Single-cell measurement of higher-order 3D genome organization with scSPRITE

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Although three-dimensional (3D) genome organization is central to many aspects of nuclear function, it has been difficult to measure at the single-cell level. To address this, we developed ‘single-cell split-pool recognition of interactions by tag extension’ (scSPRITE). scSPRITE uses split-and-pool barcoding to tag DNA fragments in the same nucleus and their 3D spatial arrangement. Because scSPRITE measures multivay DNA contacts, it generates higher-resolution maps within an individual cell than can be achieved by proximity ligation. We applied scSPRITE to thousands of mouse embryonic stem cells and detected known genome structures, including chromosome territories, active and inactive compartments, and topologically associating domains (TADs) as well as long-range inter-chromosomal structures organized around various nuclear bodies. We observe that these structures exhibit different levels of heterogeneity across the population, with TADs representing dynamic units of genome organization across cells. We expect that scSPRITE will be a critical tool for studying genome structure within heterogeneous populations.

In eukaryotes, linear DNA is packaged in a 3D arrangement in the nucleus. This includes organization of DNA regions from the same chromosome (chromosome territories), which are further subdivided into megabase-sized, self-associating TADs, based on gene activity (active/inactive or A/B compartments), and local interactions between regulatory elements (enhancer–promoter loops). Additionally, DNA regions from multiple chromosomes are organized around nuclear bodies that form higher-order structural units.

Genome organization in a single nucleus affects various nuclear functions, including DNA replication, transcription, and RNA processing. Indeed, genome structure is known to dynamically change between cell types and in individual cells across time to reflect differences in biological state. For example, during the cell cycle, DNA structure undergoes dramatic rearrangement from open chromatin in interphase to highly condensed metaphase chromosomes. Similarly, gene expression levels are heterogeneous among populations of cells, suggesting that there might be differences in enhancer–promoter contacts present in individual cells in the population.

Currently, most methods used to study nuclear organization measure ensemble structures across millions of cells and can obscure critical information about the genome organization of any given cell. For example, measuring cells across the cell cycle and averaging their DNA contacts would mask cell cycle-dependent dynamics. Additionally, several studies showed that observation of genome structures such as TADs in single cells do not always match structures predicted from ensemble measurements. Accordingly, genome organization observed in bulk assays might not accurately reflect specific structures that exist within biological populations.

The two main techniques for measuring genome architecture of single cells are microscopy and single-cell Hi-C (scHi-C). Microscopy provides the capability to study a broad range of genomic interactions in single cells but is generally limited to measurements of a small number of loci simultaneously and does not provide a genome-wide view. In contrast, scHi-C provides a genome-wide view of nuclear structure in single cells, but it requires specialized equipment (for example, robotics), generates data for low cell numbers, and is limited to low-resolution structures (~10-Mb resolution per cell). Additionally, because scHi-C relies on proximity ligation to measure interactions, it has limited ability to capture long-range and higher-order interactions, such as those organized around nuclear bodies.

To address these technological gaps, we developed scSPRITE to provide comprehensive, high-resolution, genome-wide maps of DNA structure from thousands of single cells. scSPRITE measures both inter- and intra-chromosomal interactions and dramatically increases the number of detected DNA contacts per cell relative to existing methods. To demonstrate its utility, we measured 3D genome structures in 1,000 individual mouse embryonic stem cell (mESC) nuclei and observed chromosome territories, A/B compartments, and TADs in hundreds of single nuclei. We identified higher-order structures in hundreds of single cells, including inter-chromosomal contacts around centromeres, nucleoli, and nuclear speckles. Notably, we identified cell-to-cell heterogeneity in mESC genome structure at different levels of resolution, including at promoter–enhancer contacts of the key pluripotency gene, Nanog. Together, these observations demonstrate that scSPRITE accurately measures genome structure and provides insights into genome organization. We expect that this approach will enable future studies examining the relationship between genome organization and nuclear function in individual cells.

Results
scSPRITE maps 3D structure in thousands of individual cells. To understand 3D genome organization in individual cells, we extended our previously described SPRITE protocol to enable...
single-cell measurements. scSPRITE works as follows: We dissociate cells into a single-cell suspension, crosslink DNA and protein complexes in situ, isolate and permeabilize nuclei, digest DNA using a restriction enzyme and perform two sets of split-and-pool barcoding to (1) tag DNA fragments contained in the same nucleus and (2) tag the 3D spatial arrangement of these fragments (Fig. 1a).

To map all DNA fragments originating from one nucleus, we performed split-and-pool barcoding to generate a unique cell-specific barcode (cell barcode) for all DNA molecules contained in a single nucleus. Briefly, we distributed permeabilized nuclei across a 96-well plate (~200,000 nuclei), where each well contained a unique DNA barcode tag, and performed ligation such that all DNA molecules in the same nucleus were labelled with the same tag. We then pooled nuclei and repeated the split-and-pool process twice more to ensure that the number of barcode combinations (96^2 = 884,736) exceeded the cell number (Methods). Because single nuclei can form aggregates in suspension, we filtered nuclei to remove potential clumps before proceeding to the next step (Extended Data Fig. 1a).

To verify that this approach accurately tags DNA contained in a single nucleus, we tested this first set of split-and-pool barcoding in permeabilized nuclei in a mixed population of human (HEK293T) and mouse (mESC) cells. After split-pool barcoding and sequencing, we clustered reads into groups based on their cell barcodes and computed the percentage of reads that aligned exclusively to the mouse or human genome (Methods). We found that only 3.4% of cells contained reads from both species (Fig. 1b and Extended Data Fig. 1b), indicating that most cell barcodes represent single cells. Because we cannot identify collisions that lead to mixing in the same species, we extrapolate a total collision rate (~10%) from the detected collisions.

Having developed an approach to accurately tag DNA in a single nucleus, we next sought to map these DNA fragments relative to each other in 3D space. To do this, we withdrew a small fraction of the single cell-tagged nuclei (~1,500 nuclei) and sonicated them to generate spatial clusters of chromatin. We then performed three additional rounds of split-and-pool barcoding, such that all DNA fragments contained in a spatial cluster obtained the same barcode combinations, whereas molecules in distinct spatial clusters obtained different combinations. After sequencing, we identified DNA molecules within the same spatial complex by matching all six barcode sequences and all complexes arising from the same nucleus, by matching the first three barcodes (Fig. 1a, Extended Data Fig. 1c and Methods).

To validate the method, we applied scSPRITE to mESCs because their genome structure has been extensively studied\textsuperscript{14,15}, and they display known functional heterogeneity\textsuperscript{24-26}. We sequenced ~1,500 single cells and analytically excluded cell barcodes that were likely to represent cell aggregates using the detected collision rates measured from the previously described mixing experiment (Fig. 1c and Extended Data Fig. 1b). To focus on the most informative single cells, we restricted our analysis to the 1,000 cells containing the highest number of spatial clusters per cell (Methods).

To confirm that spatial barcoding in scSPRITE accurately measures known genome structures, we merged individual cell barcodes from scSPRITE (referred to as ensemble scSPRITE) and compared heat maps to those previously generated by bulk SPRITE in mESCs\textsuperscript{2} (Fig. 1d). We found that these maps are highly similar across all levels of resolution (Pearson correlation r = 0.92, 1 Mb genome-wide; r = 0.97, 200 kb on chr2; r = 0.95, 40 kb across chr6: 48–54 Mb).

Together, our results demonstrate that scSPRITE tags single cells with minimal collisions and accurately measures 3D organization at different levels of resolution. Although we analyzed 1,500 single cells in this experiment, the number of cells analyzed by scSPRITE can be adjusted by modifying the number of rounds of split-and-pool barcoding such that the number of barcode combinations exceeds the number of single cells (>100-fold excess; Methods).

scSPRITE measures multiway interactions in single cells. Because each individual cell contains a single genome, and contacts detected in multiple cells cannot be pooled together (as they are in bulk measurements), single-cell genome structure methods need to maximize the number of contacts detected in each cell. This is the main challenge and limitation for all single-cell genomic methods.

Currently, existing single-cell genome structure methods (for example, scHi-C) use proximity ligation and are, therefore, limited to measuring pairwise DNA contacts\textsuperscript{27-30}. Although these measurements are averaged across multiple cells, this is not possible in a single cell because a specific DNA region can be measured only once per allele. Accordingly, even with perfect efficiency, pairwise methods would be unable to measure all possible contacts present in a given structure (Fig. 1e and Extended Data Fig. 1e). In contrast, scSPRITE captures multiway contacts among DNA molecules, which dramatically increases the structural resolution that can be obtained for an individual cell. This is because the maximal number of interactions that can be captured increases quadratically with the size of a complex\textsuperscript{21} (Extended Data Fig. 1d). For example, if a crosslinked complex contains four DNA fragments, the maximum number of contacts that can be observed by pairwise methods is two, whereas the maximal number of pairwise contacts that can be identified with multiway contacts is six (Extended Data Fig. 1e).

Indeed, we observe an increase in the number of pairwise contacts detected for each cell using scSPRITE (average of 34,992,080 per cell) compared to scHi-C\textsuperscript{27} (average of 375,470 per cell), even though the number of sequencing reads per cell is ~10-fold lower for scSPRITE (average of 83,318 per cell) than for scHi-C (average of 751,172 per cell) (Fig. 1f and Supplementary Note 1). We observe uniform coverage across all 1-Mb bins in virtually all cells and across all 100-kb bins in more than 80% of cells (Fig. 1g) with almost no bias toward any chromosome (with the exception of chromosome 8 due to a trisomy in our cell line; Methods) (Fig. 1h). Notably, we observe low variability in genomic coverage across the analyzed cells (median absolute deviation (MAD) = 14, MAD range: 0–49, median = 35; Extended Data Fig. 1f).

scSPRITE detects chromosome territories and compartments. To determine which DNA structures can be observed in single cells, we generated DNA contact maps from each of the 1,000 individual cells. For every structure identified in the ensemble data, we computed a normalized detection score that reflects how well each single cell contact map resembles this structure compared to a randomized contact map. Briefly, for each structure, we calculated an observed detection score, which defines whether each pair of genomic bins in a structure were in contact. A cell that contains all possible pairwise contacts in a given structure would have a detection score of 1, whereas a cell containing none of the expected pairwise contacts would have a detection score of ~1. We normalized these observed scores to a distribution of scores generated by randomly permuting the locations of each structure (Methods and Supplementary Note 4).

We focused on genomic structures that were previously reported to occur in single cells—chromosome territories and A/B compartments\textsuperscript{4}. Chromosome territories are structures containing high frequencies of intra-chromosomal interactions with minimal inter-chromosomal interactions (Fig. 2a). First, we looked at the contacts between chromosome 1 (chr1) and chr2 and detected clear separation of contacts into chromosome territories in both the ensemble data (Fig. 2a) and in more than 75% of single cells (score > 0; Fig. 2b and Extended Data Fig. 2a). Next, we quantified detection scores for every pair of chromosomes in every cell (Extended Data Fig. 2b and Supplementary Table 1). Although some chromosomes show stronger self-interactions than others, all chromosomes organize into territories (average score = 0.08, s.d. = 0.06; Fig. 2c).
Fig. 1 | scSPRITE—a single-cell method to map DNA structure genome-wide. a, Schematic of scSPRITE protocol. b, Validation of in-nuclei barcoding step on mixed cell population (human–mouse cells); the number of reads for each identified cell barcode ID is plotted. Threshold of >95% single-species reads was applied to identify mouse- or human-only cells; cell barcodes >1,000 reads are plotted. c, Number of contacts (blue), reads (red) and DNA clusters (gray) plotted for the 1,500 cells. Dashed lines represent filtration steps: left of the dashed lines—cell aggregates estimated based on detected collision rate from Fig. 1b; right of the dashed lines—cells with low number of reads/contacts. d, Comparison of merged scSPRITE (upper diagonal, ‘ensemble scSPRITE’) and bulk SPRITE (lower diagonal). Chromosome territories across all chromosomes at 1-Mb resolution (left); A/B compartments on chromosome 2 at 200-kb resolution (middle); TADs within an 18-Mb region of chromosome 6 at 40-kb resolution (right). e, Schematic illustration of multiway interactions (SPRITE-derived methods) and pairwise interactions (proximity ligation methods) and examples of heat maps. f, Number of contacts (top) and number of reads (bottom) obtained from scSPRITE (blue) and scHi-C (gray). g, Genomic coverage per 1-Mb, 100-krb, 40-kb and 10-kb bins in individual 1,000 cells. h, Average number of reads per single cell in 1-Mb bins of each chromosome (n = 1,000 cells). Average (dots) and s.d. (bars) are shown; asterisk marks chromosome with detected trisomy.
and Extended Data Fig. 2c; see Methods for chr8). We observe that 95% of cells contain well-defined territories (Fig. 2d,e), and only a small fraction of cells (<50 cells) do not contain observable chromosome territories (Fig. 2d and Extended Data Fig. 2d) and might reflect cell states containing distinct organization, such as mitotic chromosomes.

Genomes are further divided into A/B compartments, which are intra-chromosomal structures defined by open (A) or closed (B) chromatin states (Fig. 2f). To measure A/B compartment pattern in single cells, we first focused on a region on chr2 that has a well-defined B-A-B compartment switch observed in the ensemble scSPRITE data (Fig. 2f). We calculated the detection score for that region in individual cells and observed segregation of DNA into A/B compartments in more than 65% of single cells (score > 0; Fig. 2g and Extended Data Fig. 2e). Next, using our ensemble data, we defined all regions that correspond to a compartment switch (B-A-B or A-B-A) genome-wide (224 regions; Supplementary Table 2) and quantified their detection scores for each cell (Extended Data Fig. 2f). We observed that individual regions are more variable in single cells than chromosome territories (average score = 0.03, s.d. = 0.06; Fig. 2h and Extended Data Fig. 2f) but are still present in ~95% of cells (Fig. 2i). We looked more closely into three regions with different average detection scores (Region 1, score = 0.12, s.d. = 0.16; Region 2, score = −0.01, s.d. = 0.14; Region 3, score = −0.10, s.d. = 0.11) and observed that the variability in the A/B compartment structure in single cells is indeed higher for the regions with lower detection scores (Fig. 2) and Extended Data Fig. 2g) (i.e., Region 3 > Region 2 > Region 1). This suggests that the detection score metric that we developed is useful to identify cells and regions of variable structures. We observe a small detection bias toward active regions (A compartments, 45% of observed reads versus 39% of expected reads) (Extended Data Fig. 2h).

Together, our results demonstrate that scSPRITE can detect known genomic interactions, such as chromosome territories and A/B compartments in single cells, and can be used to measure structural variability between individual cells.

Inter-chromosomal hubs are organized around nuclear bodies. The nucleus is further organized around various nuclear bodies that form higher-order inter-chromosomal contacts25. These contacts have not been previously explored in single cells at the genome-wide scale because existing single-cell proximity ligation methods are limited in their ability to detect inter-chromosomal contacts14,15,17,26. scSPRITE measures, on average, an almost tenfold increase in the proportion of inter-chromosomal contacts per cell than scHi-C (54% and 6%, respectively) (Fig. 3a), which makes it a suitable method to study higher-order organization in the nucleus. We focused on three types of known inter-chromosomal structures: inactive regions associated with nucleoli, active chromatin around nuclear speckles, and centromeric and peri-centric regions organized into chromocenters.

Inactive DNA hubs are known to organize around the nucleolus, a nuclear body that is formed around transcription of ribosomal DNA (rDNA) regions27. In mESCs, regions on chr12, chr15, chr16, chr18 and chr19 contain rDNA clusters that form nucleolar organizing regions (NORs). We first explored contacts between two NOR-containing regions on two pairs of chromosomes (chr18/chr19 and chr12/chr19) that were previously reported to form strong interactions in mESCs28. We observed similar interaction patterns between these regions in the ensemble SPRITE data and in individual cells (score > 0 in 54% and 61% of cells, respectively) (Fig. 3b,c and Extended Data Fig. 3a,b). We compared the frequencies of contacts detected by scSPRITE (specifically how often these two regions are in the same cluster) to the frequencies of their co-occurrence at the same nucleolus measured by microscopy (where the nucleolus is visualized by nucleolin immunostaining and DNA regions are visualized by DNA fluorescence in situ hybridization (DNA FISH))29. We focused our analysis specifically on 1-Mb regions targeted by DNA FISH probes (three NOR-containing chromosome pairs and two control chromosome pairs) and observed a strong correlation between these datasets (R² = 0.88; Extended Data Fig. 3c), indicating that single-cell measurements generated by scSPRITE are similar to those observed by microscopy. Similarly, we observed a strong correlation between scSPRITE and SPRITE data for these NOR regions (R² = 0.88; Extended Data Fig. 3d). To look at genome-wide interactions of NORs, we quantified the percentage of single cells that contain each nucleolar contact (Extended Data Fig. 3e) and observed that, on average, 38% of cells contained each nucleolar pair (Extended Data Fig. 3e). The most frequent contacts are formed between NORs on chr18 (33–10 Mb) and chr19 (25–28 Mb or 29–37 Mb), which are both observed in more than 50% of cells, and the least frequent contacts are observed between chr15 (67–71 Mb) and chr18 (57–60 Mb), which are observed in less than 20% of cells (Extended Data Fig. 3e). In all cases, we observed that NORs interact more frequently than random non-NOR-containing regions (Fig. 3d).

Next, we looked at active nuclear hubs organized around nuclear speckles—structures enriched in pre-mRNA splicing factors27,30. First, we focused on the previously reported inter-chromosomal interactions formed by precise regions of mouse chr2/chr4 and chr2/chr5 (ref.) and observed these contacts in 53% and 38% of cells, respectively (score > 0; Fig. 3e,f and Extended Data Fig. 3f). Next, we quantified the percentage of single cells that contain each pair of interacting speckle regions (Extended Data Fig. 3h).
We detected speckle interactions in an average of 34% of cells (Extended Data Fig. 3h), with interactions between regions on chr4 (128–142 Mb) and chr5 (112–126 Mb) observed in more than 50% of cells and between chromosome 2 (117–181 Mb) and chromosome 13 (55–58 Mb) observed in less than 10% of cells (Extended Data Fig. 3h). When we calculated the frequency of contacts per 1-Mb bin of every interacting speckle region, we observed that most speckle regions interact more frequently than random regions but less frequently than NORs (Fig. 3d).

Finally, we explored centromeric and peri-centromeric heterochromatin (PCH) regions. Centromeres and peri-centromeres are long stretches of repetitive DNA essential for chromosome stability and segregation and have been shown to come into close proximity to form inter-chromosomal structures called chromocenters (Fig. 3g). Because PCH regions are not mapped in the genome, we focused our analysis on the first 10 Mb of each chromosome. First, we made single-cell contact maps and calculated detection scores for two pairs of PCH regions (chr1/chr11 and chr4/chr11).
Fig. 3 | scSPRITE identifies inter-chromosomal structures genome-wide in hundreds of single mESCs. 

(a) Quantification of inter-chromosomal contacts from the top 1,000 cells by scHi-C (gray) and scSPRITE (blue). The dashed lines represent the mean percentage of inter-chromosomal contacts.

(b) Nucleolar interaction between chr18 and chr19: illustration (left) and ensemble scSPRITE heat map (right); contact map at 1-Mb resolution.

(c) Nucleolar interaction detection scores for 1,000 cells (middle). Box plot where whiskers represent the 10th and 90th percentiles; box limits represent the 25th and 75th percentiles; the black line represents the median; red dots represent single-cell examples (n = 1,000 cells). Representation of structures with max score (+1) and min. score (−1) (left). Single-cell examples (right), plotted as number of DNA clusters at 1-Mb resolution.

(d) Frequency of NOR (blue), speckle (red) and PCH (green) higher-order interactions in comparison to randomly shuffled regions of the same size (gray) in 1,000 individual cells.

(e) Speckle interaction between chr2 and chr4: illustration (left) and ensemble scSPRITE heat map (right); contact map at 1-Mb resolution.

(f) Speckle interaction detection scores for 1,000 individual cells (middle). Box plot where whiskers represent the 10th and 90th percentiles; box limits represent the 25th and 75th percentiles; the black line represents the median; red dots represent single-cell examples (n = 1,000 cells). Representation of structures with max score (+1) and min. score (−1) (left). Single-cell examples (right), plotted as number of DNA clusters at 1-Mb resolution.

(g) Chromosome regions of higher-order structure.

(h) Interaction values per 1-Mb bins.

(i) Mean interaction value of inter-chromosomal PCH contacts (normalized to number of reads per region) for each pair of chromosomes. NOR-containing chromosomes are shown in bold.

The results of these analyses demonstrate that scSPRITE can capture various higher-order contacts reflecting inter-chromosomal interactions across multiple cells and involving structures of different sizes and transcriptional output (active versus inactive hubs). We note that centromere-proximal and nucleolar contacts were not detectable even in the ensemble scHi-C data (Extended Data Fig. 3i). Although the ensemble scHi-C was able to identify speckle interactions, the single-cell interaction maps lacked information on these structures (Extended Data Fig. 3i).

TADs are heterogeneous across individual cells. TADs are intra-chromosomal structures in which contiguous regions of the genome have been shown to interact more with themselves than with surrounding regions. However, these observations are based mainly on bulk measurements, and whether TADs exist in single cells has been debated. Specifically, it is unclear whether the inability to observe TADs in single cells reflects technical limitation
strong evidence for TADs in the ensemble scSPRITE dataset on a region of chromosome 4 (124.8–126.7 Mb) where we observed some of these TADs in every single cell. First, we focused our analysis on a region of chromosome 4 (124.8–126.7 Mb) where we observed strong evidence for TADs in the ensemble scSPRITE dataset (Fig. 4a). Using the genomic locations defined from the ensemble data, we detected TAD-like structures in more than 75% of single cells (score $>0$; Fig. 4b and Extended Data Fig. 4c), suggesting that most individual cells contain this specific TAD structure with the same boundaries.

To explore the heterogeneity of TAD structures in single cells, we performed two analyses. First, we looked at the average representation of all TADs in each cell by averaging the TAD detection score for each region (identified in the ensemble dataset) in each individual cell; we found that most cells contain TADs (95% of cells with score $>0$; Fig. 4c). Second, we explored whether individual TADs are more or less variable across individual cells by averaging the TAD detection score for individual TADs across cells. We found that most TADs are highly variable between cells (65% of cells with score $<0$; Fig. 4d) and noticed that highly variable TADs are not randomly distributed but cluster in shared genomic regions (variable TAD regions; Fig. 4e and Extended Data Fig. 4d).

Fig. 4 | TADs are heterogeneous units present in the genomes of individual mESCs. a, TAD structure between 124.8 Mb and 126.7 Mb of chr4: illustration (left) and scSPRITE heat map (right); pairwise contact map at 40-kb resolution. b, TAD detection scores for 1,000 cells (middle). Box plot where whiskers represent the 10th and 90th percentiles; box limits represent the 25th and 75th percentiles; the black line represents the median; red dots represent single-cell examples ($n = 1,000$ cells). Representation of structures with max score (+1) and min. score (−1). Single-cell examples (right), plotted as number of DNA clusters at 40-kb resolution. c, Normalized detection scores across all 1,000 cells per each TAD detected in ensemble scSPRITE data; red line marks score $= 0$. d, TAD detection scores across all TADs detected in ensemble scSPRITE data per single cell; red line marks score $= 0$. e, TAD detection scores across 1,000 cells (clustered based on score similarity pattern); columns represent the strength of TAD detection scores for all TADs detected across chr4 in ensemble scSPRITE; gray bar indicates the variable region described in Extended Data Fig. 4c. f, Ensemble heat maps across the 39.4–41.4 Mb region of chr4 made by subtracting the normalized contacts in Group 2 from Group 1 (Fig. 4f). g, Difference contact map across 39.4–41.4 Mb of chr4 made by subtracting the normalized contacts in Group 2 from Group 1 (Fig. 4f). Insulation scores for cells in Group 1 (purple) and Group 2 (green) are plotted.

of current single-cell methods (for example, low-resolution structures) or if these DNA structures are not present in individual genomes. Because scSPRITE generates higher-resolution structures in individual cells, we asked whether it can detect TADs in single cells.

We first defined all TADs present in mESCs using the ensemble scSPRITE data (Supplementary Table 3), which are similar to TADs defined from Hi-C data (Pearson correlation $r = 0.70$; Extended Data Fig. 4a,b). We used these genomic coordinates to score each of these TADs in every single cell. First, we focused our analysis on a region of chromosome 4 (124.8–126.7 Mb) where we observed strong evidence for TADs in the ensemble scSPRITE dataset (Fig. 4a). Using the genomic locations defined from the ensemble data, we detected TAD-like structures in more than 75% of single cells (score $>0$; Fig. 4b and Extended Data Fig. 4c), suggesting that most individual cells contain this specific TAD structure with the same boundaries.
studies are required to define if these cell-to-cell variabilities and similarities define structural differences at the level of TADs in single cells. More specifically, detection TAD-like genome organization in individual cells and identifying which regions of the genome are present in different cells in the population.

At least two distinct—and mutually exclusive—structural states in most major structural changes between these two groups of cells are defined in those cells are distinct from the ensemble population. When focusing exclusively on cells that contain this alternative TAD, we found that the A/B compartments boundary of the ensemble-defined A/B compartment (Fig. 4f). We confirmed that these distinct structural states are not explained by differences in cell cycle (Extended Data Fig. 4g). When focusing exclusively on cells that contain an alternative TAD that spans the 120.0–121.0-Mb region in chr5. Shown are cells containing (red) or lacking (blue) the contact between the Tbx3 locus and SE –300 kb. Each position refers to a 40-kb bin. Asterisks denote statistical significance (P < 0.0001, unpaired two-sided t-test with Welch’s correction) between the two groups at the specified positions (n = 1,000 random bootstrap groups for each of the two groups). Error bars represent 1 s.d. Schematic illustrating differences in structure when a gene of interest lacks (left) or contains (right) the long-range enhancer interaction. ChIP-seq, chromatin immunoprecipitation with sequencing.

organization at that region (Fig. 4f). Specifically, we detected a population of cells that contain an alternative TAD that spans the boundary of the ensemble-defined A/B compartment (Fig. 4f and Extended Data Fig. 4f). When focusing exclusively on cells that contain this alternative TAD, we found that the A/B compartments defined in those cells are distinct from the ensemble population (Fig. 4f,g). We confirmed that these distinct structural states are not explained by differences in cell cycle (Extended Data Fig. 4g) or by other major structural changes between these two groups of cells (Extended Data Fig. 4b). This suggests that this region is present in at least two distinct—and mutually exclusive—structural states in different cells in the population.

Together, our results demonstrate that the scSPRITE method can detect TAD-like genome organization in individual cells and identifies structural differences at the level of TADs in single cells. More studies are required to define if these cell-to-cell variabilities and region-to-region differences are functionally relevant and if they are characterized by other features such as transcription, specific chromatin marks or weak insulation boundaries (Supplementary Note 2).

scSPRITE detects heterogeneity across long-range contacts. We next asked if scSPRITE could detect structural changes that reflect biologically significant long-range DNA contacts, such as the interactions between promoters and super enhancers (SEs) or between regions enriched in polycomb group proteins (PCGs)36. SEs are large domains enriched in H3K27 acetylation that are thought to modulate gene expression by forming loops with promoters37. Bulk genome-wide studies have shown that SEs can form long- and short-range interactions with the same promoter38,39, but it remains unclear whether these interactions occur simultaneously in the same cell. Similarly, DNA regions bound by PCGs have been shown

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**Fig. 5 | Heterogeneous structural states formed by Nanog and Tbx3 loci in individual mESCs.**

**a.** Representation of the Nanog locus and its DNA interactions with SEs: 122.2–122.8-Mb region in chr6 with corresponding ChIP-seq tracks for H3K27ac and H3K4me3; Nanog–SE interaction (black line).

**b.** Representation of Tbx3 locus and its DNA interactions with Lhx5: 120.0–121.0-Mb region in chr5 with the corresponding ChIP-seq tracks for H3K27me3 and H3K4me3; Tbx3–Lhx5 interaction (black line).

**c.** Normalized contact frequency plot between Nanog locus and 122.2–122.8-Mb surrounding region in chr6. Shown are cells containing (red) or lacking (blue) the contact between the Nanog locus and SE –300 kb. Each position refers to a 40-kb bin. Asterisks denote statistical significance (P < 0.0001, unpaired two-sided t-test with Welch’s correction) between the two groups at the specified positions (n = 1,000 random bootstrap groups for each of the two groups). Error bars represent 1 s.d.

**d.** Normalized contact frequency plot between Tbx3 locus and 120.0–121.0-Mb surrounding region in chr5. Shown are cells containing (red) or lacking (blue) the contact between the Tbx3 locus and Lhx5. Each position refers to a 40-kb bin. Asterisks denote statistical significance (P < 0.0001, unpaired two-sided t-test with Welch’s correction) between the two groups at the specified positions (n = 1,000 random bootstrap groups for each of the two groups). Error bars represent 1 s.d.

**e.** Schematic illustrating differences in structure when a gene of interest lacks (left) or contains (right) the long-range enhancer interaction. ChIP-seq, chromatin immunoprecipitation with sequencing.
to interact across long distances to regulate gene expression; however, it remains unclear how heterogenous these long-range interactions are in a population of cells.

We focused on two examples of long-range interactions in mESCs: (1) the Nanog locus, a key pluripotency factor in embryonic stem cells whose promoter interacts with multiple enhancers over a broad range of distances (up to 300 kb)\(^37,40\) (Fig. 5a); and (2) the Tbx3 locus, a transcription factor involved in the maintenance of pluripotency\(^41\) whose locus interacts with another PcG-enriched gene, Lhx5 (760 kb downstream) (Fig. 5b).

We selected cells with coverage over the Nanog and Tbx3 regions of interest and split them into two groups based on whether we observed a contact between the target locus and the long-range enhancer (300 kb upstream for Nanog and 760 kb downstream for Tbx3) (Extended Data Fig. 5a). We computed the frequency of contacts between the target locus and all 40-kb bins for each group of cells (Fig. 5c,d). We noticed that, in the group with long-range interactions detected, short-range interactions were significantly weaker (P < 0.001) and, on average, three times less frequent (Fig. 5c,d). Additionally, we observed that detected long-range interactions span across a TAD border identified in the ensemble dataset for both Nanog and Tbx3 examples (Extended Data Fig. 5a,d). We confirmed that the observed structural differences were not caused by technical differences (for example, number of reads in each group of cells) (Extended Data Fig. 5b,e) or different cell cycle phases (Extended Data Fig. 5c,f).

Our results demonstrate that, in cells where either the Nanog or Tbx3 locus contacts the long-range region, the locus is less likely to form a contact with the short-range region (and vice versa) (Fig. 5e). Surprisingly, we detected long-range and short-range interactions in a similar number of cells, suggesting that both of these states are present at similar frequencies in mESCs. Whether such heterogeneity is a more global occurrence or restricted to specific loci (for example, transcription factors regulating pluripotency), and what (if any) functional role these distinct structures might play, remain to be determined (Supplementary Note 3).

Discussion

We have described scSPRITE, a method to generate high-resolution, genome-wide maps of 3D DNA organization in thousands of single cells. scSPRITE expands the toolkit of genome-wide, single-cell sequencing-based methods with an approach that enables high-resolution structural views across a broad spectrum of DNA interactions into high-throughput contact maps of the entire genome. In contrast to existing methods, scSPRITE does not require specialized equipment, techniques or training, and provides increased resolution from a lower number of sequencing reads across a larger number of cells. Because of this, we expect that it will expand the availability of single-cell genome structure measurements to any molecular biology laboratory. Additionally, we expect that scSPRITE can be scaled to work with as few as hundreds or as many as several thousands of cells simultaneously.

Our results reveal several novel insights about the heterogeneity of genome organization in mESCs. Specifically, we detected long-range higher-order interactions of both active (nuclear speckle) and inactive (centromeres and nucleolar contacts) chromatin regions as well as heterogenous organization of TADs and enhancer-promoter contacts between individual single cells. We note that our experiments were performed in a population of mESCs cultured using a ‘2 inhibitor’ (2i) cocktail that is thought to promote ground state pluripotency and display more homogenous expression profiles across single cells\(^22\,24\) (Supplementary Note 3). Nonetheless, our results suggest that, even in these conditions, nuclear organization can be heterogeneous. Whether these cell-to-cell differences in 3D structure affect gene expression or have other functional significance remains to be determined.

Although our initial study focused on mESCs, scSPRITE can be applied to different cell types or homogenized tissues that are composed of mixed cell populations. One of the current challenges with studying complex tissues (for example, brain) or disease states (for example, tumors) is the heterogeneity of their cellular composition. The application of scSPRITE to such cell populations will enable studies of intrinsically heterogeneous systems and provide an accurate global view of their 3D genome organization. Accordingly, we expect that scSPRITE will provide the field with a path toward understanding the relationship between 3D genome organization and genome function in single cells.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at [https://doi.org/10.1038/s41587-021-00998-1](https://doi.org/10.1038/s41587-021-00998-1).

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Methods

Cell types and culture conditions. We developed scSPRITE using mouse and human cells, focusing primarily on mESCs because their genome structure has been extensively studied. We seeded a male embryonic stem cell line (hps5 derived from V6.6 embryonic stem cell line, provided by K. Plath) and cultured them in serum-free 2i/LIF medium as previously described. We suspect that this mESC line displays trisomy in chromosome 8 because the average number of reads aligning to chr8 is about 33% greater than the average number of reads across the other chromosomes (Fig. 1e).

HEK293T, a female human embryonic kidney cell line transformed with the SV40 large T antigen, was obtained from the American Type Culture Collection (no. CRL-1573) and cultured in complete media consisting of DMEM (no. 11960-092, Gibco, Life Technologies) supplemented with 10% FBS (Seradigm Premium Grade III FBS, VWR), 1× penicillin–streptomycin (Gibco, Life Technologies), 1× MEM non-essential amino acids (Gibco, Life Technologies), 1 mM sodium pyruvate (Gibco, Life Technologies) and maintained at 37°C under 5% CO₂. For maintenance, 80,000 cells were seeded into 10 ml of complete media every 3–4 d in 10-cm plates.

scSPRITE protocol. Cell crosslinking. Media from mESCs was removed and washed once with 1× PBS. Cells on the 10-cm plates were then trypsinized using 2 ml of 0.025% trypsin-EDTA (pre-warmed to 37°C). Plates were incubated for 3 min, and the trypsined cells were mixed by pipetting to break up any clumps. We added 8 ml of pre-heated wash solution (DNEM/F12 + BSA, pre-warmed to 37°C) to the cell plate to inactivate trypsin and wash cells to a conical tube. Cells were centrifuged at 330g for 3 min, and the supernatant was discarded. Cells were washed once with 1× PBS at a ratio of 4 ml of PBS per 1×10⁶ cells and centrifuged again at 330g for 3 min. After the wash, 4 ml of 2 M glycine was added per 1× PBS and allowed to incubate for 2 min while mixing at 1,600 r.p.m. for 30 s every 5 min.

Cell lysis and nuclei preparation. Crosslinked cells were thawed from –80°C and were kept on ice during the cell lysis procedures. Initially, 1.4 ml of lysis buffer #1 (50 mM HEPEs pH 7.4, 1 mM EDTA pH 8.0, 1 mM EGTA pH 8.0, 140 mM NaCl, 0.25% Triton X-100, 0.5% IGEPAL CA-630, 10% glycerol, 1 mM dithiothreitol (DTT), 1 mM spermidine and 1 mM adenosine triphosphate (ATP)) was added per 1×10⁶ cells. The cell solution was mixed thoroughly by pipetting to remove clumps. The cells in DSG solution were gently shaken for 45 min at room temperature. After incubation with DSG, 200 µl of 2.5 M glycine was added per 1 ml of DSG solution previously added to quench the reaction, and the tube was gently shaken for 5 min at room temperature. Cells were then centrifuged at 1,000g for 4 min, and the supernatant was discarded. Cells were washed with 1× PBS at a ratio of 4 ml of PBS per 1×10⁶ cells and centrifuged again at 1,000g for 4 min. After the wash, 4 ml of 1% formaldehyde (16% wt/vol formaldehyde ampules, Life Technologies, no. 28908, prepared in pre-warmed (37°C) 1× PBS) was added per 1×10⁶ cells to the conical tube, and the solution was mixed thoroughly by pipetting to remove clumps. The cells in formaldehyde solution were then gently shaken for 10 min at room temperature. After incubation with formaldehyde, 200 µl of 2.5 M glycine was added per 1 ml of DSG solution previously added to quench the reaction, and the tube was gently shaken for 5 min at room temperature. Cells were then centrifuged at 1,000g for 4 min, and the supernatant was removed. Cells were twice washed with cold 1× PBS + 0.5% BSA (wt/vol) solution, and centrifugation was done at 4°C. Cells were once washed with 1× PBS at 900g for 3 min, and the supernatant was removed. Cells were then resuspended in a 1.5 ml of 10 mM Tris-HCl pH 8, 1.5 mM EDTA, 1.5 mM EGTA, 200 mM NaCl, 1×10⁶ cells and centrifuged at 900g for 4 min at 4°C, and the supernatant was removed. Next, 40 µl of 20% Triton X-100 was added to the same tube to quench the reaction, and the solution was left mixing on the same instrument at 1,200 r.p.m. for 60 min at 37°C. Lastly, 30 µl of 5,000 U ml⁻¹ HpyCH4V (NEB, no. R0820L) was added to the same tube to allow for DNA to be digested in-nuclei. In nuclei digestion, the reaction was performed for 4 h at 37°C while shaking at 1,200 r.p.m. HpyCH4V is a 4-base pair (bp) restriction enzyme that performs blunt-end cutting at TCGA sequences. This particular enzyme was chosen because it was able to perform in-nuclei enzymatic restriction digestion and eliminated the need to perform any additional DNA strand repair steps after restriction digest. After 4 h of restriction digest, the average DNA fragment size was 823 bp.

After digestion, nuclei were pelleted at 900g for 2 min, the supernatant was removed and nuclei were washed three times with 1× PBS, 1× EdTA, 1 mM MgEGTA and 0.1% Triton X-100 solution at 90°C for 2 min. After the washes, the nuclei concentration was adjusted by loading 6 µl of nuclei suspension to a disposable hemocytometer (4-Chip Disposable Hemocytometer, Bulldog Bio, no. DHC-N420). After determining nuclei concentration, 5×10⁶ nuclei were transferred by pipetting into a new 1.5 ml LoBind Eppendorf tube. In this new tube, 25 µl of d-tail reaction buffer and 10 µl of Klenow fragment were added to the nuclei (both reagents were part of NEBNext d-tailing Module (NEB, no. E6053L)). The tube was filled to 250 µl using nucleus-free H₂O, and d-tailing was performed in-nuclei at 37°C for 90 min while shaking at 1,200 r.p.m. the reaction was then stopped with the addition of 200 µl of 1× PBS, 50 mM EDTA, 50 mM MgEGTA and 0.1% Triton X-100. The nuclei pellet was spun down at 900g for 2 min and washed twice using 400 µl of 1× PBS, 1× EdTA, 1× MgEGTA and 0.1% Triton X-100 solution. After the washes, the nuclei were resuspended in fresh 1× PBS, 1× EdTA, 1× MgEGTA and 0.1% Triton X-100 solution, and nuclei concentration was determined again using the hemocytometer, as described previously.

In-nuclei combinatorial barcoding. To uniquely identify DNA sequences originating from the same cell, combinatorial barcoding was performed in-nuclei (Fig. 1a). In our specific experiments, we used three rounds of combinatorial barcoding in the following order: ‘DNA phosphate modified’ (DPM), ‘odd’ tagging and ‘even’ tagging (these tags are described in the original SPRITE paper). The resulting tags were pre-loaded onto a 96-well plate, with each well containing 2-µl of a uniquely barcoded cocktail (PIC) was added per 1×10⁶ cells. We note that the probability that any two cells will have a ‘collision’ is defined by a Poisson distribution with a mean (λ) defined by the number of cells divided by the number of barcode combinations. The probability of observing two or more cells with an expected cell ‘collision’ (where cells obtain the same barcode from the same cell, combinatorial barcoding was performed in-nuclei (Fig. 1a).

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Sonication. Next, 1,500 nuclei were placed into a Covaris microtube-15 and filled to 15 µl using lysis buffer #3. The Covaris tube was placed in the Covaris M220 Focused-ultrasonicator, and sonication was performed for 2 min on specific settings (water temperature 6°C, incident power 30 W, duty cycle 3:3) to release DNA complexes from nuclei. The tube was then removed from the instrument and set at a temperature of 10°C for 5 min. At this step of the protocol, it is important to proceed with all the sonicated nuclei, as sampling them further will lead to a loss of nuclei fragments and will prevent the analysis of DNA structure in single cells. We also recommend adjusting the sonicated number of cells to sequencing abilities to achieve satisfactory coverage per cell.

N-hydroxysuccinimide beads coupling. After sonication, sample containing crosslinked DNA complexes was coupled to N-hydroxysuccinimide (NHS) beads as previously described. Briefly, NHS-Activated Magnetic Beads (Life Technologies, no. 88826) were activated for coupling. First, 600 µl of NHS beads were washed with and resuspended in 1 ml of Fusion buffer (0.2% sodium 10 mM Tris–HCl pH 7.5, 0.5 mM EDTA, 0.5 mM EgTA and 0.1% Triton X-100, filled to the final volume with RLT (Qiagen, no. 79216)). This reaction was performed for 60 min with mixing at 1,600 r.p.m. for 30 s every 5 min. Afterwards, the reaction was stopped by adding 60 µl of RLT2 buffer to remove residual beads and 1 ml of 1M HCl, and the supernatant was withdrawn and placed in a 1.5-ml LoBind Eppendorf tube. The tube was placed on a DynaMag-2 magnet, and the supernatant was removed. The beads were washed once with 600 µl of ice-cold 1M HCl, and the supernatant was removed again and replaced with 600 µl of ice-cold 1x PBS. After removing 1x PBS, the beads were resuspended in 500 µl of 1x PBS + 0% SDS. Additionally, 85 µl of 1x PBS + 0% SDS was added to the previously sonicated nuclei solution, mixed, and added to the bead solution. The complexes were then coupled to NHS beads on an Eppendorf ThermoCycler C overnight at 4°C while shaking at 1,200 r.p.m. After coupling, the flowthrough was removed, and 600 µl of NHS beads were withdrawn and placed in a 1.5-ml LoBind Eppendorf tube. The tube was placed on a DynaMag-2 magnet, and the supernatant was removed. The beads were washed once with 600 µl of ice-cold 1M HCl, and the supernatant was removed again and replaced with 600 µl of ice-cold 1x PBS. After removing 1x PBS, the beads were resuspended in 500 µl of 1x PBS + 0% SDS. Additionally, 85 µl of 1x PBS + 0% SDS was added to the previously sonicated nuclei solution, mixed, and added to the bead solution. The complexes were then coupled to NHS beads on an Eppendorf ThermoCycler C overnight at 4°C while shaking at 1,200 r.p.m. After coupling, the flowthrough was removed, and 600 µl of 1M Tris–HCl pH 7.5, 0.5 mM EDTA, 0.5 mM EgTA and 0.1% Triton X-100 was added to the beads to quench the remaining NHS groups; this was done at 4°C at 1,200 r.p.m. After sonication the beads were washed, the flowthrough was removed, and the beads were washed twice in cold RLT2 buffer (0.2% sodium laureyl sarcosinate, 1 mM EDTA, 1 mM EgTA, 10 mM Tris–HCl pH 7.5, 0.1% Triton X-100, 0.1% NP-40, filled to the final volume with RLT (Qiagen, no. 79216)). This was followed by three washes in M2 buffer (50 mM NaCl, 20 mM Tris–HCl pH 7.5, 0.2% Triton X-100, 0.1% NP-40, 0.2% sodium deoxycholate). The beads were then resuspended in a mix of M2 buffer and H2O (58% M2, 42% H2O) to attain a total volume of 1,125 µl of M2 buffer, H2O and beads.

Spatial barcoding/complex-specific barcoding. Next, spatial barcoding of the DNA complexes on beads was performed as described previously. First, the bead solution was well-mixed and loaded into each well of a 96-well plate (11.2 µl of bead solution per well). Each well of the plate contained 2.4 ml of M2 buffer, H2O and beads. The reaction was performed for 60 min with mixing at 1,600 r.p.m. for 30 s every 5 min. Afterwards, the reaction was stopped by adding 60 µl of RLT2+ buffer to each well before pooling the solutions of each well into a 25-ml reservoir. Each well was then resuspended once with 100 µl of RLT2+ buffer to remove residual beads and pooled into the same 25-ml reservoir. The solution was then transferred to a 15-ml conical tube, which was then placed on a magnet to remove most of the RLT2+ buffer from the beads. With about 2 ml of RLT2+ buffer remaining, the beads were resuspended and transferred to a LoBind 1.5-ml Eppendorf tube, which was placed on a DynaMag-2 magnet to remove the remaining RLT2+ buffer from the beads. This process of split-pool barcoding on beads was repeated until the three additional tags were added. After the last round of split-pool barcoding was completed, the beads were resuspended in 600 µl of Myk buffer (20 mM Tris–HCl pH 8.0, 0.2% SDS, 100 mM NaCl, 10 mM EDTA, 10 mM EgTA, 0.5% Triton X-100) after the washes.

We performed three rounds of spatial tagging because it provided sufficient barcode combinations to uniquely label DNA complexes coming from each individual cell. Briefly, the mouse genome contains 2.5 × 108 nucleotides, which, when divided per the average fragment size of DNA after digestion (823 bp), results in 5.04 × 109 DNA fragments per cell. If we do three rounds of barcoding, we provide 884,736 number of combinations, which exceeds the number of DNA molecules 3.4 times. Notably, during scSPRITE barcoding, we distributed clusters of DNA molecules, not single molecules, so the actual number of barcode combinations will exceed the number of spatial clusters much more than our calculation.

Library preparation. To ensure that we capture all information coming from single cells, we need to sequence all DNA molecules that were bound to the beads. The bead solution was split equally into ten LoBind Eppendorf 1.5-ml tubes, with each tube containing 60 µl of beads in Myk buffer. Next, an additional 32 µl of Myk buffer and 8 µl of proteinase K (NEB, no. P81075) were added to each tube. All ten tubes were placed on an Eppendorf ThermoCycler C, and reverse crosslinking proceeded overnight at 60°C while shaking at 1,200 r.p.m. Next, the tubes were placed on a DynaMag-2 magnet, and the Myk and proteinase K solution was transferred to ten new LoBind Eppendorf tubes. The beads from each of the tubes were washed once with 20 µl of H2O and then transferred to the same tube containing each respective Myk and proteinase K solution. DNA from each of the tubes were purified using the Clean & Concentrator-5 columns (Zymo, no. D4004) using 5x binding buffer to increase yield. Purified DNA from each column was eluted in ten new Eppendorf 1.5-ml tubes using 12 µl of H2O. Each of the tubes were filled to 30 µl with 15 µl of Qubit FastStart High-Fidelity 2x Master Mix (NEB, no. M0395S), 1.5 µl of 20x EvaGreen (Biotium, no. 31000-T), 1.25 µl of 25 µM indexed Illumina primers and 0.3 µl of H2O. Real-time PCR amplification proceeded for 14 cycles, which was when the libraries entered exponential amplification but had not plateaued. After amplification, each of the libraries was diluted four-fold before running on a 1% agarose gel (Life Technologies, no. G402001) with an E-Gel 1-Kb Plus DNA Ladder (Life Technologies, no. 10480890) as a reference. After the run, the gel was cut between 300- and 1,000-bp marks to remove primer dimers, small non-specific amplicons and long DNA amplicons. Libraries from the gel were purified using a Gel Purification Kit (Zymo, no. D4002) as described by the manufacturer, and 20 µl of H2O was used to elute libraries off the column.

To estimate the number of unique molecules in our libraries, the molarity of our libraries was determined using the concentration of our library from Qubit 3.0 Fluorometer (using the Qubit dsDNA HS Assay Kit) and the average library size (bp) using an Agilent TapeStation 2200 (using the Agilent High Sensitive D1000 Screen/tape and reagent kit) to estimate the yield of each library. This, in addition to estimated losses during library purification, allowed us to estimate the number of unique molecules in our libraries. The libraries were sequenced with a read depth of 2.4X to ensure that we are able to map the DNA contained in each cluster.

Sequencing analysis pipeline. The full barcode sequence was identified by combining the DNA tag sequence label with the begins of Read 1 and the reads from the scSPRITE library made from ~1,500 cells. From the FastQC report, we observe a normal distribution of GC content per sequence (Read 1: normal distribution between ~15% and 71%; Read 2: normal distribution between ~27% and 59%).

Alignment and filtering of reads. The trimmed reads containing the full six-barcode sequence were mapped to pre-indexed mm9 reference genome using STAR 2.6.1 using the following parameters: --outFilterMultimapNmax 50 --outFilterScoreMin OverLread 0.30 --outFilterMatchNminOverLread 0.30 --outFilterIntronMotifs None--aligntronMax 50000--alignMatesGapMax 1000--genomeLoad NoSharedMemory–outReadsUnmapped Fastx–alignIntronMin 80–alignSJDBoverhangMin 140. All reads that were aligned to a mm9 non-unique region of the genome were removed by excluding alignments that mapped to regions by the ComputeGenomeMask program (read length = 35 nucleotides). After these filtration steps, all BAM files that corresponded to the same sample but contained different Illumina primers at sequencing were pooled together before cluster identification (Extended Data Fig. 1c).

Cluster barcode and cell barcode identification. To identify SPRITE clusters, all reads that contained the same six-barcode sequences were grouped together into a single cluster. All reads containing the same six-barcode sequences that started at the same genomic position were excluded to remove possible PCR duplicates. This led to 161,989,473 remaining reads. Once identified, a SPRITE cluster file was generated where each line contained the cluster barcode name and corresponding
in-nuclei barcoding), we performed an in-nuclei part of the scSPRITE experiment that are mixed together during scSPRITE (from crosslinking until the end of the full scSPRITE procedure (nuclei isolation, DNA digestion and in-cell barcoding), we focused only on cell barcodes from further analysis. Reads were sorted into individual cells based on cell-specific barcodes, and we focused only on cell barcodes that had more than 1,000 reads per cluster.

Next, we calculated the percentage of reads that aligned to each genome for each identified cell barcode. We categorized cell barcodes as mouse- or human-derived when they contained more than 95% single-species reads and as mixed when they contained less than 95% single-species reads. We then calculated the fractions of human-only, mouse-only and mixed-cell barcodes (Fig. 1b) and reported the percentage of mixed-cell barcodes. Detected collusion rate was further used to estimate thresholds used for cell filtering (Extended Data Fig. 1c and Methods) and to calculate total collision rate. Total collision rate represents an estimation of all possible collisions and relies on the assumption that cells from the same species show similar collision rates as cells of the same species. The best alignment was taken into consideration, and if reads align equally well, they were considered as multimers and removed from further analysis. Reads were sorted into individual cells based on cell-specific barcodes, and we focused only on cell barcodes that had more than 1,000 reads per cluster.

Next, we calculated the size distribution of DNA clusters per cell and remove large clusters (>10,000 reads per cluster) from further analysis. We previously reported that clusters larger than >10,000 reads per cluster contain less information about higher-order chromosome structures (heterochromatin, TADs) and most likely contain big chunks of nucleic that are composed of several chromosomes. We consider them less informative for the type of DNA interactions/structures (background) and, therefore, remove them from the further analysis. Excluding all reads in the >10,000-read clusters led to 83,318,292 remaining reads that were used for all downstream analyses.

Human–mouse mixing experiment. To determine the percentage of single cells that are mixed together during scSPRITE (from crosslinking until the end of in-nuclei barcoding), we performed an in-nuclei part of the scSPRITE experiment using cell types from different species—mouse and human. We perform only the in-nuclei barcoding step because we previously showed that the spatial barcoding step used in bulk SPRITE leads to minimal collisions if the total number of NHS beads is in an excess to the total number of clusters in a sample (their mixing human–mouse experiment detects more than 99% of reads aligning to one species). mESCs (bbs) and human cells (HEK293T) were harvested and resuspended into a single-cell solution, and then 30 × 10^6 cells per each cell type were mixed together in equal quantities and crosslinked, digested, DA-tailed and barcode-in-nuclei as described above. Additionally, for the experiments described in Extended Data Fig. 1b, we mixed equal numbers of mouse and human cells after crosslinking but (1) before digestion or (2) after digestion, or (3) we proceeded to the next step without non-nuclei barcoding done (ODPM – odd + even) toward the number of pairwise contacts between any two bins, we also generated pairwise contact frequency matrices for genome-wide (1-Mb resolution) and A/B compartment (200-kb resolution) for both ensemble scSPRITE and SPRITE but using clusters containing fewer than 1,000 reads per cluster. Downweighted pairwise contact frequency matrices were used for TAD comparison (40-kb resolution) for both ensemble scSPRITE and SPRITE but using all clusters.

Generation of single-cell heat maps from scSPRITE. Similarly to ensemble scSPRITE, single-cell contact frequency matrices were generated at 1-Mb and 40-kb resolution. Contact frequency cell barcode files were identified by grouping clusters together that contained the same DPM, first even barcode combinations and 2 × 10^6 cells. Next, nuclei were filtered through a 10-µm filter (pluriStrainer); 300 nuclei were removed as a new sample and reverse crosslinked; and we proceeded as described above. Next, 10% of the total purified libraries were sequenced using MiSeq; reads were then aligned to combined human/mouse genome using STAR alignment (hg19 and mm10 reference genomes). The best alignment was taken into consideration, and if reads align equally well, they were considered as multi mappers and removed from further analysis. Reads were sorted into individual cells based on cell-specific barcodes, and we focused only on cell barcodes that had more than 1,000 reads per cluster.

Note 4. Detection scores for 3D genome structures. Detection scores were calculated to identify various 3D genome structures in single cells. These structures included chromosome territories, A/B compartments, TADs, centromere interactions, nuclear speckle interactions and nuclearic interactions. Each score reflects how clearly defined a given structure is in a single cell. The scores were calculated using a binary contact matrix for each cell, which defined whether each pair of genomic bins was in contact in that cell. For example, a clearly defined chromosome territory in a single cell consists of chromosomes interacting more with themselves than with each other (illustration in Fig. 2g).

To normalize detection scores, an expected detection score was calculated for each 40-kb genome bin (using the ‘insolation’ file output by the matrix2insulation.pl script).

Detection scores for 3D genome structures. Detection scores were calculated to identify various 3D genome structures in single cells. These structures included chromosome territories, A/B compartments, TADs, centromere interactions, nuclear speckle interactions and nuclearic interactions. Each score reflects how clearly defined a given structure is in a single cell. The scores were calculated using a binary contact matrix for each cell, which defined whether each pair of genomic bins was in contact in that cell. For example, a clearly defined chromosome territory in a single cell consists of chromosomes interacting more with themselves than with each other (illustration in Fig. 2g).

To normalize detection scores, an expected detection score was calculated for each 40-kb genome bin (using the ‘insolation’ file output by the matrix2insulation.pl script).
chromosomes, excluding combinations between the same chromosomes (for example, chr1-chr1), amounting to 171 combinations (Supplementary Table 1) from binary matrices at 1-Mb resolution (171 combinations because chrA-chrB = chrB-chrA).

- **Compartments:** observed intra-compartment contacts / (total possible intra-compartment contacts) / (total possible inter-compartment contacts). Genome-wide scores were calculated for all 224 regions across all chromosomes in which we detected a compartment switch in the ensemble dataset (for example, chr1 has 21 regions and chr3 has zero regions) (Supplementary Table 2). A compartment switch is defined as a transition between A to B or B to A to B' compartments. Scores were calculated from binary matrices at 1-Mb resolution.

- **TADs:** observed intra-TAD contacts / (total possible intra-TAD contacts) / (total possible inter-TAD contacts). Genome-wide TAD scores were calculated from 1-Mb from TAD boundary regions, and these were calculated for all 2,602 TAD boundary regions that we detected in our ensemble dataset (Supplementary Table 3). Scores were calculated from binary matrices at 40-kb resolution.

- **Centromere interactions:** observed centromere–centromere contacts / (total possible centromere–centromere contacts) / (total possible centromere–non-centromere contacts). Centromere interactions were defined as interactions between positions 3 Mb and 13 Mb of each chromosome.

- **Nuclear speckle interactions:** observed speckle–speckle contacts / (total possible speckle–speckle contacts) / (total possible speckle–non-centromere contacts). Nuclear speckle interactions were defined as interactions among the following nuclear speckles regions: chr2 (164–174 Mb, 177–181 Mb), chr4 (128–142 Mb, 147–155 Mb), chr5 (112–126 Mb), chr6 (123–127 Mb), chr11 (95–103 Mb, 115–121 Mb), chr13 (55–58 Mb), chr15 (76–79 Mb) and chr17 (25–30 Mb).

- **Nucleolar interactions:** observed nucleolar–nucleolar contacts / (total possible nucleolar–nucleolar contacts) / (total possible nucleolar–non-nucleolar contacts) / (total possible non-nucleolar–nucleolar contacts) / (total possible non-nucleolar–non-nucleolar contacts). Nucleolar interactions were defined as interactions among the following nucleolar regions: chr12 (5–17 Mb, 25–32 Mb), chr14 (3–6 Mb, 67–71 Mb), chr16 (5–8 Mb), chr18 (3–10 Mb, 13–24 Mb, 25–33 Mb, 39–42 Mb, 57–60 Mb) and chr19 (11–24 Mb, 25–28 Mb, 29–37 Mb, 48–53 Mb, 58–61 Mb).

Calculation of MAD scores for scSPRITE. For each single cell in scSPRITE, we calculated the number of reads in each 1-Mb bin for every chromosome genome-wide (chr1–19). Once these reads were counted, we calculated the MAD value for each cell based on the number of reads in each 1-Mb bin genome-wide to determine the variability of coverage.

Analysis of higher-order structures. Comparison of inter-chromosomal versus inter-compartment contacts. The percentage of inter-chromosomal and inter-compartment contacts can separately be calculated from cells that were analyzed from the single-cell data in scHi-C and from the filtered 1,000 cells from scSPRITE (cluster size threshold of <100,000 reads per cluster). For scHi-C, because every cluster is a pairwise contact, we counted the number of pairwise contacts that were intra-chromosomal contacts (two contacts in the same chromosome) and inter-compartmental contacts (two contacts coming from different chromosomes). For scSPRITE, we counted all pairwise contacts per cluster, where the number of pairwise contacts per cluster can be expressed as a binomial coefficient of ‘n choose 2’, where n is the number of reads per cluster. From this, we then counted the number of intra-chromosomal and inter-compartmental contacts. This was repeated for all clusters in each cell. The percentage of inter-compartmental contacts was determined by dividing the number of inter-compartmental contacts by the sum of the number of intra- and inter-compartmental contacts.

Frequencies of higher-order inter-chromosomal interactions. To determine the frequency of centromeric, speckle or nucleolar interactions in single cells, we used following metrics: (1) percentage of cells that contain a given interaction in each 1-Mb bin and (2) normalized mean interaction value. The percentage of cells containing centromere-proximal, speckle or nucleolar interactions is determined by looking through the filtered 1,000 single cells, focusing on clusters below 10,000 reads per cluster and counting the number of cells containing at least one interaction between all 1-Mb bins (i and j) in the given centromere, speckle and nucleolar regions, respectively. The genomic regions of these higher-order structures were defined previously in the section titled ‘Detection scores for 3D genome structures’. To determine the expected frequency of cells that would contain these interactions by chance, we generated random genomic regions that were size matched to each feature. We generated 1,000 random permutations of each feature. For each permutation, we computed the percentage of single cells containing any of these random bins. We get the normalized mean interaction value by first calculating an interaction matrix between all 1-Mb genomic bins, where the values in the interaction matrix were the percentage of cells containing an interaction between each pair of 1-Mb bins, and then calculating the mean value for pairs of regions in this interaction matrix representing centromere-proximal, speckle or nucleolar regions. For example, to determine the mean interaction value for cells containing an interaction between the pro-nuclear regions on chromosome 1 and chromosome 2, we calculated the mean value in this interaction matrix for chromosome 1 positions 3,000,000–13,000,000 with chromosome 2 positions 3,000,000–13,000,000.

Higher-order structures in scHi-C data. Ensemble and single-cell contact maps from scHi-C were plotted to visualize centromere, speckle and nucleolar interactions. The single-cell barcode from scHi-C that was referenced was ‘hpy 2.1-1CDES-1CDES_p10.H9-adj’.

DNA FISH comparison with ensemble scSPRITE analysis. For the FISH analysis, we focused on the same chromosomal loci pairs that were originally analyzed in SPRITE. These pairs include two control chromosomal pairs and four NOR chromosomal pairs. The chromosomal loci pairs are listed below:

- **Control 1:** chr3 (15–16 Mb) and chr15 (4–5 Mb)
- **Control 2:** chr5 (15–16 Mb) and chr19 (18–19 Mb)
- **NOR 1:** chr12 (6–7 Mb) and chr15 (4–5 Mb)
- **NOR 2:** chr15 (4–5 Mb) and chr18 (3–4 Mb)
- **NOR 3:** chr18 (3–4 Mb) and chr19 (18–19 Mb)

We first compared contact frequency values from the loci pairs listed above between ensemble scSPRITE and SPRITE to determine how well the two methods correlated with each other. For both ensemble scSPRITE and SPRITE, we generated 1-Mb resolution, genome-wide, pairwise contact frequency maps using clusters containing fewer than 10,000 reads per cluster. These contact maps were normalized using Hi-Corrector. Using both the ensemble scSPRITE and SPRITE normalized contact frequency maps, we then pulled out the contact frequency value from each of six loci pairs, plotted their values using a scatter plot and calculated the coefficient of determination (R²).

We generated 1-Mb resolution, genome-wide contact frequency matrices for each single cell using clusters containing fewer than 10,000 reads per cluster. For each chromosomal loci pair, a cell contained that loci pair interaction if there was at least one read in the bin containing both loci. To calculate the percentage of cells for each loci pair, we divided the number of cells containing the loci pair interaction by the total number of cells used in the analysis.

Calculation of percentage of reads coming from A/B compartments. To get the expected percentage of reads that fell into either the A or B compartments in our ensemble dataset, we used the data from the ensemble, genome-wide compartment switch analysis. We counted the number of 1-Mb bins that were classified as being in either A or B compartments genome-wide (except for chr3 and chrX). To get the expected percentage of reads in A or B compartments in our ensemble dataset, we then used the number 1-Mb bins in A or B compartments, respectively, by the total 1-Mb bins counted.

To get the percentage of reads in A or B compartments in single cells, we looked into the genome-wide reads (with the exception of chr3 and chrX) in each single cell file. From there, we sorted reads into A or B compartments depending on the data from the ensemble, genome-wide compartment switch analysis. Once sorted, we then divided the number of reads that fell into A or B compartments by the total number of reads counted for that cell to determine the percentage of reads in A or B compartments, respectively.

Contact maps of regions with heterogeneous structures. To identify regions of heterogeneity, we manually looked through a genome-wide heat map using the ensemble scSPRITE dataset to look for emerging TAD-like structures in between designated A/B compartments and TAD regions based on the previously identified A/B regions and TAD boundary regions, respectively. Once we identified a given chromosome, 40-kb weighted, single-cell contact maps were made for that specific chromosome, where the contact frequency values in each 40-kb bin are weighted by cluster size. In the 40-kb single-cell maps, the two 40-kb bins that made up the maximum interaction of the pseudo-TAD structure in the ensemble dataset were used to look for this same interaction in the single-cell dataset (further referred to as ‘bin A’ and ‘bin B’).

Long-range interactions. Detection of heterogeneity in long-range interactions. For the interactions studied in this paper (Phc1 at the Nanog locus and Lhx5 at the Tbx3 locus), we used the 40-kb bins containing the locations of the Phc1 enhancer and Nanog promoter and the locations of the Lhx5 gene and the Tbx3 gene as identified previously. In every single cell, we first identified cells containing a contact anywhere along bin A and bin B in that chromosome to ensure that coverage was accounted for. For Phc1 and Nanog, bins A and B are chr6 122,280,000–122,320,000 bp and chr6 122,640,000–122,680,000 bp, respectively. For Tbx3 and Lhx5, bins A and B are chr3 120,120,000–160,160,000 bp and chr3 120,880,000–120,920,000 bp, respectively. On average, we detect a contact in 1/3 of the total cell number, which we think is technical and due to non-sufficient coverage of every region per cell. Once the cells with coverage were identified, we identified and grouped cells in this set that contained or lacked the interaction at
the intersection of bin A and bin B. For the SE–promoter interaction at the Nanog locus, we identified 308 cells with read coverage, of which 159 cells contained the Nanog–Phc1 contact and 149 cells lacked the Nanog–Phc1 contact. For the SE–promoter interaction at the Tbx3 locus, we identified 301 cells with read coverage, of which 152 cells contained the Tbx3–Lhx5 contact and 149 cells lacked the Tbx3–Lhx5 contact.

For the cells with and without an interaction over the A/B compartment boundary (Fig. 4f), a similar approach was done as described above for grouping. We first identified cells based on coverage anywhere along bin A and bin B (that is, chrA 40,080,000–40,200,000 and chrA 40,920,000–41,000,000 bp). Unlike the promoter–enhancer examples where a known bin contains the promoter and enhancer loci, this information does not exist for contacts over a compartment boundary. Therefore, we provided a wider bp range to sort the cells into those two groups. Of the 379 cells identified with read coverage, 199 cells contained an interaction over the A/B compartment, and 170 cells lacked this interaction.

Virtual 4C analysis. To identify contacts with a specific locus, such as in the Nanog and Tbx3 examples, we first calculated a contact frequency matrix for all pairs of genomic bins at 40 kb resolution. Using the bins that were grouped into sets either containing or lacking interactions with a specific locus, we combined each cell’s individual contact frequency matrix to create an ensemble contact frequency matrix for each set. Each ensemble contact frequency matrix was normalized by Hi-Corrector44. To convert this contact matrix to a one-dimensional profile of contacts, we simply used the values in the row of the contact matrix corresponding to the locus of interest.

Significance and variance estimation. To determine the variance and significance of the observed contacts between these two groups, we performed a bootstrap method. Specifically, we generated random groups of cells by sampling with replacement from the initially defined groups. This approach allows us to estimate how much of the observed signal is dependent on individual cells in the population and how stable these estimates are across cells in the group. We generated 1,000 random bootstrap groups for each of the two groups and computed the average and s.d. across these permutations. To define the significance of differences between these two groups, we computed a P value using the unpaired two-sided t-test with Welch’s correction between the bootstrap values in group A versus group B.

Comparison of cells with and without SE–promoter contact. We compared the number of reads and contacts from the cells containing and lacking the SE–promoter contact from the Nanog and Tbx3 examples to determine if there was any bias that contributed to differences in their respective virtual 4C plots. For the Nanog–Phc1 example, we focused our analysis on the cells that contained or lacked the Nanog–Phc1 contact, as described previously. For each cell, we went through every cluster and calculated the number of genome-wide reads and contacts from each cluster. We then summed the number of reads and contacts from all the clusters in each cell and then repeated this process for all the cells in the two groups. We then used the Kolmogorov–Smirnov test to calculate statistical significance between the two groups. This same analysis was repeated for the Tbx3–Lhx5 example.

Chromatin immunoprecipitation with sequencing data. We downloaded the call sets from the ENCODE portal (https://www.encodeproject.org/) with the following identifiers: H3K27ac ENCSR000CDE, H3K4me3 ENCSR000CBG and H3K27me3 ENCSR000CFN.

Cell cycle analysis. We computationally sorted the cells into M, G1, G2 or S phases of cell cycle based on the parameters described previously46. After categorizing the cells by phase, we calculated the percentage of cells in each corresponding cell cycle phase in the sets that contained or lacked a particular interaction.

For the SE–promoter interaction at the Nanog locus, 152 of the 159 cells (95.6%) containing the Nanog–Phc1 contact and 149 of the 149 cells (97.3%) lacking the Nanog–Phc1 contact were sorted into cell cycle phases. For the SE–promoter interaction at the Tbx3 locus, 146 of the 152 cells (96.1%) containing the Tbx3–Lhx5 contact and 148 of the 149 cells (99.3%) lacking the Tbx3–Lhx5 contact were sorted into cell cycle phases. For the chrA A/B heterogeneity example, 195 of the 199 cells (98.0%) containing the interaction of the A/B compartment boundary and 166 of the 170 cells (97.6%) lacking this interaction were sorted into cell cycle phases. The other cells were identified as ‘Unknown’ and were not included in the cell cycle plot.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The datasets (Figs. 1–5 and Extended Data Figs. 1–5) generated and analyzed in the current study are available in the Gene Expression Omnibus repository under accession number GSE145353 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE145353).

Code availability

scSPRITE software is available at https://github.com/caltech-bioinformatics-resource-center/Guttman_Ismagilov_Labs.

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Author contributions

M.V.A. conducted the experiments to develop and validate the method, conceptualized and performed the analyses and wrote the manuscript. J.W.J. contributed to and supervised the experiments to develop and validate the method, conceptualized and performed the analyses and wrote the manuscript. N.O. conceptualized and performed analysis to validate the method, developed the pipeline for the workflow of scSPRITE sequencing data and contributed to writing the manuscript. M.S.C. contributed to the experiments to develop the method. C.A.L. developed a pipeline to sort cells by cell-specific barcodes. S.A.Q. contributed to the experiments to validate and develop the method. D.A.S. contributed to conceptualize scSPRITE and to the experiments to develop the method. R.F.I. conceptualized scSPRITE and supervised the experiments and the analysis to develop the method. M.G. conceptualized scSPRITE, supervised the experiments and the analysis to validate the method and wrote the manuscript. For detailed author contributions, please see Supplementary Note 5.

Competing interests

This paper is the subject of a patent application filed by Caltech. R.F.I. has a financial interest in Talis Biomedical Corp. S.A.Q. and M.G. are inventors on a patent owned by Caltech on SPRITE. The remaining authors declare no competing financial interests.

Additional information

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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | scSPRITE generate single cell maps with high genomic coverage.  

**a.** Quantification of cell aggregation. Top: number of cells in clumps pre- and post-filtration (singlets, doublets, triplets, etc). Bottom: microscope images (10x) of cells pre- and post-filtration step, scale bar 100 µm.

**b.** Validation of in-nuclei barcoding step of the protocol on mixed cell population (human-mouse cells): no mixing (top middle and top right), mixing before crosslinking (bottom left), mixing after crosslinking (bottom middle), and mixing after in-nuclei restriction digest (bottom right).

**c.** Schematic of the computational analysis pipeline for processing scSPRITE data.

**d.** Theoretical number of contacts measured by SPRITE-derived methods and Hi-C-derived methods over increasing numbers of DNA molecules per complex.

**e.** Maximum number of pairwise interactions that can be obtained from proximity ligation (Hi-C-derived methods) and complex barcoding (SPRITE-derived methods).

**f.** Genome-wide coverage for the filtered 1,000 cells: the median (black triangular points) and median absolute deviation (MAD) (green circular points) values were calculated per cell using the number of reads per 1Mb bin genome-wide (chr1-19).

**g.** Genomic coverage of 20 random cell barcodes; 1Mb bin per chromosome.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Known chromosomal structures can be measured genome-wide in hundreds of single mESCs by scSPRITE.

a. Additional single cell examples of chromosome territory structure between chr1 and chr2; plotted as number of DNA clusters at 1 Mb resolution. Box plot represents normalized detection scores between chr1 and chr2, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells).

b. Chromosome territory scores across 1000 cells (clustered based on similarity pattern). Columns represent chromosome territory detection scores for all pairs of chromosomes with the reference chromosome. Arrows represent chromosome territory scores between chr1 and chr2, which were analyzed in this paper.

c. Quantification of chromosome territory scores with respect to each chromosome. Boxplots show the range of chromosome territory scores, the average score (black line), and individual pairs of chromosome territory scores (grey dots).

d. Box plot represents average chromosome territory detection scores from all genome-wide (chr1-19) chromosome pairs, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells).

e. Additional single cell examples of A/B compartments detected within 0-55Mb in chr2; plotted number of DNA clusters at 1 Mb resolution (right). Box plot represents normalized detection scores between 0-55Mb in chr2, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells).

f. Representation of compartment switching scores across 1,000 cells (clustered based on score similarity pattern). Columns represent the strength of compartment switching detection scores for compartments that switched from “B-to-A-to-B” or “A-to-B-to-A” genome-wide (chr1-19). Arrows represent compartment switching scores for chr2 1-55 Mb, chr8 22-37 Mb, chr10 58-70 Mb, and chr17 8-45 Mb, all of which were analyzed in this paper.

g. Additional single cell examples of compartment switching from region 1, region 2, and region 3 (right). For each region’s box plot: whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells).

h. Expected (right) and observed (left) coverage of reads in the A and B compartment.
Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | Higher-order structures are identified genome-wide in hundreds of single mESC by scSPRITE method. a. Additional single cell examples of nucleolar interactions detected between chr18 and chr19; plotted number of DNA clusters at 1Mb resolution; detection scores below contact map (right). Box plot represents normalized detection scores between chr18 and chr19, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). b. Nucleolar interaction between chr12 and chr19; detection scores for 1000 cells (middle). Box plot where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). Representation of structures with max score (+1) and min. score (-1) (left) and ensemble scSPRITE heatmap (middle); contact map at 1Mb resolution. Single cell examples (right); plotted number of DNA clusters at 1Mb resolution. c. Relative correlation of the percent of cells from scSPRITE vs DNA-FISH containing inter-chromosomal interactions at specified 1Mb regions targeted by DNA-FISH probes. Control chromosomes (grey points) and nucleolar associating chromosomes (black dots) are plotted. d. Relative correlation of the contact frequency from scSPRITE vs the contact frequency from SPRITE containing inter-chromosomal interactions targeted by DNA-FISH probes. Control chromosomes (grey points) and nucleolar associating chromosomes (black dots) are plotted. e. Frequency of cells containing inter-chromosomal nucleolar contacts (normalized to number of reads per region) for each pair of nucleolar associating chromosomes. f. Single cell examples of speckle interaction detected between chr2 and chr5; plotted number of DNA clusters at 1Mb resolution. Box plot represents normalized detection scores between chr2 and chr5, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). g. Additional single cell examples of speckle interactions detected between chr2 and chr4; plotted number of DNA clusters at 1Mb resolution. Box plot represents normalized detection scores between chr2 and chr4, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). h. Frequency of cells containing inter-chromosomal speckle contacts (normalized to number of reads per region) for each pair of speckle associating chromosomes. i. Additional single cell examples of centromere-proximal interactions detected between chr1 and chr11; plotted number of DNA clusters at 1Mb resolution. Box plot represents normalized detection scores between chr1 and chr11, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). j. Single cell examples of chr4 and chr11 centromere-proximal regions interacting together; plotted number of DNA clusters at 1Mb resolution. Box plot represents normalized detection scores between chr4 and chr11, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). k. Frequency of cells containing inter-chromosomal centromeric contacts (normalized to number of reads per region) for each pair of chromosomes. l. Higher-order structures representation from scHi-C data - centromere-proximal interactions, speckle interactions, and nucleolar interactions; Pairwise contact map from ensemble 1,000 cells (left), pairwise contact map from their best single cell (right).
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | TADs are heterogeneous units present in the genomes of individual mESCs. a. Genome-wide correlation of insulation scores between ensemble scSPRITE and Hi-C' from mouse ES cells at 40 kb resolution. b. Insulation score profile of ensemble scSPRITE (red) and Hi-C' (blue) at 40 kb resolution at chr1 65-95 Mb. c. Additional single cell examples of TAD-like structures between 124.8-126.7Mb of chr4; plotted number of DNA clusters at 40 kb resolution; detection scores below contact map. Box plot represents normalized detection scores between 124.8-126.7Mb of chr4, where whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median, red dots represent single cell examples (n = 1000 cells). d. TAD detection scores across 1,000 cells (clustered based on score similarity pattern) in chr2 (left) and chr18 (right). Columns represent the strength of TAD detection scores for all TADs detected across chr2 or chr18, respectively, in ensemble scSPRITE. e. TAD detection scores across 1,000 cells between 38.5-48.56 Mb of chr4. Each line represents the strength of TAD detection scores in this given region from a single cell. Cells are either in Group 1 or 2 in Fig. 4f or not used. f. Ensemble heatmap from all 1000 cells between 39.4-41.4Mb of chr4 representing strong TADs detected in bulk (blue lines), and weak emerging TADs (green line) over the A/B boundary. g. Fraction of cells in each cell cycle phase from the set of single cells containing (left) or lacking (right) the contact between the boundary region (Fig. 4f). h. Difference contact map across a control region 84.8-88.4 Mb of chr4 made by subtracting the normalized contacts from cells in Group II from Group I (Fig. 4f). Insulation scores for cells in Group I (dark grey) and Group II (light grey) are plotted.
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | Structural heterogeneity in long-range interactions is revealed by scSPRITE. **a.** Ensemble heatmaps across 122.2-122.8 Mb region in chr6 representing cells containing (top) or lacking (bottom) the contact between the Nanog locus and the -300 Kb SE. Blue square shows the contact. **b.** Number of genome-wide reads (left) and number of genome-wide contacts (right) for groups of cells with and without the Nanog-SE interaction. For each box plot, whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median (with = 159 cells, without = 149 cells). No statistical significance between the two groups were seen based on the Kolmogorov–Smirnov two-sided test. **c.** Fraction of cells in each cell cycle phase from the set of single cells containing (left) or lacking (right) the contact between the Nanog locus and the SE 300kb upstream of Nanog. **d.** Heatmaps between 119.24-121.28Mb in chr5 of pooled cells either containing (top) or lacking (bottom) the contact between the Tbx3 locus and Lhx5. Blue square shows the contact. **e.** Number of genome-wide reads (left) and number of genome-wide contacts (right) for groups of cells with and without the Tbx3-Lhx5 interaction. For each box plot, whiskers represent the 10th and 90th percentiles, box limits represent the 25th and 75th percentiles, black line represents the median (with = 152 cells, without = 149 cells). No statistical significance between the two groups were seen based on the Kolmogorov-Smirnov two-sided test. **f.** Fraction of cells in each cell cycle phase from the set of single cells containing (left) or lacking (right) the contact between the Tbx3 locus and the Lhx5.
Reporting Summary

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Statistics

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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection  No software was used

Data analysis  cworld [https://github.com/dekkerlab/cworld-dekker]; Snakemake pipeline for sequencing data [https://github.com/caltech-bioinformatics-resource-center/Guttman_Ismagilov_Labs]

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- Accession codes, unique identifiers, or web links for publicly available datasets
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GEO Accession # GSE154353 [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE154353]. Secure token for viewing the GEO Accession is “gqknzyawozdyh”. Github link: https://github.com/caltech-bioinformatics-resource-center/Guttman_Ismagilov_Labs.
Field-specific reporting

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☒ Life sciences  ☐ Behavioural & social sciences  ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/n-refining-summary-flat.pdf

Life sciences study design

All studies must disclose these points even when the disclosure is negative.

Sample size: Around 1500 nuclei were originally loaded for in-nuclei cell-specific barcoding and subsequent sequencing. During in-silico analysis, to ensure we selected single cells for downstream analysis, the top 3.4% percent of cells, as determined from the results of the human-mouse mixing experiment, were removed. From the remaining files, to ensure that the analyses containing the highest coverage per cell were analyzed, we then selected the top 1000 cell barcode files containing the most number of clusters per cell for downstream single-cell analysis.

Data exclusions: All data, including any data removed during in-silico analyses, are being made available.

Replication: Because single cell measurements are done, each cell is a replicate.

Randomization: None - No treatment groups were present such that randomization is necessary.

Blinding: None - No treatment groups were present such that blinding is necessary.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

n/a Involved in the study
☒ Antibodies
☒ Eukaryotic cell lines
☒ Palaeontology and archaeology
☒ Animals and other organisms
☒ Human research participants
☒ Clinical data
☒ Dual use research of concern

Methods

n/a Involved in the study
☒ ChiP-seq
☒ Flow cytometry
☒ MRI-based neuroimaging

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s): The male bsp5 mouse embryonic stem cell line was provided by K. Plath. HEK293T, a female human embryonic kidney cell line transformed with the SV40 large T antigen was obtained from ATCC (#CRL-1573).

Authentication: Not applicable for cell line authentication

Mycoplasma contamination: Cells were routinely tested for mycoplasma using a kit from Millipore-Sigma (MP0025-1KT)

Commonly misidentified lines: No commonly misidentified cell lines

(See ICLAC register)