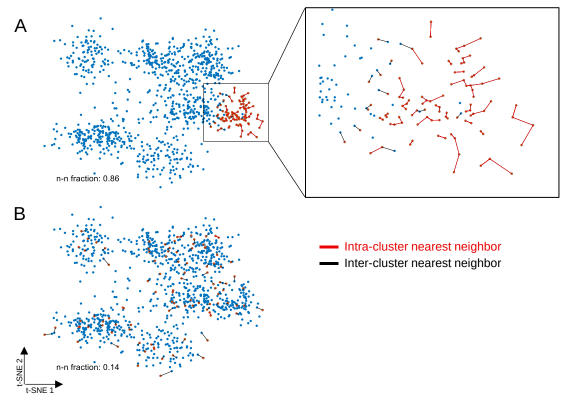
$$\|\boldsymbol{x}_{i} - \boldsymbol{x}_{j}\| = 1 - \frac{(\boldsymbol{x}_{i} - \overline{\boldsymbol{x}}_{i})^{T}(\boldsymbol{x}_{j} - \overline{\boldsymbol{x}}_{j})}{\sqrt{(\boldsymbol{x}_{i} - \overline{\boldsymbol{x}}_{i})^{T}(\boldsymbol{x}_{i} - \overline{\boldsymbol{x}}_{i})}\sqrt{(\boldsymbol{x}_{j} - \overline{\boldsymbol{x}}_{j})^{T}(\boldsymbol{x}_{j} - \overline{\boldsymbol{x}}_{j})}}$$
(17)

1047 where  $\overline{x}$  denotes the average value of  $x_i$ .

Perplexity values were varied from 30 to 200. Note that t-SNE is a probabilistic approach since points are initially distributed randomly in two dimensions. Therefore, multiple runs on the same data set might result in varying results.

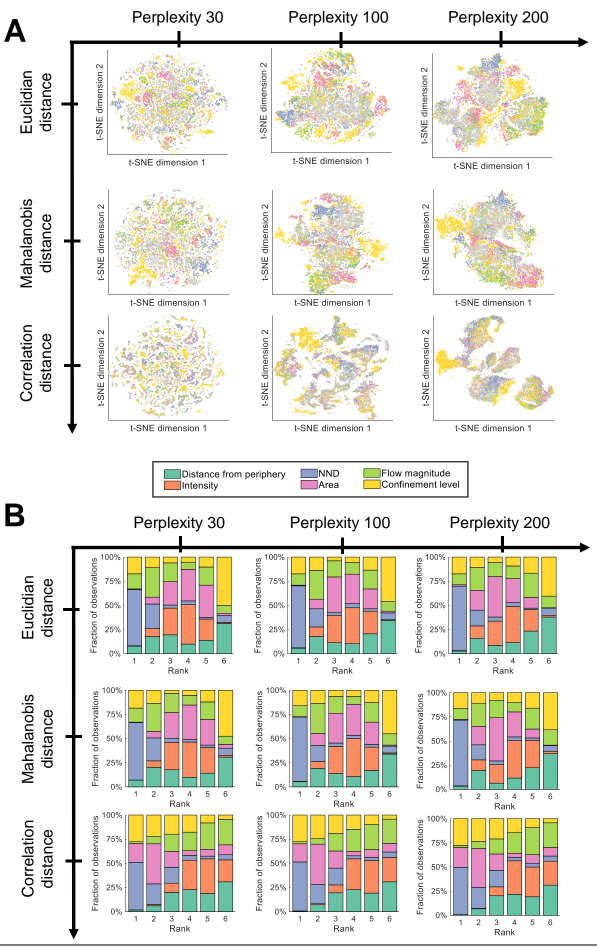
Exemplary t-SNE maps are shown in Supplementary Figure 10A for the tested perplexity values 1051 and distance metrics. Maps are colored corresponding to Figure 6D. The embedding of points in 1052 two dimensions varies greatly among the scenarios. However, the probability of two points being 1053 1054 nearest neighbors is largely conserved and thus ranking of input parameters yield similar results across the employed scenarios (Supplementary Figure 10B). Rankings are especially robust when 1055 the distance metric is the Euclidian distance or the Mahalanobis distance. When the correlation 1056 distance is employed, the rankings slightly change. Especially, the area of blobs seems to be more 1057 prominent than the flow magnitude, in contrast to rankings when one of the other distance metrics 1058 is used. However, rankings change only to a small extent among the different distance metrics and 1059 perplexity values and the presented results are therefore free of artifacts of t-SNE or algorithm-1060 dependent parameters. 1061

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Supplementary Figure 9: Clustering illustration of points within a subset based on nearest-1065 neighbors in t-SNE maps. Dimension reduction using t-SNE results in point clouds in which 1066 nearest neighbors in high-dimensional space are conserved and mapped as nearest neighbors in two 1067 dimensions. Points at which a number of interest exhibits unusually high/low values are identified 1068 (red points). Mapping of two points as nearest neighbors in reduced t-SNE space can be because 1069 nearest neighbors belong to the same subset of high/low values of a variable of interest. To quantify 1070 this empirical characteristic, the number of nearest neighbors within the same subset in t-SNE space 1071 is counted relative to the total number of nearest neighbors of all points within the subset. A) A 1072 subset mostly contains its own nearest neighbors (86% of nearest neighbors of all points in the 1073 subset are contained within the subset). The magnification shows nearest neighbor connections 1074 within the subset (red lines) and between points in the subset and points not contained in the subset 1075 (black lines). B) Points within the subset are distributed over the whole t-SNE space and thus do 1076 not form a grouped region. The fraction of nearest neighbor links within the subset is small (14%) 1077 compared to the clustered case. 1078

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1081 Supplementary Figure 10: t-SNE for different distance metrics and perplexity values. We

1082 tested the influence of different distance measures between the high-dimensional data points as well

as variations in the perplexity from 30 to 200 and found that our data are robust to changes in these parameters within the explored range. A) Exemplary t-SNE maps and B) feature ranking

1085 considering various distance metrics and values for the perplexity parameter applied to our data.