Protocol

Two-parameter single-molecule analysis for measurement of chromatin mobility

This protocol provides a two-parameter analysis of single-molecule tracking (SMT) trajectories of Halo-tagged histones in living adherent cell lines and unveils a chromatin mobility landscape composed of five chromatin types, ranging from low to high mobility. When the analysis is applied to Halo-tagged, chromatin-binding proteins, it associates chromatin interaction properties with known functions in a way that previously used SMT parameters did not.

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HIGHLIGHTS

Single-molecule imaging to assay/compare motions of chromatin and nuclear factors

Protocol walks the user through imaging, detection, and tracking of single molecules

Two-parameter analysis allows assay of motions of chromatin-interacting molecules

Translates motions into interaction with functionally diverse chromatin domains
Protocol
Two-parameter single-molecule analysis for measurement of chromatin mobility

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SUMMARY
This protocol provides a two-parameter analysis of single-molecule tracking (SMT) trajectories of Halo-tagged histones in living adherent cell lines and unveils a chromatin mobility landscape composed of five chromatin types, ranging from low to high mobility. When the analysis is applied to Halo-tagged, chromatin-binding proteins, it associates chromatin interaction properties with known functions in a way that previously used SMT parameters did not.

For complete information on the use and execution of this protocol, please refer to Lerner et al. (2020).

BEFORE YOU BEGIN
The wide range of chromatin compaction states visible in super-resolution and electron microscopy (Ou et al., 2017; Ricci et al., 2015) elicits a variety of mechanical and electrostatic constraints over the motions of chromatin domains in living cells (Dion and Gasser, 2013; Nozaki et al., 2017). Single-Molecule Tracking (SMT) allows the assessment of the displacements of individual Halo (derived from haloalkane dehalogenase) fusion proteins in living cells, with a high spatiotemporal resolution (Liu and Tjian, 2018).

The following protocol performs the characterization of Halo-tagged histone H2B trajectories in living cells with two parameters: the radius of confinement, estimating the area explored by the molecule, and the average displacement, representing the average frame-to-frame jump (Lerner et al., 2020). Though positively correlated, the two parameters present systematic deviations allowing the resolution of five chromatin mobility groups corresponding to functionally different chromatin domains (Lerner et al., 2020).

The protocol then uses the measured chromatin mobility landscape as a benchmark and scales the mobility of diverse chromatin-binding proteins, thus assaying the interactions with different chromatin domains while scanning the genome (Lerner et al., 2020).
Construct Halo fusion expression vectors

**Timing:** 3–4 days

This protocol has been used and validated for various Halo-tagged nuclear proteins: histone H2B, transcription factors, heterochromatin regulators, lamins; and in a variety of somatic adherent cell lines, as well as pluripotent iPSCs (Lerner et al., 2020).

1. Clone the open reading frame (ORF) encoding the protein of interest in a vector containing a Halo cassette (see Key resources table), at the N- or C-terminal end.

**Notes:**

Fusion with Halo, a relatively large protein (33 kDa) might impair the biological function of the protein of interest. To circumvent possible functional interferences, we recommend using a flexible linker sequence between Halo-tag and the protein of interest. In the publication of reference (Lerner et al., 2020), we used a polyGlycine/Serine linker of the following sequence: SGGGSGGGSGGGSGGGGS (Iwafuchi et al., 2020).

The Halo-tagged protein should present a proper nuclear and subnuclear localization (e.g., nucleoli, chromocenters, lamina, ...). Please note that some transcription factors display a normal partial cytoplasmic localization (Raccaud et al., 2019). We recommend validation of the biological activity of Halo fusion proteins through additional approaches: western blot, luciferase reporter assays, Electro Mobility Shift Assays, chromatin-immunoprecipitation, ...

Other labeling strategies for SMT in living cells such as SNAPtag or CLIPtag (Crivat and Taraska, 2012) are also compatible with this protocol.

For the recommended reagents and kits for this step, see Key resources table.

Deliver the vectors into living cells

**Timing:** 48 h

2. Plate cells in the imaging set up compatible with the single-molecule microscopy device (coverslips, 8 well plates, ...). We recommend plating cells to reach a confluency of 70%–80% on the day of imaging. Too high confluency might result in cell overlays which can affect the imaging quality.

3. Using the appropriate transfection system, deliver the expression vectors in the cells.

4. Grow the cells in the adequate cell culture medium for 48 h.

**Notes:**

Using a Tetracycline-inducible lentiviral expression vectors (such as TETO-FUW) ensures the good delivery of the vector in a wide range of cell types, allows a controlled expression of the transgene, and can conveniently generate stable cell lines expressing the Halo-tagged protein of interest (see Troubleshooting). In the case of cells endogenously expressing the protein of interest, we recommend using expression levels of the Halo-tagged protein near those of the endogenous protein. This can be achieved through clonal selection, with expression levels assayed by western blot, FACS, or RT-qPCR. Alternatively, insertion of the Halo cassette at the endogenous locus through a CRISPR-Cas9 system is a recommended strategy when possible. Other strategies for moderating the transgene expression level are also available (Liu et al., 2018).
We recommend using Phenol red-free medium at all steps of cell culture, to reduce the background during imaging.

We also encourage running a western blot with an antibody specific of the protein of interest to verify the stability of the Halo fusion (see Troubleshooting).

**Computing setup**

© Timing: 1 h

Install MATLAB and upload MATLAB script package.zip (see Key resources table). Extract the .zip file in the folder of your choice. In the home section of MATLAB, hit “Set Path.” Select “Add with sub-folders” and browse to the folder containing the extracted files. Hit “Save.” MATLAB can now find and run all the scripts necessary for this analysis, notably scatplot.m, SLIMfast.m, SMT_classifier.m, Two_Parameter_SMT.m, densitydifferences.m, msdanalyzer.m.

**Notes:**

A digest version of this protocol can also be found in the Scripts, Lerner-Zaret, as STEP BY STEP simplified protocol.pdf. The protocol does not require an extensive knowledge of MATLAB; however, we recommend performing the tutorial available at Matlab.com, particularly the section about setting up paths and running scripts. We ran this analysis successfully on a computer with ad minima a 2.70 GHz Processor and 6 GB of RAM. Higher specifications can lead to reduced times of analysis.

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Recombinant DNA     |        |            |
| pENTR4/HALO         | Eric Campeau | Addgene W876-1 |
| Chemicals, peptides, and recombinant proteins | | |
| Janelia Fluor 549 Halo ligand | Promega | GA1110 |
| Critical commercial assays | | |
| MinElute PCR purification kit | QIAGEN | 28004 |
| MinPrep             | QIAGEN | 27106 |
| MAXIPREP            | QIAGEN | 12263 |
| Gel extraction kit  | QIAGEN | 28706 |
| Gibson Assembly kit | NEB    | ES510 |
| T4 DNA ligase high concentration | NEB    | M0202T |
| Software and algorithms | | |
| Fiji is Just ImageJ (Fiji) version 2.0.0-rc/1.51f | ImageJ | https://imagej.net/ |
| MATLAB              | MathWorks | https://www.mathworks.com/products/matlab.html |
| MATLAB script package.zip containing: | This paper | 2D_tracking_matlab_package Mendeley https://doi.org/10.17632/hxh4ttbptk.1 |
| Step by step simplified protocol.pdf, a digest version of the protocol | This paper | STEP BY STEP simplified protocol.pdf |
| @msdanalyzer, containing scripts used for the analysis of T-MSD curves | (Tarantino et al., 2014) | https://tinevez.github.io/msdanalyzer/ |

(Continued on next page)
MATERIALS AND EQUIPMENT

The JF549 Halo ligand should be stored as a 5 μM stock solution in DMSO, at −20°C and protected from the dark, and aliquoted in small volumes to avoid repetitive freeze/thaw cycles. The ligand can be thawed the day of the experiment and diluted directly at 5 nM in the cell culture medium (see Halo ligand section).

Single-molecule imaging for this protocol is done with a microscope performing Highly Inclined and Laminated Optical sheet (HILO) illumination for optimal background-to-noise ratio, and with a camera compatible with a 100 Hz acquisition. Other imaging methods such as lattice light sheet (Chen et al., 2014a) have not been tested for this protocol. In (Lerner et al., 2020), we used a Nanoimager S from Oxford Nanoimaging Limited (ONI), comporting a temperature and humidity controlled chamber, a scientific Complementary metal-oxide–semiconductor (sCMOS) Hamamatsu Orca Flash 4 V3 camera with a 2.3 electrons rms read noise at standard scan, a 100X, 1.49 NA oil-immersion objective and a 561 nm green laser. Alternatively, we used a Nikon Eclipse Ti microscope equipped with a 100X oil-immersion objective lens (NA, 1.4), and EMCCD (iXon3, Andor) (Chen et al., 2014b).

STEP-BY-STEP METHOD DETAILS

Halo ligand staining

© Timing: 30 min
At this step, cells are incubated with the fluorescent Halo ligand, which penetrates the cells specifically and covalently bind to the Halo-tag. Janelia Fluorophore 549 (JF549, see Key resources table) has been broadly used for single-molecule applications and provides an excellent background/signal ratio. The Halo fusion protein is detected with a laser excitation optimal for JF549 (549 nm).

1. Incubate the cells for 15 min with 5 nM of fluorescent Halo ligand.
2. Remove the medium containing the fluorophore in excess and replace with fresh medium (free of Phenol Red). Repeat three times.
3. Leave the cells in the incubator for 15 min to release unbound fluorophore.
4. Remove the medium and replace with fresh medium (free of phenol red).

△ CRITICAL: We recommend concealing the sample from the light as much as possible to avoid photobleaching and leaving the light off in the cell culture hood.

Single-molecule imaging

△ Timing: several hours

This is the step of detection of fluorescent single molecules of the Halo-tagged protein of interest, using HILO imaging.

5. Set the sample in the observation chamber, ideally allowing the control temperature and CO₂ levels. If no CO₂ control is possible, we recommend imaging for relatively short times (<1 h), alternating between samples in and out from a cell incubator to avoid impairing cell function.
6. Using a laser compatible with JF549 fluorophore excitation (peak at 549 nm) and starting at a moderate power coupled with a 100 ms exposure, detect the JF549 fluorescence in the sample. Adjust focus and illumination angle for optimal signal.
7. Take a single snapshot of the nucleus/nuclei to be acquired. It can be used for further detection of nuclear structures (e.g., nucleoli, condensates, …), as well as for assaying the intensity of fluorescence prior to single-molecule acquisition.
8. Set the camera exposure time at 10 ms and increase progressively the laser power, until single molecules are visible. Photobleaching the nucleus might be necessary and take several seconds (Methods Video S1).
   a. Adjust the illumination angle, z-axis and laser power to obtain an optimal molecule density and background-to-signal ratio (see Troubleshooting and Methods Videos S1 and S2).
   b. As the elevated laser power causes fast photobleaching, start recording the single molecules as soon as possible. Typically, 5,000 frames are recorded (total acquisition duration: 50 s).

Notes:

For step 6, we recommend starting with a low laser power. Typically, we use values ranging from 50 to 500 W/cm² for imaging. If you observe rapid photobleaching, decrease the laser power.

For step 8, the optimal range for acquisition lies between imaging as seen in Methods Videos S1 and S2. We recommend acquiring between 20–30 individual nuclei to ensure having enough data for further analysis. Our protocol was established using a 10 ms exposure for single-molecule imaging. Exposure times between 5 and 30 ms should be compatible with this protocol but might lead to differences in the accuracy of the measure of the average displacement and radius of confinement leading to two-dimensional plots different of the examples provided here.
Single-molecule tracking analysis I: tracking

© Timing: several hours

This is the first step of computational analysis for the single-molecule imaging data. The input is a stack of .tif files generated at the single-molecule imaging step (Figure 1A).

We preferentially use tracking software SLIMfast.m (Normanno et al., 2015; Teves et al., 2016), based on the tracking algorithm Multiple Target Tracing (MTT) (Sergeé et al., 2008), both publicly available (please also cite these publications whenever referring to our protocol). Alternatively, this protocol is also compatible with tracking files generated by TrackMate (ImageJ), which can be directly used as inputs in the following section.

For each .tif stack, this step outputs a .txt tracking data file containing a series of successive spatio-temporal coordinates (x, y, t) for each single molecule trajectory (or motion track) over the time of acquisition. The tracking step here is using the MATLAB-based SLIMfast.m script (Teves et al., 2016), a modified version of MTT (Sergeé et al., 2008), both publicly available. Please cite these publications whenever referring to this protocol.

9. Start MATLAB and run SLIMFast.m

10. In SLIMFast, hit Load > Batch, and select the .tif stacks to analyze. Once all the files loaded, adjust the Optics parameters: fluorophore (emission for JF549=580 nm), objective (Numerical Aperture, NA) and camera (Pixel size) used during single-molecule imaging (Figure 1B and Methods Video S3).

Methods video S3. Detection of molecules by SLIMfast.m, related to single-molecule tracking analysis I (step 10)
This video displays how to add the tracking script package to the path, and how to detect molecules in successive frames of the .tif single-molecule imaging stacks.

11. Hit START. A progress bar (“localization”) will pop up, representing the progressive detection of particles in the successive frames of each .tif stack (Methods Video S3). Once detection is finished, a signal (“Done!”) pops up. This step produces several output files sharing the name of the initial .tif stack, among which a .mat file.

12. In SLIMfast, Load > Particle data and select an individual .mat file. An overlay of all particles detected over the course of acquisition is displayed (Figure 1C and Methods Video S4).

   a. To track the molecules frame-by-frame, in the overlay menu, hit OPT > Tracking, and change the Max. expected Diffusion Coefficient to 3 μm²/s.
   b. Hit “GEN TRAJ” in the overlay window. A “Processing Frame” progress bar pops up. Once completed, the output files can be saves in the directory of choice. This step generates a .txt file with the same name than the initial .mat file, which will be converted as .csv and used as an input for the two-parameter mobility analysis (Figure 1D).

Methods video S4. Single-molecule tracking by SLIMfast.m, related to single-molecule tracking analysis I (step 12)
This video displays how to upload the file containing the successive position of molecules in the .tif stacks, and to run the MTT tracking algorithm to establish trajectories.

ıld Pause Point: Step 11 can be timely and can take up to several hours to complete for large data sets. For example, for 50 cells, using the minimal configuration indicated in the Computing setup section, it can take up to 3–4 h.
For step 10: tracking algorithm MTT performs the detection of single particles by fitting a two-dimensional Gaussian to the Point Spread Function (PSF), over sliding windows of 7 x 7 pixels.
The shape and size of the PSF is function notably of the emission wavelength of the fluorophore in use (580 nm for JF549) and on the Numerical Aperture (NA) of the objective used for imaging (1.45 for the Oni Nanoimager S). In our experiments (Lerner et al., 2020), the PSF was approximately of 0.8. To discriminate false and true detected particles, a probability of false alarm (PFA) is set by default at $10^{-6}$ by the tracking algorithm (Sergé et al., 2008). The Count/photons value is a technical feature intrinsic to the camera used to estimate the number of detected photons which does not affect the tracking parameters. Please note that all parameters can be freely modified by the user if desired.

For step 12, the Maximal expected Diffusion Coefficient value reflects the maximal frame-to-frame displacement expected for single particles, and was set as most accurately reflecting the motions of single particles seen in our experimental conditions. If desired, the user can input different values, which might be relevant if using exposure times different than 10 ms (see previous section).

**Single-molecule tracking analysis II: motion track classification**

- **Timing:** several hours

The two-dimensional mobility analysis is only performed on chromatin-interacting molecules. Thus, during the second step of data analysis, free-diffusing molecules are identified and filtered out by classifying trajectories generated by SLIMfast.m on the basis of their mean-squared displacement (or MSD, Figure 2A, Ernst and Köhler, 2012). The tracking .txt file generated at the previous step are converted to .csv and used as input.

The SMT_Motion_classifier.m script performs the following operations:

The SMT_Motion_classifier.m script performs the following operations: (1) discarding of trajectories shorter than 5 frames to optimize the classification; (2) computing the T-MSD curve (or MSD=$f(t_{lag})$, where $t_{lag}$ is the delay in seconds) for each individual trajectory (Figure 2B); (3) fitting of each full T-MSD curve with a power law distribution (Michalet and Berglund, 2012) the scaling component $\alpha$ (Figure 2B and see Ernst and Köhler, 2012); (4) $\alpha$ -based classification of trajectories: $\alpha \leq 0.7$: confined; $0.7 < \alpha < 1$: Brownian; $\alpha \geq 1$: directed (Figure 3A); (5) in parallel, detection of “Butterfly” trajectories with mixed confined and directed/Brownian behaviors (“Butterfly”), on the basis of a displacement step (or jump) at least 1.5 times superior to the average displacement, and of a sum of displacements bigger than 8 times the average frame-by-frame displacement (Figure 3B); (6) splitting of Butterfly trajectories into confined and Directed/Brownian segments based on geometrical properties: Brownian/directed segments have at least 30% of its points outside of a polygon (convex hull) formed by the points of previous and posterior segments of the trajectory (Figure 3C); (7) classification of confined trajectories (including Butterfly confined segments) based on their average frame-to-frame displacement. Trajectories with average displacement below 100 nm are considered as confined, corresponding to molecules interacting with chromatin (Figure 3D).

For each imaged nucleus, seven “msd_results” .mat files are generated as outputs. They correspond to each class of trajectories: All, Brownian, Directed, Confined with high (confined_High_Jump) or Low (confined_Low_Jump) average displacement, Butterfly Confined (segments_confined) and Directed (segments_directed) segments.

13. Gather the batch of .txt files generated by SLIMfast.m at the previous step (each .txt file corresponds to one initial .tif stack) in a unique folder, which will be made directory for MATLAB (Methods Video S5).

14. Run csv_converter.m (Methods Video S5), which converts each .txt file into a .csv file containing the following information (column-by-column): iteration (1-n, with n number of rows of the csv
15. Run SMT_Motion_Classifier.m. Select the batch of tracking .csv files to analyze. To display additional figures, or change parameters, please refer to the User Manual provided in the MATLAB script package. After execution, the script outputs a new folder starting with “MSD_results” and containing 7 .mat files (starting with “msd_results”).

Pause Point: Step 15 can be timely for large datasets and depending on your computing power, this can be used as a Pause point for an hour or more (see Troubleshooting). For example, for 100,000 trajectories, using the minimal configuration indicated in the Computing setup section, the script can take up to 2–3 h.

Notes:

Thresholds for detection of butterfly trajectories were set empirically, optimizing the values of thresholds for an optimal classification, as assayed by visual inspection of the confined and directed parts. For a first-time analysis, we recommend using the defaults parameters. Nevertheless, advanced users can freely modify all parameters of the data analysis in lines 25–54 of the SMT_Motion_Classifier.m, to vary the conditions.
Figure 3. Basis of trajectory classification by SMT_motion_classifier.m

(A) The scaling component α-T-MSD curve is used to keep confined trajectories, corresponding to chromatin associated molecules, and discard facilitated and pure diffusion trajectories, corresponding to molecules non-associated with chromatin.

(B) Composite ("butterfly") trajectories, presenting confined and diffusing components, are identified on the basis of two criteria: a displacement (d) larger than the average frame-to-frame displacement (Δd), and a sum of displacements (Σd) larger than 8 times Δd.

(C) "Butterfly" trajectories are segmented into their confined and Brownian/directed parts. Each number represents the successive time of detection of the single molecule. Example of a fragment considered as confined (from time 5 to 6) and directed (from time 6 to 7) are displayed. The Brownian/directed segment must fulfill the condition that a minimum percentage of its points lie outside the polygon (convex hull) formed by the points of previous and posterior segments of the trajectory.

(D) To discard confined tracks with high displacements, a threshold for a maximal frame-to-frame displacement of 100 nm is set. T-MSD curves are shown here before and after filtration for all confined tracks in a single nucleus expressing FOXA1-HALO. T-MSD curves after filtration (right panel) present a confined aspect only.
Users also have the option not to use the Confined segments of the butterfly tracks in the analysis, just by selecting the Confined_low_jump.mat file in the following part.

For step 13, we provide sample .csv files with our MATLAB script package

**Single-molecule tracking analysis III: measure of the radius of confinement and of the average displacement, and mapping of the data density**

© Timing: several hours

The third step of single-molecule data analysis consists of assessing the two parameters used for characterizing the mobility of individual molecules. The Two_Parameter_SMT.m script is run on trajectories of nuclear proteins in a chromatin-bound state, defined as the pool of confined trajectories and of the confined segments of the Butterfly trajectories (Figure 3 and Figure 4A).

To measure the radius of confinement (RC), the Two_Parameter_SMT.m script fits each T-MSD curves with a circle confined diffusion model (Wieser and Schütz, 2008) to provide (1) the corresponding radius of confinement RC; (2) the diffusion coefficient at short time scales D; (3) a constant offset corresponding to the localization precision limit (Figure 4B).

The average displacement (AVE) is computed by measuring the average Euclidean distance between two consecutive positions of the molecule in each motion track (Figure 4C).

The different mobility chromatin domains are arbitrarily defined based on the correlation between the two parameters, and based on the data density: very low (vLMC) and low (LMC) mobility chromatin are seen as two marked deviations from the line of proportionality. Along the correlation line, Intermediate (IMC) and High (HMC) are distinguished based on the bimodality of the distribution of the radius of confinement: IMC presents lower RC than HMC (Figure 5A). The limits for radius of confinement (RC) and average displacement (AVE) are set as follows (Figure 5A): very Low Mobility Chromatin (vLMC): 0.020 <RC< 0.030 & 0.015 <AVE< 0.029; Low Mobility Chromatin (LMC): 0.035 <RC< 0.05 & 0.020 <AVE< 0.035; Intermediate Mobility Chromatin (IMC): 0.015 <RC< 0.0375 & 0.029 <AVE< 0.036; High Mobility Chromatin (HMC): HMC 0.0375 <RC< 0.06 & 0.0375 <AVE< 0.06; very High Mobility Chromatin (vHMC) 0.055 <RC< 0.3 & 0.06 <AVE< 0.3.

The outputs generated by the script are (1) trackingdata.csv files, in which each row represents a motion track, with its corresponding RC (column 1), average displacement (column 2) and track duration (column 3); (2) two representations of the radius of confinement versus average displacement as assigned to each motion track: a scatter plot (Figure 4D), and a density scatter plot (Figure 4E) (see Scatplot.m). The scripts also generate a 3D mobility plot with x: radius of confinement, y: average displacement, z: data density as measured by scatplot.m.

16. Run Two_Parameter_MSD and hit “add.” Go to the appropriate directory and open Butterfly_segments_confined.mat and confined_Low_jump_0.1.mat for the set of data to analyze. Hit “Done Selecting Files” and wait for the script to execute.

17. Save the figures generated by the script under the desired format.

**Pause Point:** Step 16 can be timely for large datasets and depending on your computing power, this can be used as a Pause point for an hour or more. For example, for 100,000 trajectories, using the minimal configuration indicated in the Computing setup section, the script can take up to 30 min to 1 h.

**Notes:**
In our model, IMC and LMC were distinguished on the basis of the mobility of heterochromatin constituents. In fact, heterochromatin regulators bind preferentially to vLMC and LMC (Lerner et al., 2020). IMC is bound by transcription factors and heterochromatin regulators and might be constituted by composite chromatin domains.

Boundaries in radius of confinement and average displacement chosen for each population were chosen as non-overlapping squares covering the best each mobility populations. The user can modify these limits as desired in the script.

As representing the circle best encompassing the motion track, rather than encompassing it strictly, the measurement of the radius of confinement is independent of the track duration (Lerner et al., 2020).

For point 14., the user might also use only confined_Low_jump.mat files, in the case no butterfly tracks have been detected.
Figure 5. Comparing the mobility of a protein of interest with a mobility of reference
(A) Example of a reference mobility plot (H2B-HALO, n = 80,000) and of the mobility plot for a protein of interest (FOXA1-HALO, n = 80,000). The squares represent the boundaries set for each chromatin mobility group. Differences in data density are measured by normalizing the data density in the reference plot by the maximal value in the plot of interest. vL, very low mobility chromatin; L, low mobility chromatin; I, intermediate mobility chromatin; H, high mobility chromatin; vH, very high mobility chromatin.

(B) Density values in the plot of reference and of interest are first normalized by the corresponding maximal density value in the plot (noted max(density)). Then, the Density_differences m script overlaps the two-dimensional density plots of reference (e.g., histone H2B in red) and of interest (in green), and subtracts the average density values of the 5 nearest points of reference to each density of interest. Density differences are then multiplied by –1 for intuitiveness. Then positive values for density differences indicate increased binding, while negative values indicate decreased binding.

(C) Data density differences between FOXA1 and H2B in the different mobility groups. Data are represented as means ± SEM. vL, very low mobility chromatin; L, low mobility chromatin; I, intermediary mobility chromatin; H, high mobility chromatin.

Single-molecule tracking analysis IV: comparison of data density respective to histone H2B

© Timing: several hours
This step consists in the quantitative measurement of differences between a reference two-dimensional mobility plot (e.g., histone H2B, wildtype protein, ...) and the two-dimensional mobility plot of a protein of interest (e.g., transcription factor, mutant) (Figure 5A). The trackingdata.csv files generated at the previous step are used as input for this step.

The Density_Differences.m script performs the following operations: (1) measuring the local density in the different chromatin mobility populations of the two-parameter (RC and AVE) scatter plot with scatplot.m and assigning a density number to each trajectory (each individual point in the scatter plot). (2) Normalizing density values by the maximum density value for each condition (Figures 5A and 5B). (3) Computing the density differences between the protein of interest (any Chromatin-Binding Protein) and the reference (Figure 5B), by subtracting the average density of the five nearest points to the density of each trajectory of interest (Figure 5B). (4) Multiplying the result by $1/C_0$ for intuitiveness: in each chromatin mobility population, positive and negative values thus mean that the protein of interest is more or less dense than the reference in the considered chromatin mobility domain, respectively (Figure 5C).

The script generates 3 outputs plots: one individual density plot for the reference and the protein of interest, and a histogram plot displaying the average and standard error in 4 mobility population. The very high mobility population is excluded because the sparsity of the data in that region gives rise to high errors in the measurement.

18. On line 9 of the Density_differences.m script, indicate the number of trajectories to analyze. We recommend running the script on the same number of randomly downsampled data, with $n \geq 10,000$.
19. Run the Density_differences.m script, and first select the trackingdata.xlsx file corresponding to the reference plot (e.g., histone H2B). Then select the trackindata.xlsx file for the protein of interest.
20. Save the figures under the desired format.

EXPECTED OUTCOMES

The two-parameter mobility plot for histone H2B was similar to Figure 4E in several differentiated cell lines (Lerner et al., 2020), but might be dependent on the cell context. For example, in the cited reference, ES cells presented a histone H2B mobility plot of higher mobility than other cell types, with mainly High Mobility Chromatin, and mitotic cells presented an overall decreased chromatin mobility (Lerner et al., 2020). These assessments from the two-parameter analysis agree with independent assessments of chromatin structure (Meshorer et al., 2006; Ou et al., 2017).

QUANTIFICATION AND STATISTICAL ANALYSIS

For the radius of confinement measurement, to discard fitting errors, the trajectories with squared norm of the residual (or RSS) higher than $10^{-5}$. In addition, fitting of T-MSD curves can lead to aberrant values for radius of confinement (e.g., bigger than the radius of the nucleus). Thus, the threshold set for maximal radius of confinement is 300 nm, consistently with the maximal size of chromatin domains seen in super-resolution microscopy (Ricci et al., 2015) and in single-molecule imaging (Nozaki et al., 2017). Less than 5% of trajectories were consistently discarded by these filters.

The fitting of T-MSD curves with a circle diffusion model (Ernst and Köhler, 2012) also provides a constant offset due to the localization precision limit inherent to all the localization-based microscopy methods. In our case, we estimate a localization precision of 13 nm. Tracks shorter than 5 steps were arbitrarily excluded, as it seems a reasonable cutoff to reduce the standard error in the measurement of the average displacement.
When measuring density differences between a two-dimensional plot of reference and for a protein of interest, the plot displays the mean and the standard error of the mean in each chromatin mobility population. Statistical significance was assayed by one-way ANOVA in the original publication (Lerner et al., 2020).

LIMITATIONS
Our protocol has been carried out only on adherent cells. Non-adherent cells pose a challenge for single-molecule imaging because of their constant movements in the medium, even when sedimented. That can be addressed by coating the support with components allowing immobilization of the cell.

TROUBLESHOOTING
Problem
The protein of interest not compatible with a Halo fusion. In our experience, this is particularly true for very large protein (>100 kDa) and resulting in the detection of Halo signal both in the nucleus and the cytoplasm.

Potential solution
This can possibly be circumvented by tagging the other extremity (N- or -C-terminal) of the protein. Alternatively, trying several linkers between the protein and the Halo-tag might render the fusion possible.

Problem
Elevated fluorescence levels, resulting in high background, overlapping molecules, or artifacts (e.g., peri-nuclear inclusions of Halo fluorescence).

Potential solution
Use a low expression strategy for the Halo-tagged protein. For example, in the publication of reference we used Tetracyclin inducible promoters in the absence of activator and tetracycline. Other strategies for low expression of the protein can also be used (Liu et al., 2018). Alternatively, decreasing the concentration of Halo ligand, or incubation time is a good strategy for decreasing background levels.

Problem
No single-molecule are visible during the imaging step.

Potential solution
If no fluorescent signal can be seen at a 100 ms exposure, the problem is probably connected to the settings of the microscope. The 3 key parameters for signal optimization are: (1) laser power, (2) illumination angle, (3) z-position. These need to be finely tuned to find optimal imaging conditions. These parameters might have to be change when switching to another field.

Problem
The SMT_Motion_Classifier script is stuck on “Detecting Butterfly Motions.”

Potential solution
This step can be timely as many operations are simultaneously performed and can sometimes take more than several hours to finish but will not crash. If the number of tracks analyzed is over 100,000, we recommend selecting less of the .csv files, which will make the analysis faster.
RESOURCE AVAILABILITY
 Lead contact
 Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact and corresponding author, Dr. Jonathan Lerner at jler@pennmedicine.upenn.edu.

Materials availability
 This study did not generate any new unique reagents.

Data and code availability
 Scripts used in this study are available on Mendeley https://doi.org/10.17632/hxnhtttxpk.1

SUPPLEMENTAL INFORMATION
 Supplemental Information can be found online at https://doi.org/10.1016/j.xpro.2020.100223.

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AUTHOR CONTRIBUTIONS
 Conceptualization, J.L., P.G.G., K.Z., and M.L.; methodology, J.L., P.G.G., K.Z., Z.L., and M.L.; software, P.G.G., Z.L., J.L., and R.M.; validation, J.L. and P.G.G.; formal analysis, J.L. and P.G.G.; investigation, J.L.; resources, K.Z., Z.L., and M.L.; data curation, J.L. and P.G.G.; writing – original draft, J.L. and P.G.G.; writing – review & editing, J.L., P.G.G., and K.Z.; visualization, J.L. and P.G.G.; supervision, K.Z., M.L., and Z.L.; funding acquisition, J.L., K.Z., and M.L.

DECLARATION OF INTERESTS
 The authors declare no competing interests.

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