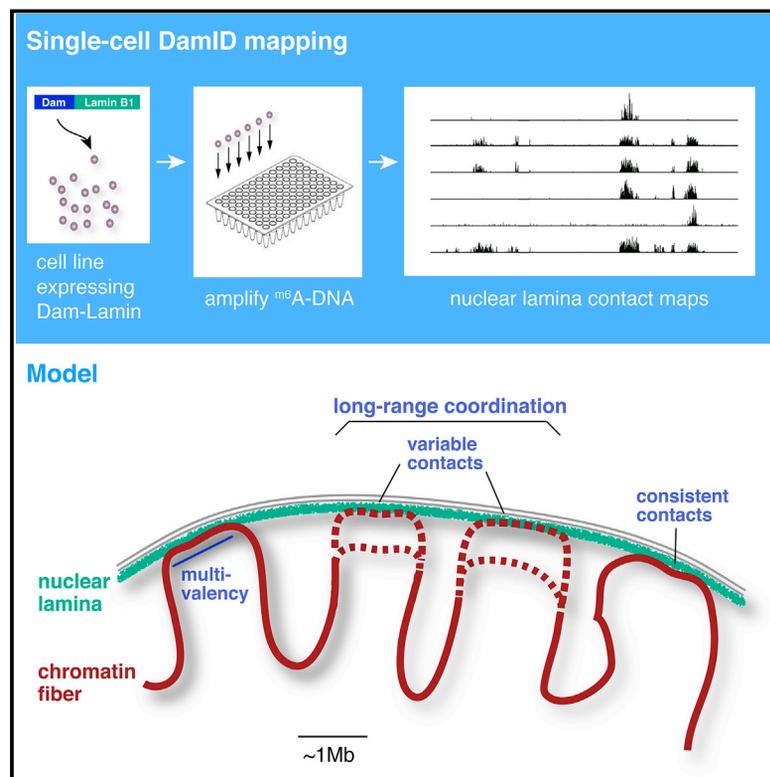


# Genome-wide Maps of Nuclear Lamina Interactions in Single Human Cells

## Graphical Abstract



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## In Brief

A modified DamID method enables the mapping of genome-wide nuclear lamina interactions in single human cells, providing insight into the cell-to-cell variation in the interphase chromosome architecture and suggesting extensive intra-chromosomal coordination of nuclear lamina contacts.

## Highlights

- Modified DamID method maps genome-nuclear lamina interactions in single human cells
- Nuclear lamina contact frequency is locus specific
- Stable contact sites are extremely gene poor, suggesting a structural role
- Single-cell maps point to multivalent interactions and long-range coordination

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# Genome-wide Maps of Nuclear Lamina Interactions in Single Human Cells

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## SUMMARY

Mammalian interphase chromosomes interact with the nuclear lamina (NL) through hundreds of large lamina-associated domains (LADs). We report a method to map NL contacts genome-wide in single human cells. Analysis of nearly 400 maps reveals a core architecture consisting of gene-poor LADs that contact the NL with high cell-to-cell consistency, interspersed by LADs with more variable NL interactions. The variable contacts tend to be cell-type specific and are more sensitive to changes in genome ploidy than the consistent contacts. Single-cell maps indicate that NL contacts involve multivalent interactions over hundreds of kilobases. Moreover, we observe extensive intra-chromosomal coordination of NL contacts, even over tens of megabases. Such coordinated loci exhibit preferential interactions as detected by Hi-C. Finally, the consistency of NL contacts is inversely linked to gene activity in single cells and correlates positively with the heterochromatic histone modification H3K9me3. These results highlight fundamental principles of single-cell chromatin organization.

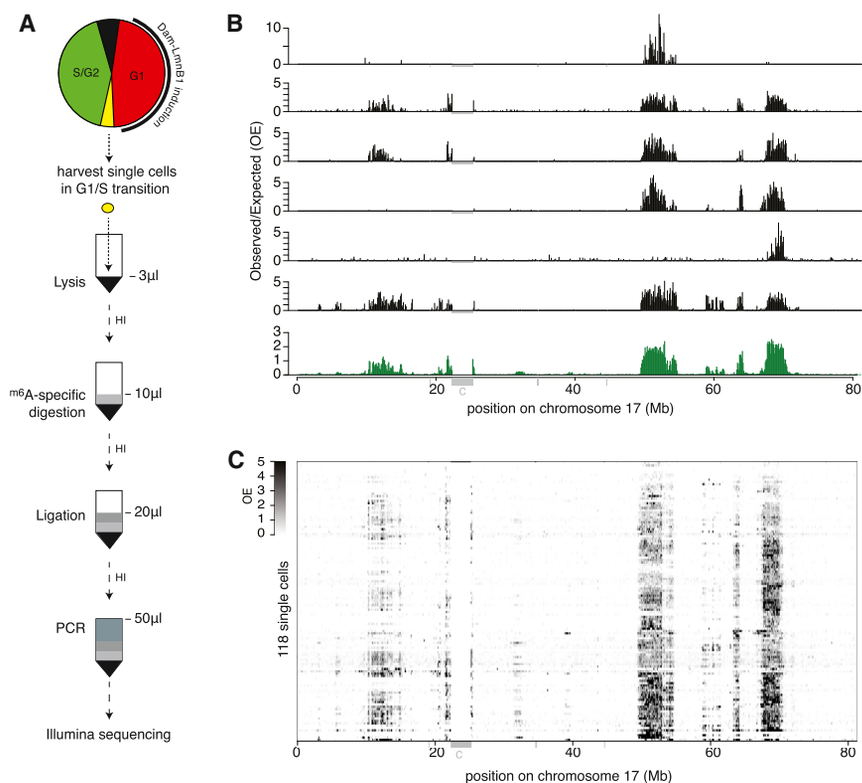
## INTRODUCTION

An important unresolved question in eukaryotic genome biology is how chromosomes are spatially organized inside interphase nuclei. Current evidence suggests that this organization is driven by probabilistic principles (Bickmore, 2013; Cavalli and Misteli,

2013; Gibcus and Dekker, 2013). Systematic fluorescence in situ hybridization (FISH) experiments have revealed that in a homogeneous cell population the nuclear positions of chromosomes are variable with respect to each other and relative to the periphery (Bolzer et al., 2005). However, this positioning is not entirely random; for example, in human lymphoid cells, chromosome 18 (chr18) tends to be located near the periphery, while chr19 shows a preference for the nuclear interior (Croft et al., 1999; Cremer et al., 2001).

At a smaller scale, certain individual genomic loci visualized by FISH also exhibit preferences for specific nuclear landmarks, such as the nuclear envelope (Marshall et al., 1996; Kosak et al., 2002) and nucleoli (Manuelidis and Borden, 1988; Ochs and Press, 1992), but usually with some degree of random variation. This variability is directly illustrated by in vivo tagging experiments in which loci contacting the nuclear lamina (NL) were tracked over mitosis in a clonal human cell line (Kind et al., 2013). This demonstrated that a sizeable subset of loci that were associated with the NL in mother cells relocated to the nuclear interior in daughter cells, indicating that, at least to some degree, genome contacts with the NL are intrinsically variable.

Complementary to these single-cell microscopy approaches are genome-wide mapping techniques that query the chromosome organization in large pools of cells (van Steensel and Dekker, 2010). For example, the 4C, 5C, and Hi-C technologies generate maps of the pair-wise spatial proximity of genomic loci (de Wit and de Laat, 2012; Dekker et al., 2013). Such maps have revealed global patterns that indicate that mammalian interphase chromosomes are partitioned into domains that are roughly 200 kb–2 Mb in size (Lieberman-Aiden et al., 2009; Dixon et al., 2012; Nora et al., 2012; Rao et al., 2014). Computational models of chromosome polymer folding fitted to 5C and Hi-C data generally suggest that interphase chromosomes adopt multiple configurations that vary from cell to cell



**Figure 1. Single-Cell DamID Mapping of Genome-NL Interactions**

(A) Schematic representation of the single-cell DamID procedure. HI, heat inactivation step. (B) NL contact maps for chr17 in six individual cells (black) and the average profile of 118 single cells (bottom track, green). OE, observed over expected read count ratio in contiguous 100-kb segments. (C) Grayscale representation of OE values for 118 single cells on chromosome 17. The gray bars underneath the axes in (B) and (C) mark unmappable regions; c, centromeric region. See also Figure S1.

ing, provide insights into the nature of LAD-NL interactions and uncover principles of cell-to-cell variation in chromosome architecture.

## RESULTS

### Single-Cell DamID Methodology

As a model system to develop single-cell DamID, we chose the human myeloid leukemia cell line KBM7, which is haploid for all chromosomes, except for chr8 and a small part of chr15 (Kotecki et al., 1999; Bürckstümmer et al., 2013). Even though this haploid state is unusual for human

(Lieberman-Aiden et al., 2009; Baù et al., 2011; Kalhor et al., 2012; Giorgetti et al., 2014; Nagano et al., 2013).

Another genome-wide approach to studying chromosome architecture is the mapping of interactions with the NL, mostly by means of the DamID technology (Guelen et al., 2008; Peric-Hupkes et al., 2010; Meuleman et al., 2013). The NL provides a very large surface area for potential contacts with the genome. Indeed, DamID studies have estimated that as much as ~35% of the mammalian genome can interact with the NL in any tested cell type, although it has remained unclear how much of the genome contacts the NL in a single cell. Genome-NL interactions occur through about 1,100–1,400 discrete lamina-associated domains (LADs), which have a median size of ~0.5 Mb and are scattered across all chromosomes. Most genes in LADs are expressed at very low levels, and results of various tethering experiments (Finlan et al., 2008; Reddy et al., 2008; Kind et al., 2013; Therizols et al., 2014) point to a reciprocal relationship between gene positioning at the NL and a repressive chromatin state.

The large size of LADs and their prevalence throughout the genome strongly suggest that LAD-NL interactions play an important role in interphase chromosome architecture. Insights into the single-cell behavior of these interactions will thus enhance our fundamental understanding of chromosome organization. Here, we report a modified version of the DamID technology that is sensitive enough to generate genome-wide maps of NL contacts in single human cells at a resolution of ~100 kb, which is well below the median size of LADs. We generated a total of 395 of these single-cell maps. These maps, complemented by Hi-C analysis and super-resolution microscopy imag-

somatic cells, it facilitates the interpretation of single-cell genome-wide maps because there is no need to discriminate the homologous chromosomes that are present in diploid cells.

We developed a DamID protocol for single cells as summarized in Figure 1A. We created a KBM7 clone (#14) that expresses an inducible fusion protein consisting of DNA adenine methyltransferase (Dam) and Lamin B1 (LmnB1) (Kind et al., 2013), as well as the Fucci two-color fluorescent reporter system to monitor the cell-cycle stage (Sakaue-Sawano et al., 2008). We then induced Dam-LmnB1 protein expression (Figures S1A and S1B), and 15 hr later we collected single cells at the onset of the S phase by fluorescence-activated cell sorting (FACS) (Figure S1C). This design ensured that the harvested cells had expressed Dam-LmnB1 for most of their recent G1 phase (which lasts on average ~14.9 hr, an estimate based on doubling time and FACS data), providing sufficient time for the accumulation of adenine methylation on the genomic loci that contact the NL. Importantly, LmnB1 was strictly confined to the nuclear periphery before and after Dam-LmnB1 induction (Figure S1D).

The single FACS-sorted cells were captured directly in a small volume of lysis buffer in a 96-well plate. Subsequent sample processing consisted of only a few steps (Figure 1A): digestion with DpnI, which is highly specific for Dam-methylated GATC sequences, followed by adaptor ligation (Figure S1E) and a total of 26 cycles of PCR amplification. A key difference with the conventional DamID protocol is that these steps are all done in the same well by sequential addition of reagents, without intermediate purification or concentration steps that could lead to loss of DNA. Gel electrophoresis showed that approximately half of the

wells yielded a clear smear of amplified DNA (Figure S1F). These PCR products were prepared for multiplexed Illumina sequencing by ligation of indexed adaptors, subsequently pooled, and sequenced.

### Single-Cell NL Interaction Maps

In total, we generated 118 single-cell DamID maps from KBM7 clone #14. After applying quality filters (Figure S1G), we obtained a median of  $5 \times 10^5$  reads per cell that could be mapped to a unique genomic location (Figure S1H). On average, 92% of the mapped locations started with a GATC motif, which is the recognition sequence of both Dam and DpnI, indicating that the detection is highly specific. For each cell, we binned these reads in 100-kb contiguous genomic segments, and then we calculated for each segment an observed over expected (OE) score based on the number of recovered unique reads, the theoretical maximum number of mappable unique reads in the segment, and the total genome-wide read count obtained from each cell. OE scores  $>1$  indicate more Dam-LmnB1 methylation than may be expected by random chance.

Essentially, all cells showed a striking domain pattern of OE scores along most chromosomes, as illustrated for chr17 in Figures 1B and 1C. This pattern is reminiscent of the LAD profiles previously published for populations of cells. However, clear differences can be observed between individual cells (Figure 1B), often with entire megabase-sized domains missing, which is suggestive of cell-to-cell variation in NL contacts.

We note that the chosen segment size of 100 kb is a compromise between resolution and noise. Because the median human LAD size is approximately 0.5 Mb (Guelen et al., 2008), segments of 100 kb are expected to capture most of the LAD organization. Indeed, the OE scores in adjacent segments within each of the 118 single cell samples have a Pearson correlation coefficient of  $0.70 \pm 0.06$  (mean  $\pm$  SD), indicating that neighboring segments report NL interactions in a partially redundant fashion with acceptable noise levels. For reference, at 100 kb resolution our previously published Dam-LmnB1 profiles from pools of human Tig3, hESCs, and HT1080 cells (Guelen et al., 2008; Kind et al., 2013; Meuleman et al., 2013) show a correlation of 0.88–0.90 between neighboring segments. The single-cell correlation of OE scores between adjacent segments is not related to the number of reads per cell (Pearson's correlation coefficient of 0.06,  $p = 0.51$ ), indicating that the latter does not impose a limit to the quality of our data at 100 kb resolution.

### Validation of Single-Cell Maps

To further gauge the overall quality of these data, we first reconstructed a population profile by averaging the maps of the 118 single cells and then compared it to a conventional microarray-based DamID profile generated from a large pool of KBM7 cells (Figure 2A). The highly similar domain patterns and an overall Spearman rank correlation coefficient of 0.90 demonstrate that the new protocol captures the same regions of interaction as the previous well-validated protocol.

However, in conventional DamID, a Dam-only control is typically included to normalize for chromatin accessibility and other potential biases (Vogel et al., 2007; Guelen et al., 2008). This normalization is not possible in single-cell DamID because the

Dam-LmnB1 and Dam-only profiles cannot be obtained from the same cell. Nevertheless, we established a Dam-only and Fucci-expressing KBM7 clone and mapped the adenine methylation patterns in 26 single cells. The resulting patterns are very different from the Dam-LmnB1 profiles (Figures 2B and 2C). In general, regions that have no detectable Dam-LmnB1 signal show clear Dam-only signals; hence, these regions are not intrinsically undetectable. Conversely, regions with very high Dam-only signals generally do not show a Dam-LmnB1 signal, indicating that there is no strong bias for accessibility in our Dam-LmnB1 maps. We conclude that leaving out the Dam-only normalization is acceptable for single-cell Dam-LmnB1 profiling.

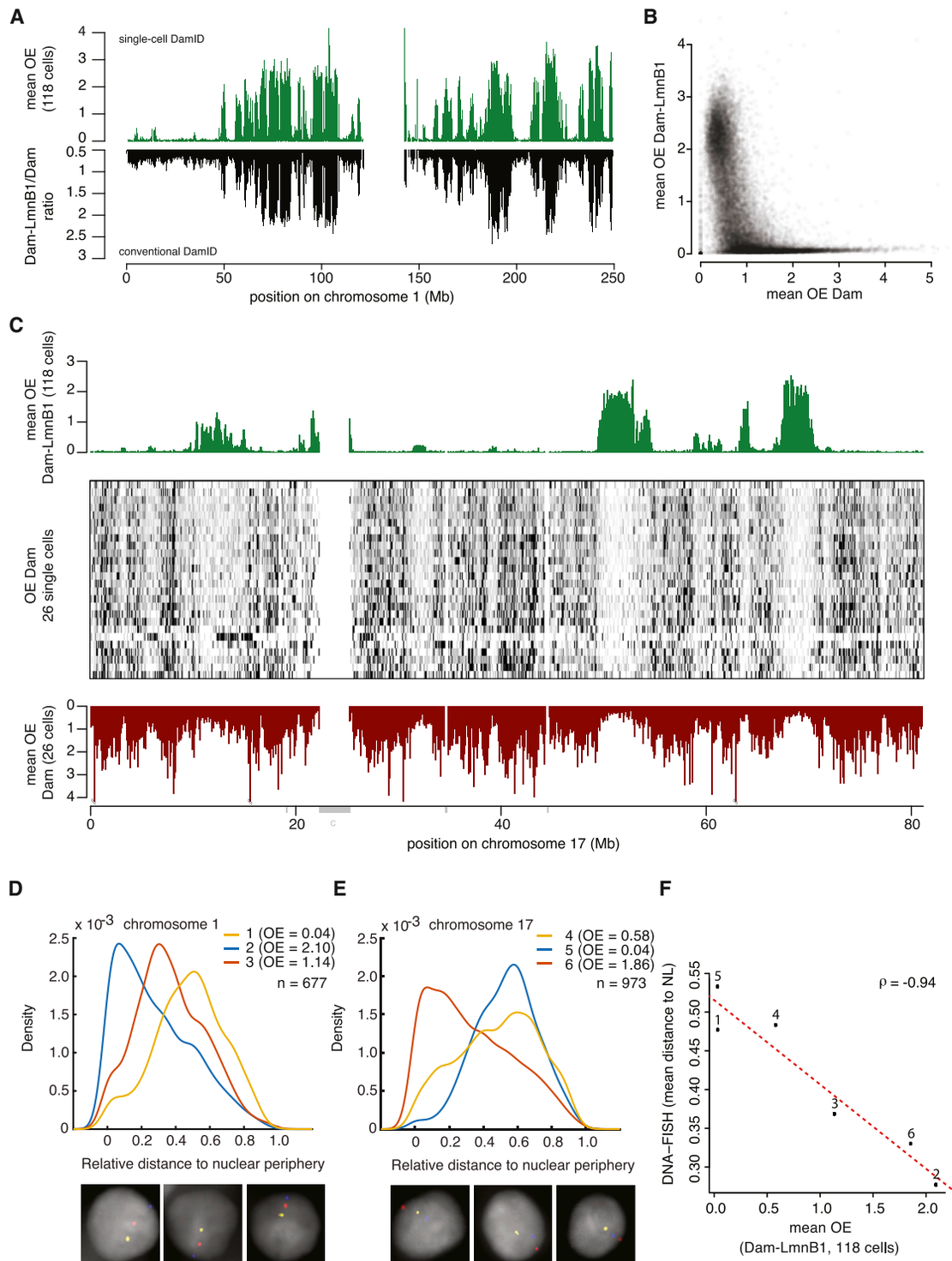
Next, we performed multi-color fluorescent in situ hybridization (FISH) with probes for six genomic loci covering a broad range of average OE scores (Figures 2D–2F). Analysis of hundreds of nuclei revealed a good correspondence between the average distance to the periphery according to FISH and the average OE scores (Spearman's  $\rho = 0.94$ ), confirming that our single-cell Dam-LmnB1 profiles provide a view of the spatial organization of the genome relative to the NL. Together these data indicate that single-cell DamID using Dam-LmnB1 generates NL interaction profiles with low noise levels, suitable resolution, acceptable bias, and with good correspondence to localization by FISH.

### Cell-to-Cell Variability and Consistency of Genome-NL Associations

Visual inspection of the collection of single-cell maps suggested that some regions interact more frequently with the NL than others (Figures 1B and 1C). In order to analyze this systematically, we first converted the OE scores for each cell to a binary NL contact map. For this, we used an OE score cutoff of 1, motivated by the bimodal distribution of OE scores (Figure S2A) that suggests that loci are either in a “contact” or “no-contact” state. We then calculated for each 100-kb segment the NL contact frequency (CF), defined as the proportion of cells in which this segment contacted the NL. This data processing does not lead to a substantial loss of information content, because the Spearman rank correlation coefficient of the average radial position of the six FISH probes with CF is 0.90 (as compared to 0.94 for OE values, see above), and CF values correlate strongly with the average OE scores (Figure S2B). CF values are also highly robust with respect to sequencing read depth, because subsampling of the single-cell data to the one-but-lowest read count ( $1.04 \times 10^5$  reads; an average 3.1-fold downsampling) does not affect CF values (Figure S2C).

Strikingly, CFs vary widely across the genome (Figures 3A and 3B). About 23% of the 100-kb segments show no detectable contact with the NL in any of the 118 cells and thus are very stably located in the nuclear interior. Conversely,  $\sim 15\%$  of the segments have CFs  $>80\%$ , representing loci that are consistently located at the NL. About 34% of the segments have contact frequencies in the range of 20%–80% and thus show high cell-to-cell variability in their NL associations. The remaining loci (29%) only occasionally contact the NL ( $0 < CF < 20\%$ ).

These different classes of loci are scattered throughout the genome (Figure 3A), although the smaller chromosomes tend



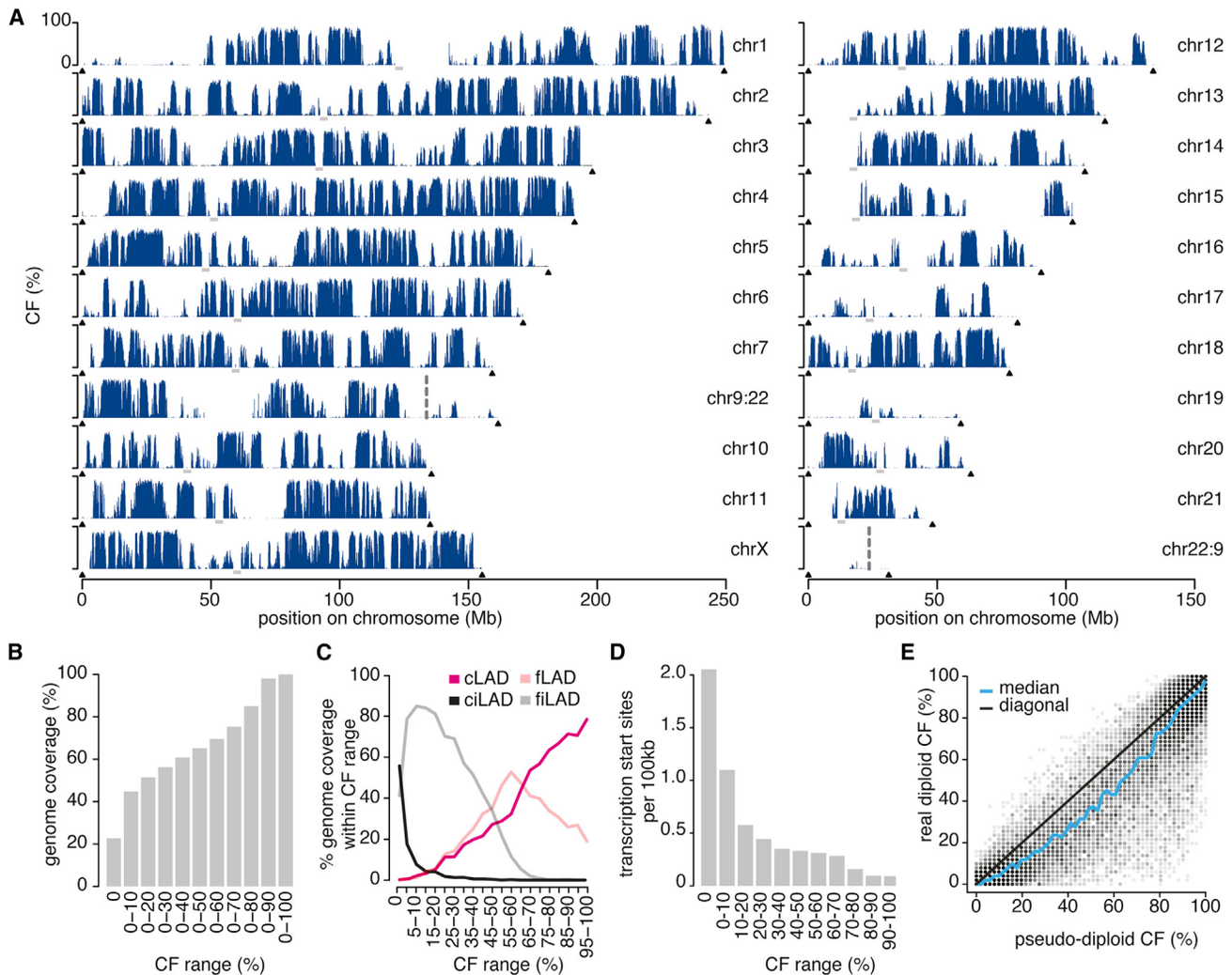
**Figure 2. Validation of Single-Cell DamID Maps**

(A) Comparison NL contact map for chr1 representing the average of 118 single-cell profiles (top profile) and a conventional DamID map generated with a population of  $\approx 1\text{--}2 \times 10^5$  cells (bottom profile). The genome-wide correlation between the two methods is 0.90 (Spearman's  $\rho$ .)

(B) Average OE score of 118 single-cell Dam-LmnB1 samples (y axis) versus the average OE score of 26 single-cell Dam-only samples (x axis).

(C) Comparison of average OE scores obtained with Dam-LmnB1 (top track, average of 118 cells, same as in Figure 1B) and Dam-only (bottom track, average of 26 cells). OE scores for the individual Dam-only cells are shown as grayscale-encoded rows in the center frame. Gray bars underneath the bottom axis mark unmappable regions; c, centromeric region.

(legend continued on next page)



**Figure 3. NL Contact Frequencies Are Linked to Developmental Dynamics, Gene Density, and Ploidy**

(A) Estimated contact frequency maps for all chromosomes in clone #14 cells. KBM7 cells carry a balanced translocation between chr9 and chr22 (Bürckstümmer et al., 2013); vertical dotted lines mark the junctions. Centromeric regions are indicated by gray bars; telomeres are marked by black triangles. chr8 is not shown because it is diploid.

(B) Cumulative histogram of genome-wide CF values.

(C) Distribution of genomic segments with indicated CFs over constitutive (c) and facultative (f) LADs and inter-LADs (iLADs).

(D) Average number of transcription start sites per 100-kb segment, plotted as a function of CF.

(E) Comparison of CFs in diploid cells and pseudo-diploid cells. The latter are simulated by combining equal numbers of sequence reads from pairs of haploid cells. See also Figure S2 and Table S1.

to have a lower density of stable NL contacts than larger chromosomes. An exception to this rule is chr18, which harbors many regions with high CFs. This contrasts in particular with chr19, which only exhibits a few contact sites and very low CFs. This matches previous chromosome painting studies that found chr18 to be preferentially located at the nuclear periphery and chr19 in the nuclear interior (Croft et al., 1999; Cremer et al.,

2001). An intriguing pattern is visible on chrX (Figure 3A, bottom left). The distal arms of this chromosome have many stable NL contacts, while the centromere-proximal ~40 Mb show only variable contacts.

In order to confirm these CF patterns, we used an independently derived KBM7 clone that also expresses Dam-LmnB1 and the Fucci system (clone #5.5) to generate a total of 168

(D–F) Multi-color 3D DNA FISH microscopy with probes for six genomic loci covering a broad range of average OE scores distributed on chr1 ( $n = 677$ ) (D) and chr17 ( $n = 973$ ) (E). Graphs depict the distributions of radial probe positions, with zero corresponding to the nuclear edge and one to the centroid. Three representative nuclei with three-color FISH signals are displayed below the graphs; DNA staining with DAPI is shown in gray. (F) Mean radial positions of the six probes versus the mean DamID OE scores. Numbers 1–6 correspond to probe numbers in (D) and (E). The dotted line shows linear regression fit.

single-cell maps. The genome-wide CF profile of clone #5.5 is highly similar to that of clone #14 (Pearson correlation,  $r = 0.97$ ; Figures S2D and S2E). However, the absolute CF values in clone #5.5 are systematically 1.3-fold lower than in clone #14, which we attribute to a somewhat lower activity of the Dam-LmnB1 protein in clone #5.5 (Figure S2F). Nevertheless, the relative CF differences between loci are highly consistent between the two clones. We cannot rule out that the CF values of clone #14 are underestimated, but the fact that the highest CF values of clone #14 are in the range of  $\sim 95\%$ , combined with the tight linearity with clone #5.5, suggests that any underestimate in clone #14 is minor.

### Stable and Variable NL Contacts Are Linked to Degrees of Developmental Plasticity

Previously, we reported that some LADs interact with the NL in a cell-type-specific (facultative) fashion, while other LADs do so in a cell-type invariant (constitutive) manner (Peric-Hupkes et al., 2010; Meuleman et al., 2013). We investigated whether this distinct behavior is linked to differences in CF. We used a collection of conventional microarray-based DamID maps of NL interactions in nine human cell lines of diverse origin (C.A.d.G. and B.v.S., unpublished data; see the Supplemental Experimental Procedures) to classify each 100-kb segment as constitutive LAD (cLAD), constitutive inter-LAD (ciLAD), facultative LAD (fLAD), or facultative inter-LAD (fiLAD). The latter are inter-LAD regions that are not associated with the NL in KBM7 cells, but do interact with the NL in at least one other cell type.

As expected, ciLADs have low CF values (Figure 3C), consistent with their definition as not associated with the NL. Likewise, cLADs tend to have high CF values. Thus, these constitutive regions are not only invariable between different cell types but tend to be consistently positioned relative to the NL within one cell type. In contrast, facultative LADs and iLADs have mostly intermediate CF values: fLADs generally have lower CF values than cLADs ( $p < 2.2 \times 10^{-16}$ , Wilcoxon test) and fiLADs have higher values than ciLADs ( $p < 2.2 \times 10^{-16}$ ). The partial overlap in CF distributions of fLADs and fiLADs indicates that the definitions of these LAD classes are not perfect, which may be due to differences in the data types (single-cell versus population-based DamID maps; sequencing versus microarray) and resolution. These results uncover a link between CF and the consistency of NL interactions between different cell types.

We previously reported that cLADs have a  $\sim 2$ -fold lower gene density than ciLADs (Meuleman et al., 2013). CF values are more dramatically linked to gene density: regions with  $CF > 80\%$  have a  $\sim 20$ -fold lower gene density than regions with no NL contacts (Figure 3D). The few genes that are located in such high-CF regions are enriched for gene ontology categories very divergent from myeloid cell functions, among them most prominently the olfactory receptor genes (Table S1), which are rarely expressed.

A picture thus emerges in which most cLADs are relatively consistently associated with the NL, providing a structural backbone to chromosomes that is largely invariant between individual cells and also between cell types. In contrast, regions with cell-type-specific NL interactions generally interact less consistently with the NL, contributing to cell-to-cell variability in the spatial organization of chromosomes.

### Ploidy of KBM7 Cells Primarily Affects Variable NL Contacts

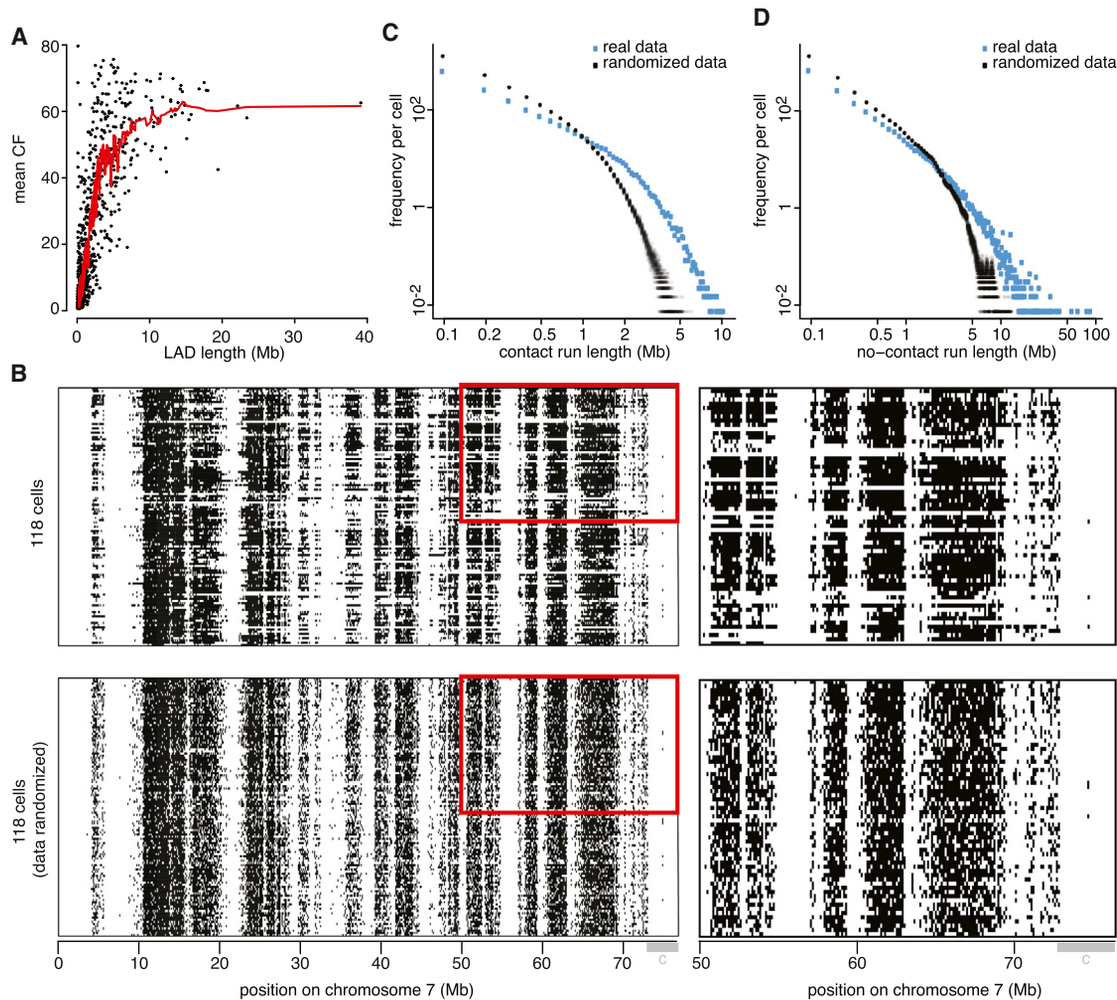
We considered the possibility that competition between LADs could explain why some LADs contact the NL in only a fraction of all cells: such LADs may have lower NL binding affinities and fail to compete with stronger LADs in some of the cells. We wondered whether this balance could be altered by changing the total amount of genomic DNA in the nucleus. To test this, we took advantage of the fact that KBM7 cells spontaneously form diploid cells at low frequency (Kotecki et al., 1999). Such diploid cells should be genetically identical to the haploid cells, except for their ploidy. We derived a clonal diploid line from clone #14, with normal cell-cycle behavior (Figure S2G), and generated a total of 51 single-cell Dam-LmnB1 contact maps.

Comparison of the single-cell maps from haploid and diploid cells is not straightforward, because our current DamID method cannot determine whether the sequence reads from diploid cells are derived from one homolog or both. We therefore constructed “pseudo-diploid” reference maps by combining equal numbers of sequence reads from pairs of single haploid cells and then merged the reads and processed the data as above, as if they were derived from a single cell. Hence, these pseudo-diploid maps are simulated maps in which two homologs of each chromosome are present, but there cannot be any biological effect of increased ploidy (because the homologs were in different cells), while technical skews due to the inability to discriminate the two homologs should be identical to those in diploid cells.

Although the overall pattern of NL contacts was similar between pseudo-diploid and diploid cells, many loci showed decreased CFs in diploid cells compared to haploid cells (Figure 3E), while increased CFs were rare. Strikingly, these changes occurred preferentially in genomic segments that have intermediate CFs in pseudo-diploid (and thus haploid) cells. This result indicates that diploidization leads to preferential loss of NL interactions of LADs that contact the NL less robustly in haploid cells. We obtained similar results with 32 single cells from a diploid clonal line derived from clone #5.5 (Figure S2H). The loss of NL contacts in diploid cells suggests increased competition for NL contacts. This may be caused by a reduced surface-to-volume ratio and an increase in the average distance of loci to the NL, both of which are expected to accompany the increase in nuclear volume due to the doubled DNA content. However, other mechanisms cannot be ruled out.

### Single-Cell Maps Point to a Multivalent Mechanism of LAD-NL Interactions

We wondered which biophysical principle could explain the differences in apparent NL affinity between genomic loci. The domain pattern of NL contacts suggests that these interactions are not mediated by focal attachments, but rather by multivalent interactions within each LAD. Because interactions with higher valency typically have a higher avidity, we predicted that long LADs interact more stably with the NL than short LADs. To analyze this, we defined LADs as continuous stretches of 100-kb segments with  $CF > 1\%$  across the 118 cells. This yielded a total of 1,358 LADs. Strikingly, the mean CF within each LAD shows a clear positive correlation with LAD length (Spearman's  $\rho = 0.81$ ;  $p < 2.2 \times 10^{-16}$ ), reaching a plateau for sizes larger



**Figure 4. Evidence for Multivalent NL Interactions**

(A) Correlation between the mean CF and the length of LADs. Here, LADs are defined as continuous regions in which all 100-kb segments have CF >1% across all 118 clone #14 cells. The curve shows running mean with bin size 15.

(B) Binarized NL contact map of the p arm of chr7 in 118 cells (top), and the same data after a random shuffling procedure that keeps CFs and the number of contacts per cell the same, to simulate the complete absence of coordination between neighboring segments (bottom). The gray bar marked “c” indicates the centromeric region. Right-hand panels are magnified views of the regions outlined by red boxes in the left-hand panels.

(C) Distribution of genome-wide NL contact run lengths in 118 single-cell datasets (blue) compared to 100 sets of randomized data (gray).

(D) Genome-wide length distribution of no-contact runs in real (blue) and 100 sets of randomized (gray) data. No-contact runs present in all 118 cells (i.e., regions that never contact the NL) were excluded in this analysis.

See also Figure S3.

than ~6–8 Mb (Figure 4A). This supports a model of multivalent local genome-NL interactions.

This multivalency model predicts that there are continuous stretches of contact in individual LADs in single cells. Indeed, such runs of contact could be frequently observed (black horizontal streaks in Figure 4B, top). For comparison, we randomly shuffled the contact data to simulate a “random button” model, in which each segment maintains exactly the same CF as in the real data, but contacts the NL independently of its neighboring segments (Figure 4B, bottom). This pattern has a much more fine-grained appearance than the original contact maps, with fewer long runs of contact. Quantitative analysis showed that long contact runs, particularly above ~1.5 Mb, occur more

frequently in the real data than in the simulated random button model (Figure 4C). This supports the multivalent interaction model.

#### Long-Range Coordination of NL Contacts within Chromosomes

Interestingly, we also observed long runs of no-contact on many single chromosomes (white horizontal streaks in Figure 4B, top). Particularly in the range of >5 Mb, such runs occur more frequently than may be expected by random chance (Figure 4D). We interpret these long runs of no-contact as large chromosomal regions (often including multiple LADs) that completely dissociate from the NL in an incidental manner. These long no-contact runs are not generally due to loss of large chromosomal

regions during sample processing, because our Dam-only single cell maps have more homogeneous methylation patterns (Figure 2C) and lack long runs that are completely devoid of mappable sequence reads (Figure S3A).

This coordinated detachment of neighboring LADs from the NL prompted us to systematically search for evidence of coordination of NL contacts, by calculating pair-wise NL contact correlations for all 100-kb segments within each chromosome, based on the binary NL contact maps of the 118 single cells. The resulting correlation matrices, which indicate how pairs of loci may coordinately attach and detach from the NL in single cells, showed remarkable plaid-like patterns on all chromosomes (Figure 5A). Along the diagonal, squares of consistently positive correlations represent units of coordinated NL contacts, which tend to be one to several megabases in size. Most of these units exhibit additional off-diagonal correlations with other units, sometimes with striking specificity (Figure 5A, examples marked with arrows and boxes). Such coordinated units can be tens of megabases apart, although the frequency decays gradually with distance (Figure 5B). Randomized contact maps yield much lower average correlation coefficients over all distances, demonstrating that the prevalence of positive correlations is not due to random chance (Figure 5B). Intra-chromosomal coordination is on average higher than inter-chromosomal coordination, and the latter is not higher than may be expected by random chance (Figure S4A). These results point to extensive coordination of NL contacts within chromosomes, often over long genomic distances.

### Coordination of NL Contacts Is Partially Linked to Physical Proximity as Detected by Hi-C

The plaid-like correlation patterns reminded us of patterns commonly observed with Hi-C (Lieberman-Aiden et al., 2009; Dixon et al., 2012), a technology that maps proximity of genomic sequences in nuclear space (Dekker et al., 2013). An intriguing possibility was therefore that regions with coordinated NL contacts could be in spatial proximity to one another. To investigate this, we generated Hi-C maps from KBM7 clone #14 cells. The resulting Hi-C interaction matrices appeared partially similar to the NL contact correlation matrices (Figure 5C).

To quantify the similarity, we calculated the correlation between the degree of NL contact coordination and the Hi-C interaction frequency as a function of genomic distance (Figure 5D). The results show that Hi-C interactions correlate moderately but significantly with the degree of NL contact coordination. This correlation is most prominent at ~1–2 Mb distance and declines gradually over longer distances, but is still statistically significant ( $p < 0.001$ ) at ~100 Mb. Somewhat surprisingly, this positive correlation appears absent among directly neighboring 100-kb segments. One possibility is that pairs of adjacent 100-kb segments have very high Hi-C interaction frequencies due to their physical linkage, regardless of any coordination of NL contacts; indeed, our analyses show that finer-scale features of Hi-C do not always correspond to coordination of NL contacts (see below). Together, these data indicate that over a broad range of linear distances, coordinated NL contacts have a tendency to be linked to close proximity in nuclear space.

Previous Hi-C studies have revealed multi-scale compartmentalization of chromatin. At the highest level, megabase-sized do-

main are segregated into two main compartments that can be identified by eigenvector decomposition of the Hi-C matrix (Lieberman-Aiden et al., 2009). The CF pattern shows a remarkably tight correlation with this compartment score (Figure S4B; genome-wide Spearman's  $\rho = -0.88$ , weighted average of by-chromosome correlations), indicating that the two main Hi-C compartments largely correspond to NL-interacting and internally located chromatin.

At a finer scale, the genome is partitioned into topologically associated domains (TADs), which are discrete domains that have many intra-domain Hi-C interactions but relatively few interactions with neighboring TADs (Dixon et al., 2012; Nora et al., 2012). We computed TAD boundary positions and compared them to the NL contact coordination patterns. We found examples in which TAD boundaries coincide with borders of units of coordinated NL contacts, but also cases in which they do not coincide (Figure S4B). We discovered that only TAD boundaries that coincide with strong transitions in CF values also mark the edges of units of NL contact coordination, while TAD boundaries located in regions with relatively uniform CF values typically do not (Figure S4C). This is not due to a general difference in TAD boundary strength, because the average TAD boundary scores were highly similar (Figure S4C, bottom). We conclude that TADs overlap only partially with units of coordinated NL contacts.

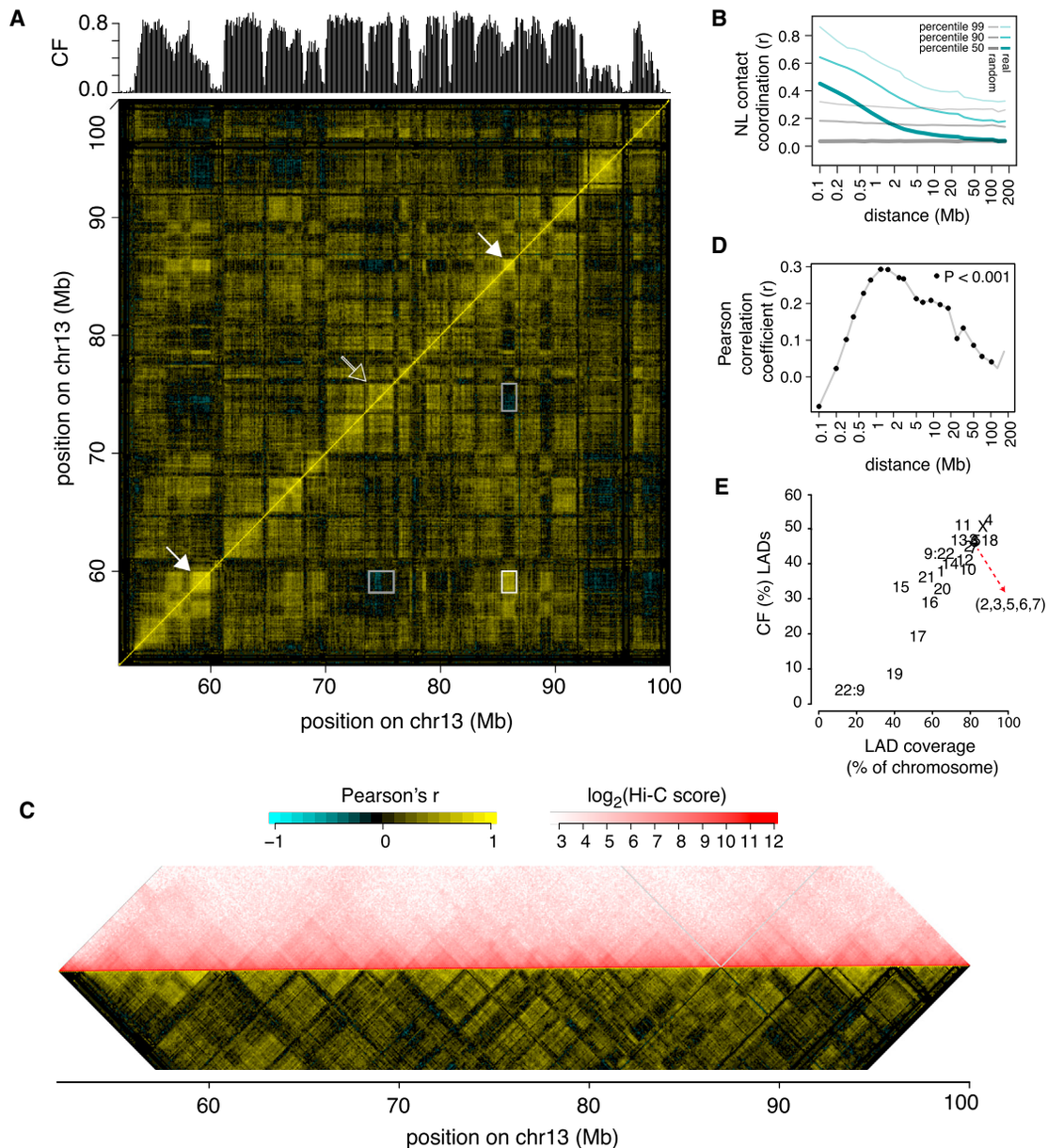
### Chromosomal CF Patterns Suggest Long-Range Cooperativity

The prevalence of intra-chromosomally coordinated NL contacts raised the possibility that multiple LADs across a chromosome associate with the NL in a cooperative manner. In support of this, we noticed a strong correlation between the average CF of all LADs along each chromosome and the overall density of LADs on the same chromosome (Figure 5E; Spearman's  $\rho = 0.85$ ,  $p = 9 \times 10^{-7}$ ). This is, for example, illustrated by the stark contrast between chr18 and chr19: the former has a very high density of LADs, of which many have consistent NL contacts, while the latter has only a few LADs that infrequently contact the NL (Figure 3A). The tight correlation between chromosomal density of LADs and their average CF is suggestive of chromosome-wide cooperative LAD-NL interactions.

### NL Contacts Often Involve Embedding of DNA in the NL

To gain more insight into the nature of NL contacts at the scale of single LADs, we visualized these contacts by ground state depletion (GSD) super-resolution fluorescence microscopy (Fölling et al., 2008) in combination with the <sup>m6</sup>A-Tracer method (Kind et al., 2013). We used a previously established clonal cell line (derived from HT1080 fibrosarcoma cells) that expresses inducible Dam-LmnB1, together with a GFP-tagged <sup>m6</sup>A-binding protein; this system allows for direct visualization of DNA that is, or has been, in contact with the NL (Kind et al., 2013). We labeled the NL in the same cells with an antibody against LmnB1 and used GSD microscopy to obtain super-resolution two-color images.

20 hr after Dam-LmnB1 induction, the <sup>m6</sup>A-Tracer signal exhibits a striking speckled pattern that is mostly confined to a zone of ~1  $\mu$ m underneath the NL (Figures 6A and 6B). The signal consists of clusters with diameters in the range of approximately

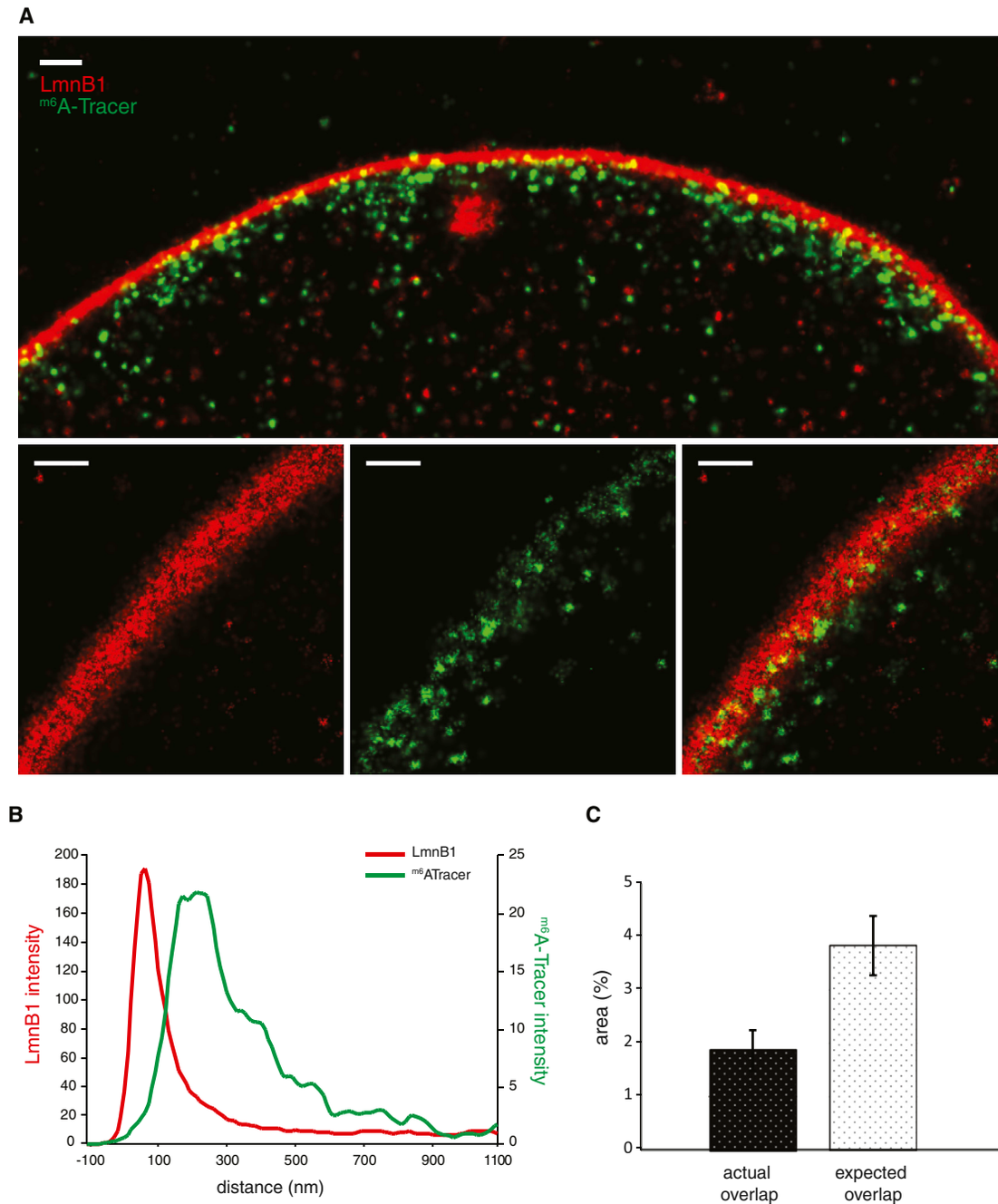


**Figure 5. Intra-Chromosomal Coordination of NL Contacts over a Wide Range of Distance**

(A) NL contact correlation matrix for part of chr13 in clone #14, showing the Pearson correlation of binary NL contacts across the 118 cells for all possible pairs of genomic segments. The color key is shown in panel (C). White arrows mark two example regions located ~26 Mb apart that exhibit coordinated contacts (white box); the open arrow indicates a region that is not positively correlated with these regions (gray boxes). (B) Genome-wide coordinated NL contacts as a function of linear distance. Median, 90<sup>th</sup> percentile, and 99<sup>th</sup> percentile Pearson's r values are shown for real (cyan) and randomized (gray) binary NL contact data from 118 single cells. (C) Comparison of NL contact coordination matrix to Hi-C interaction matrix. The matrix in (A) was turned clockwise by 45° and only a section below the diagonal is shown; the corresponding part of the Hi-C interaction matrix (white-red color scale) is juxtaposed to facilitate comparison. Gray lines mark a repetitive region. (D) Genome-wide correlation between NL contact coordination and Hi-C interactions, plotted as function of linear distance along chromosomes. Dots mark distances at which the correlation is significantly different from zero ( $p < 0.001$ ). (E) Positive correlation (Spearman's  $\rho = 0.85$ ,  $p = 8.96 \times 10^{-7}$ ) between the fraction of each chromosome covered by LADs and the average CF in these LADs. See also Figure S4.

50–300 nm. A similar pattern is seen 10 hr after Dam-LmnB1 induction (Figure S5A), although the signals are sparser than after 20 hr. Many of the m<sup>6</sup>A-Tracer signals do not touch the NL directly, indicating that they represent loci that contacted the

NL in the recent past and subsequently moved over a short distance toward the nuclear interior, as noted previously (Kind et al., 2013). Nevertheless, close association of m<sup>6</sup>A-Tracer signals with LmnB1 staining is frequently observed (Figure 6A).



**Figure 6. LAD-NL Interactions Involve Partial Embedding of Chromatin in the NL**

(A) GSD microscopy image sections perpendicular (top) and oblique (bottom) to the NL. Red: LmnB1; green: m<sup>6</sup>A-Tracer signal 20 hr after induction of Dam-LmnB1. The large red structure in the center of the top panel may be an invagination of the NL. Scale bars represent 500 nm.

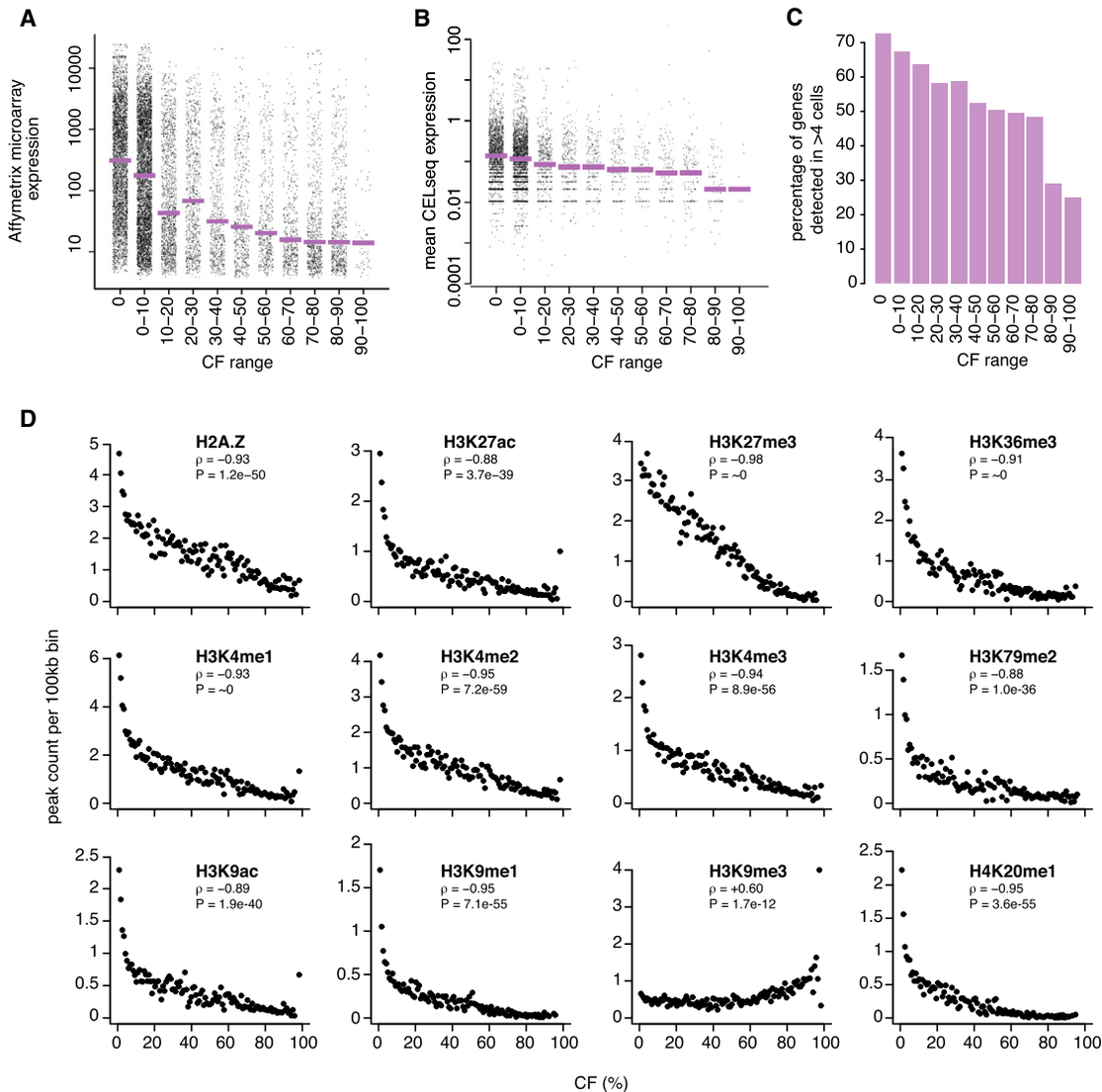
(B) Average pixel intensity of m<sup>6</sup>A-Tracer and LmnB1 signals as a function of the distance to the center of the NL. Curves show the average of four images.

(C) Quantification of the overlap of m<sup>6</sup>A-Tracer and LaminB1 signals within the confines of the NL, compared to random overlap. Data are the average of four nuclei. Error bars indicate SD.

See also Figure S5.

It is noteworthy that the edge of the LmnB1 signal at the NL is less sharply defined at the nucleoplasmic side than at the inner nuclear membrane side (Figures 6A and 6B). This could indicate that lamin filaments extend to varying degrees

into the nuclear interior, creating a somewhat fuzzy surface. Interestingly, the m<sup>6</sup>A-Tracer signals that abut the NL often appear partially embedded in this LmnB1 meshwork (Figure 6A, top). Oblique sections show that m<sup>6</sup>A-Tracer signals tend to



**Figure 7. Links of CF to Gene Expression and Chromatin Composition in Single Cells**

(A) Gene expression in pools of KBM7 cells (GEO: GSE56465; n = 7 independent clones) as a function of CF. Dots represent genes; purple bars show median values.

(B) Gene expression level in single clone #14 cells (mean of 96 single cells) as a function of CF. Dots represent genes; two genes with expression levels >100 are not shown. Purple bars show median values. Only genes with detectable expression in at least one cell are included.

(C) Fraction of genes with detectable expression in at least 5 out of 96 cells, as function of CF. Genes not detected in any of the 96 cells are not counted.

(D) Links of CF to epigenome mapping data from K562 cells (Consortium, 2012).

See also Figure S6.

occupy small pockets in the Lamin B1 signals (Figure 6A, bottom). Indeed, quantitative analysis of oblique sections showed that, within the confines of the NL, the overlap between <sup>m6</sup>A-Tracer clusters and LmnB1 is less than expected by chance (Figure 6C). This embedding is not caused by thickening of the NL due to expression of Dam-LmnB1, because the thickness of the NL is similar to that of cells in which Dam-LmnB1 is not induced (Figure S5B). Together, these results show that contact of LADs with the NL often involves embedding of chromatin in the relatively fuzzy nucleoplasmic surface of the NL.

### Links between NL Contacts, Single-Cell Gene Expression, and Chromatin State

Because most NL interactions have been linked to gene repression (Finlan et al., 2008; Guelen et al., 2008; Reddy et al., 2008; Peric-Hupkes et al., 2010), we asked how transcriptional activity of genes is linked to their CF. Analysis of publicly available gene expression profiles from pools of KBM7 cells revealed that with increasing CFs the distribution of gene expression levels shifts gradually toward lower values (Figure 7A). To investigate this further, we employed a modified CEL-seq method (Grün et al., 2014) to generate genome-wide mRNA expression profiles

from 96 single KBM7 cells. These data show that the average mRNA expression level, as well as the fraction of genes with detectable mRNA, declines with increasing CF values (Figures 7B and 7C), which is in agreement with the cell pool expression analysis. Thus, in general, the more stably a gene is associated with the NL, the less active it tends to be. We considered that genes with mid-range CFs (i.e., genes that associate with the NL in only a subset of cells) might exhibit more cell-to-cell variation in gene expression. Analysis of the CEL-seq data did not uncover such a relationship (Figures S6A and S6B), although we note that technical noise in CEL-seq data may obscure such biological noise, particularly because NL-associated genes are generally expressed at very low levels.

Finally, we investigated whether CFs are linked to the presence of specific histone modifications. While no genome-wide maps of these modifications are available for KBM7 cells, extensive datasets are available from K562 cells (ENCODE Project Consortium, 2012), which is another myeloid leukemia cell line. These data reveal clear negative correlations between CFs and 10 out of 11 tested histone marks, as well as the histone variant H2A.Z (Figure 7D), most of which have previously been linked to active transcription. Only H3K9me3, which is generally linked to gene repression (ENCODE Project Consortium, 2012), is positively correlated with CF values (Spearman's  $\rho = 0.60$ ,  $p = 1.7 \times 10^{-12}$ ). This is consistent with the previously reported role of H3K9 methylation in NL tethering (Towbin et al., 2012; Bian et al., 2013). In contrast, H3K27me3, which is also linked to gene repression, shows a strong negative correlation with CFs, suggesting that it does not play a role in NL tethering in KBM7 cells.

## DISCUSSION

### Single-Cell Protein-DNA Interaction Mapping by DamID

Here, we demonstrate that a modified DamID protocol can be used to map protein-DNA contacts genome-wide in single cells. At present, the resolution of the resulting maps is approximately 100 kb, which is suited to study proteins that form large domains along the genome (Bickmore and van Steensel, 2013). We expect that the resolution may be further improved by optimization of the DamID protocol and by deeper sequencing of samples so that more unique reads are recovered. It will be of interest to integrate single-cell DamID mapping with the recently reported single-cell Hi-C approach (Nagano et al., 2013).

### Most Chromosomes Have a “Backbone” of Consistent NL Interactions

One interesting outcome of this study is that about 15% of the genome contacts the NL in most of the cells (CF > 80%). We propose that these consistently interacting loci together form a “backbone” that may help to shape the overall chromosome architecture. Their strong overlap with constitutive LADs suggests a common backbone function in many cell types. The extremely low gene density within these loci suggests that they may have evolved to play a structural role. Not all cLADs have high CFs, but because virtually all chromosomes have multiple cLADs, the backbone may be robust due to redundancy. The precise distribution of loci with stable NL contacts may be critical; this is suggested by the previously reported strong evolu-

tionary conservation of the boundaries of cLADs (Meuleman et al., 2013). It will be of interest to investigate the consequences of deleting such stable contact sites from chromosomes.

### Intra-chromosomal Coordination of Variable NL Contacts and Spatial Proximity

Scattered between these stable contact sites are many loci that associate with the NL in only a subset of cells. Such loci may be subject to a balance between mechanisms that tether them to the NL and mechanisms that sequester them in the nuclear interior. These variable loci, which together cover about one-third of the genome, exhibit a complex pattern of coordinated NL contacts within chromosomes. This domain-like pattern presumably reflects the overall chromosomal architecture. Indeed, our Hi-C analysis shows that loci with intra-chromosomally coordinated NL contacts tend to be in close proximity in nuclear space, particularly in the 0.5–5 Mb distance range. Physical interactions between these loci may facilitate the coordination of their NL contacts. However, it is also possible that loci with coordinated NL contacts are more often in spatial proximity of one another simply because they are located in the same nuclear compartment.

### Multivalent Interactions and Embedding of Chromatin in the NL

The non-random long runs of NL contacts observed in our single-cell maps strongly suggest a multivalent mode of interaction. Considering that both the NL and chromosomes consist of polymer structures, such a mechanism is quite plausible. The Mb range over which we find this mechanism to be active appears at odds with a report claiming that a short 400-bp repetitive sequence was sufficient to target a locus to the NL (Zullo et al., 2012). One possibility is that this sequence was integrated as a long tandem repeat, which often happens in stable transfections. Another study identified three independent “peripheral targeting regions” in a human LAD, one of which could be narrowed down to 6.3 kb (Bian et al., 2013). However, this element was unable to target a free plasmid to the NL, while in genomic context its ability to interact with the NL was strongly dependent on the integration site, indicating that this element requires support from other elements.

H3K9me3 is strongly correlated with CF values and several studies have found that di- and tri-methylation of H3K9 can promote NL interactions (Towbin et al., 2012; Bian et al., 2013; Kind et al., 2013). These histone modifications tend to cover large genomic regions (Bickmore and van Steensel, 2013) and are thus probable candidates to be involved in multivalent chromatin-NL contacts.

The largest <sup>m6</sup>A-Tracer clusters (~300 nm in size) that we observed by super-resolution microscopy may contain ~100–250 kb of DNA, a rough estimate that we infer from previous polymer modeling of FISH distance measurements in mammalian nuclei (Mateos-Langerak et al., 2009; Giorgetti et al., 2014). However, most <sup>m6</sup>A-Tracer clusters are substantially smaller. We therefore suggest that a typical long (>1 Mb) contact run as observed in the single-cell maps may consist of a string of such clusters, each typically shorter than 100 kb, that contact the NL in a multivalent manner. Future improvement

of the resolution of single-cell DamID mapping may enable the identification of individual <sup>m6</sup>A-Tracer clusters at the sequence level.

## EXPERIMENTAL PROCEDURES

Detailed methods are described in the [Supplemental Experimental Procedures](#).

### Single-Cell DamID

The protocol for the detection of Dam methylation in single-cell DamID is similar to that of conventional DamID (Vogel et al., 2007) and uses largely the same reagents. Key differences are (1) use of clonal cell lines with controlled expression of Dam-LmnB1, (2) use of the Fucci system and flow sorting to collect single cells at the G1/S transition, (3) the solution used to lyse the cells, (4) performance of all enzymatic steps to detect Dam methylation in a single well of a 96-well plate, by sequential addition of reagents without any purification of the DNA, (v) 4–6 additional PCR cycles, and (5) use of Illumina sequencing instead of microarrays as readout. Multiplexing of samples was done with indexing primers as listed in [Table S2](#). A detailed description is provided in the [Supplemental Experimental Procedures](#), which also document the processing of sequencing reads.

### Hi-C, Gene Expression Analysis, and CEL-Seq

Hi-C data from clone #14 cells were generated in duplicate and processed as described (Belton et al., 2012; Lajoie et al., 2015). Gene expression profiles from pools of KBM7 cells were obtained from GEO: GSE56465. CEL-seq was done essentially as previously described (Grün et al., 2014).

### GSD Microscopy

Super-resolution microscopy was performed with a Leica SR GSD microscope (Leica Microsystems) with a Sumo Stage (#11888963) for drift-free imaging. Images were collected with an EMCCD Andor iXon camera (Andor Technology) and an oil-immersion objective (PL Apo 160x, na 1.46). Lasers used included 405 nm/30 mW (back-pumping only), 488 nm/300 mW, and 647 nm/500 mW. Between 10,000 to 50,000 frames were collected at 100 Hz for each SR image. Data were analyzed with the ImageJ ThunderStorm analysis module (Ovesný et al., 2014).

## ACCESSION NUMBERS

The accession numbers for the single-cell and conventional DamID, Hi-C, and CEL-seq reported in this paper are deposited in GEO: GSE69423, GSE69841, and GSE68596.

## SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, two tables, and five data files and can be found with this article at <http://dx.doi.org/10.1016/j.cell.2015.08.040>.

## AUTHOR CONTRIBUTIONS

J.K.: conceived of and designed the study; single-cell DamID development and experiments; data exploration; paper writing. L.P. and S.S.d.V. made equal contributions. L.P.: data processing pipeline; data analysis. S.S.d.V.: clonal cell line establishment; single-cell DamID; control experiments and experimental design. L.N.: super-resolution microscopy. S.S.D.: CEL-seq experiments and data analysis. M.B.: FISH experiments and data analysis. Y.Z.: Hi-C experiments. B.L.: Hi-C data processing. C.A.d.G.: unpublished LAD atlas; data processing support. M.A.: constructs; experimental support. G.F.: data analysis. M.I.: data analysis. L.A.M.: advisor of G.F. and M.I.; data analysis; intellectual input. K.J.: advisor of L.N.; superresolution microscopy experimental design and image analysis. J.D.: advisor of Y.Z. and B.L.; intellectual input. A.v.O.: advisor of S.S.D. and M.B.; intellectual input. B.v.S.:

conceived of and designed study; data analysis; project coordination; paper writing.

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