Fine-scale chromatin interaction maps reveal the cis-regulatory landscape of human lincRNA genes

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A number of powerful genome architecture assays, including DNA adenine methyltransferase identification (DamID)1 and chromosome conformation capture (3C)-based methods2, have greatly advanced our understanding of the three-dimensional (3D) organization of genomes but are limited in resolution by their reliance on restriction enzymes. Here we describe a method called DNase Hi-C for comprehensively mapping global chromatin contacts. DNase Hi-C uses DNase I for chromatin fragmentation, leading to greatly improved efficiency and resolution over that of Hi-C. Coupling this method with DNA-capture technology provides a high-throughput approach for targeted mapping of fine-scale chromatin architecture. We applied targeted DNase Hi-C to characterize the 3D organization of 998 large intergenic noncoding RNA (lincRNA) promoters in two human cell lines. Our results revealed that expression of lincRNAs is tightly controlled by complex mechanisms involving both super-enhancers and the Polycomb repressive complex. Our results provide the first glimpse of the cell type–specific 3D organization of lincRNA genes.

High-throughput methods based on chromosome conformation capture have greatly advanced our understanding of the three-dimensional (3D) organization of genomes but are limited in resolution by their reliance on restriction enzymes. Here we describe a method called DNase Hi-C for comprehensively mapping global chromatin contacts. DNase Hi-C uses DNase I for chromatin fragmentation, leading to greatly improved efficiency and resolution over that of Hi-C. Coupling this method with DNA-capture technology provides a high-throughput approach for targeted mapping of fine-scale chromatin architecture. We applied targeted DNase Hi-C to characterize the 3D organization of 998 large intergenic noncoding RNA (lincRNA) promoters in two human cell lines. Our results revealed that expression of lincRNAs is tightly controlled by complex mechanisms involving both super-enhancers and the Polycomb repressive complex. Our results provide the first glimpse of the cell type–specific 3D organization of lincRNA genes.

To overcome this limitation, we developed a method for mapping global chromatin interactions on the basis of random fragmentation with DNase I (DNase Hi-C; Fig. 1a). We combined this approach with DNA-capture technology to carry out a targeted, massively parallel dissection of local chromatin architecture at unprecedented resolution.

lincRNAs play key roles in various cellular and developmental processes12–14; however, the regulation of lincRNA expression remains largely elusive. We therefore applied targeted DNase Hi-C to map the chromatin configurations associated with 998 lincRNA-encoding genes (‘lincRNA genes’) in the human embryonic stem cell (hESC) line H1 and in the human chronic myelogenous leukemia cell line K562. Our method provides a paradigm for characterizing at high resolution how the 3D spatial organization of genomic loci correlates with transcriptional regulation in different cell types.

RESULTS

Development and validation of DNase Hi-C

The key difference between the conventional Hi-C protocol and DNase Hi-C is the use of DNase I instead of REs for fragmenting cross-linked chromatin. Unlike REs, which generate predictable and consistent fragment ends, DNase I generates a heterogeneous mixture of fragment ends composed of 5′ and 3′ overhangs of varying lengths as well as blunt ends. Consequently, we undertook extensive protocol modifications and optimizations to develop DNase Hi-C (Fig. 1a, Supplementary Notes 1 and 2 and Supplementary Fig. 1a).

We sought to validate the DNase Hi-C methodology by mapping chromatin contacts in H1 hESCs and K562 cells (Supplementary Table 1). Five observations demonstrate the reliability of DNase Hi-C. First, consistent with previous Hi-C studies8,9, both experimental libraries displayed the polymer-like properties characteristic of chromatin fibers; these properties were not evident in the non-cross-linked control library (Supplementary Fig. 1b). Second, even distal intrachromosomal contacts were much more frequent than interchromosomal contacts (Supplementary Fig. 1c), reflecting chromosome territories, as observed previously by cell imaging and Hi-C9,15. Third, chromosomes were segregated into open and closed compartments in both cell types (Supplementary Fig. 2a,b), as observed in previous Hi-C assays8. Fourth, chromosomes exhibited megabase-sized topological domains16 that were highly correlated with those identified previously by Hi-C16 (P < 10−500, Fisher’s exact test; Supplementary Fig. 2c). Fifth, side-by-side comparisons demonstrated that DNase Hi-C and RE-based Hi-C libraries possessed similar levels of chromatin accessibility–associated bias at both the local and large scale (Fig. 1b,c).

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Overall, with respect to biases associated with G+C content, mappability and RE sites, DNase Hi-C libraries performed slightly better than RE-based Hi-C libraries (Fig. 1d, Supplementary Fig. 1d,e and Supplementary Note 3). Notably, the small size and random distribution of DNA fragments within a DNase Hi-C library yielded better genome coverage than an RE-based Hi-C library (Fig. 1e).

Development and validation of targeted DNase Hi-C

We next combined the DNase Hi-C protocol with a method for targeted DNA sequence capture17 (NimbleGen SeqCap) to map chromatin contacts associated with genomic loci of interest (Fig. 1a, Supplementary Note 4 and Online Methods). In the DNase Hi-C library, chimeric DNA molecules contain two interacting fragments separated by an internal adaptor (Supplementary Fig. 1a). Probes designed to target regions of interest can thus be used to capture the corresponding contact partners that are brought along in tow.

In pilot studies using a 220-kb promoter-enhancer bait library designed to target 113 cis elements, including both promoters and enhancers (‘220kb P-E library’; Supplementary Table 5 and Online Methods), we proved the feasibility of targeted DNase Hi-C by demonstrating that the capture efficiency achieved by targeted DNase Hi-C was similar to that of DNA fragments in a control genomic DNA library without ligation (Supplementary Table 6). We also observed that the targeted DNase Hi-C data exhibited the expected polymer-like behavior of chromatin fibers (data not shown). We then mapped the contact profiles associated with each of the 113 cis elements in both H1 and K562 cells at multiple resolutions (2–50 kb; Fig. 2 and Supplementary Figs. 3–5). Domainograms18 showed that local contacts (<1-Mb genomic distance) were dominant for each target element (Supplementary Fig. 3), a finding consistent with previous observations in Hi-C and 4C-seq assays8,18,19. Moreover, targeted DNase Hi-C recapitulated the local 3D architectures of several genomic loci previously examined by 3C, 4C or ChIA-PET technologies, including the NANOG locus in hESCs20 (Fig. 2b and Supplementary Fig. 4) and the MYC locus in K562 cells21 (Supplementary Fig. 3). Targeted DNase Hi-C also captured known promoter-enhancer interactions, including the well-known interaction between the locus control region (LCR) and the γ-globin promoter in K562
Figure 2 | Validation of targeted DNase Hi-C. (a) Profile of targeted DNase Hi-C contacts within 250 kb of the HS2-HS3 region of the β-globin gene's locus control region in H1 and K562 cells. Red arrows indicate the position of the target in each domainogram. The geometric mean of read coverage in 5-kb sliding windows (computed in overlapping offsets of 1 kb) in each domainogram is indicated in the y axis. The color scale of each domainogram was set according to the range of geometric means in 12-kb windows (also computed with 1-kb offsets), and the contact frequency corresponding to the color scale is indicated. In the spline-fitting plots, the high-confidence bins (red dots) and those with false discovery rate (FDR) <0.05 (blue dots) and FDR <0.1 (green dots) are indicated, and the positions of merged high-confidence contacts are highlighted with pink bars. High-confidence contacts identified by targeted DNase Hi-C; DNase I hypersensitive sites (DHSs) from the UCSC Genome Browser, using data from the ENCODE Project Consortium; DNase Hi-C read coverage (at 1-base resolution); topological domains and the virtual 4C of the target region generated from H1 or K562 DNase Hi-C data; and the NCBI Reference Sequence Database (RefSeq) genes are shown. The β-globin genes are highlighted in orange. (b) Reproducibility of the contact profiles of the NANOG promoter. Intrachromosomal contacts within 250 kb of the target region are shown by domainograms. The contact frequency (geometric mean of the window coverage) corresponding to the color scale is indicated. Red arrows indicate the position of the target. The GDF3-DPPA3-NANOG locus is highlighted in green. The virtual 4C of the target region generated from H1 or K562 DNase Hi-C data and the RefSeq genes are also shown.
cells as well as the interaction between the SOX2 promoter and its 3’ enhancer in H1 ESCs (Fig. 2a, Supplementary Fig. 5 and Supplementary Note 5).

We developed a computational pipeline for identifying high-confidence contacts (Fig. 2a, Supplementary Fig. 6, Supplementary Table 7 and Online Methods). After merging adjacent contacts, we refer to a genomic locus that is in contact with a captured target locus as a ‘target partner’. In total, after merging adjacent contacts, we identified at 1-kb resolution 180 and 508 intrachromosomal local (<10 Mb) target partners associated with the 109 cis elements (among the 113 designed cis elements in the 220kb P-E bait library, 4 promoters were excluded owing to low coverage in both cell lines; see Online Methods) in H1 and K562 cells, respectively (Supplementary Tables 7–9 and Supplementary Fig. 7). We also identified long-range intrachromosomal (≥210 Mb) and interchromosomal contacts in the two cell lines (Supplementary Tables 8 and 9), but at lower resolution (10 kb); however, in this proof-of-concept study, we mainly focused on the intrachromosomal contacts (<10 Mb). Accordingly, only the uniquely mapped and nonredundant long-range (>1 kb) intrachromosomal read pairs were investigated, which account for ~1.6–8.5% of the total mapped read pairs in the targeted DNase Hi-C libraries (Supplementary Table 1). Therefore, all of the analyses described below pertain to the high-resolution local contacts (>1 kb, <10 Mb). We further validated our targeted DNase Hi-C method by comparing the high-confidence contacts (intrachromosomal, <10 Mb) identified by targeted DNase Hi-C with those identified in previous 5C (Supplementary Table 1). Also, consistent with previous observations, among the 180 targeted DNase Hi-C–identified high-confidence contacts in H1 ESCs and the 508 in K562 cells, only 60 contacts were shared by both cell types. This indicates that the target partners associated with the 109 cis elements were highly specific for cell type.

**lincRNA promoter–centered chromatin contacts**

To further establish the reliability of targeted DNase Hi-C and to demonstrate its use, we next applied targeted DNase Hi-C to investigate the detailed 3D chromatin signatures associated with the promoters of lincRNA genes in H1 and K562 cells. We designed a target library (‘lincRNA P library’; Supplementary Table 10 and Online Methods) for 1,030 distinct, well-annotated lincRNA gene promoters, and we generated targeted DNase Hi-C data sets for H1 and K562 cells, including independent replicates for each (Supplementary Table 1). We then assessed the specificity, efficiency, coverage uniformity, complexity and reproducibility of the targeted DNase Hi-C libraries (Supplementary Notes 3–5). First, we found that reads-on-target (i.e., at least one end lies within a target region) were highly enriched in the targeted DNase Hi-C libraries (Supplementary Figs. 8 and 9 and Supplementary Tables 11 and 12). The percentage of reads-on-target ranged from 7–10% for the libraries generated with the 220 kb P-E library to 35–57% for those generated with the 5-Mb lincRNA P library (Supplementary Tables 1, 11 and 13), which is similar to the reads-on-target of Capture-C libraries (10–17%) (Supplementary Table 13). However, among all the specifically captured paired-end reads in the Capture-C libraries, only about 10% (9.9–11.8%) of the reads corresponded to chromatin contacts between the target promoter regions and the rest of the genome, and about 90% (88.2–90.2%) were short-range reads with both ends located within the same target promoter region (Supplementary Table 14). In contrast, in our targeted DNase Hi-C libraries, at least 37% (37.1–76.0%) of the specifically captured paired-end reads were from chromatin contacts between the target promoter regions and the rest of the genome (Supplementary Tables 11 and 14), indicating that targeted DNase Hi-C is much more efficient than Capture-C (Supplementary Note 6).

Second, targeted DNase Hi-C libraries were highly correlated with the DNase Hi-C libraries in both cell types (Supplementary Fig. 9 and Supplementary Note 5). Third, targeted DNase Hi-C was highly reproducible (Fig. 2b, Supplementary Figs. 10 and 11, Supplementary Table 15 and Supplementary Note 5). These results suggest that our method is reliable and robust.
To characterize the lincRNA promoter contact partners, we next identified, by using 1-kb bins, 12,739 and 8,330 high-confidence intrachromosomal contacts (<10 Mb) associated with the 1,001 cis elements representing 998 lincRNA promoters and 3 positive controls in H1 and K562 cells, respectively (Fig. 3 and Supplementary Tables 7, 16 and 17). Note that 32 of the 1,030 targeted lincRNA promoters were excluded owing to low coverage in both cell lines (Online Methods). Remarkably, more than 39% of the RNA polymerase II–mediated contacts and 69% of the CCCTC-binding factor (CTCF)-mediated contacts associated with the lincRNA promoters identified in previous ChiA-PET studies\textsuperscript{21}, plus more than 38% of the significant contacts identified by the previous 5C study\textsuperscript{22}, were captured among our high-confidence contact lists (Supplementary Fig. 7). Although most lincRNA promoters were associated with <30 target partners within a 10-Mb genomic distance in both cell lines, several promoters had up to 77 partners (Supplementary Fig. 12a). After we merged adjacent significant contacts, the local intrachromosomal promoters (<10 Mb) ranged in size from 1 kb to over 100 kb (Supplementary Fig. 12b). Consistent with previous observations\textsuperscript{19}, the majority of the target partners for each lincRNA promoter occurred within the same topological domain, independent of the transcriptional status of the promoters\textsuperscript{(P < 10^{-222} in H1 ESCs, P = 2.8 × 10^{-27} in K562 cells; Fig. 3 and Supplementary Figs. 12c and 13–15). In addition, we found that 14.2% and 14.1% of the high-confidence target partners were shared by multiple lincRNA promoters in H1 and K562 cells, respectively, and that 49.2% and 36.5% of lincRNA promoters shared target partners with each other in H1 and K562 cells, respectively. These observations suggest that co-regulation of lincRNA gene expression, like that of protein-coding gene expression\textsuperscript{19,21,25,28}, is widespread. We also found that less than 20% of the target partners in H1 (1,485 of 12,739 high-confidence partners) and K562 (1,371 of 8,330 high-confidence partners) cells overlapped, indicating that the 3D organization of the lincRNA gene promoters is cell-type specific.

**Three-dimensional organization of the lincRNA promoters**

We next examined the genomic features associated with the lincRNA promoter–associated target partners. We first asked whether these lincRNA promoter–associated target partners were concentrated in cis elements as annotated by the Encyclopedia of DNA Elements (ENCODE) Consortium\textsuperscript{29,30}. We observed that, in both H1 and K562 cells, the lincRNA promoter–associated target partners were enriched in active promoters, enhancers, CTCF-binding sites, DNase I hypersensitive sites (DHSs) and formaldehyde-assisted isolation of regulatory elements (FAIRE)-defined open genomic regions (Fig. 4a–c and Supplementary Tables 18–20). However, when the regions marked by promoters, enhancers, CTCF-binding sites and DHSs were excluded from the FAIRE-defined open regions, the remaining (‘FAIRE-only’) regions were not enriched in the lincRNA promoter–associated target partners in both cell lines (Fig. 4a,b and Supplementary Table 18). Instead, FAIRE-only regions were depleted from the lincRNA promoter–associated target partners in K562 cells. In H1 cells, the lincRNA promoter–associated target partners were also enriched in poised promoters, indicating that some lincRNA gene promoters might also be in a ‘poised’ state in H1 ESCs. There were also notable differences between H1 and K562 cells. For example, the lincRNA promoter–associated target partners in K562 cells were enriched in transcribed gene bodies but not in Polycomb repressive complex (PRC)-repressed areas. Conversely, in H1 cells

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**Figure 4** | Identification of lincRNA promoter–associated cis elements. (a) Percentage of the given chromatin-state labels (Supplementary Table 18) that overlap with the lincRNA promoter–associated target partners in H1 and K562 cells. Z scores and P values were calculated using the Genome Structure Correlation. Red indicates enrichment, and blue represents depletion. The definition of each of the eight nonoverlapping segments (promoter, poised promoter, enhancer, transcribed, CTCF, FAIRE-only, repressed and dead) in H1 cells and of the seven (promoter, enhancer, transcribed, CTCF, FAIRE-only, repressed and dead) in K562 cells can be found in Supplementary Table 18 and the Online Methods. (b) Percentage of the lincRNA promoter–associated target partners that overlap with the various chromatin-state labels in H1 and K562 cells. (c) Percentage of the DNase I hypersensitive sites (DHSs) and FAIRE peak regions that overlap with the lincRNA promoter–associated target partners in H1 and K562 cells. In a–c, * indicates 3 < |Z| < 5; ** indicates |Z| ≥ 5. In each cell line, the combined targeted DNase Hi-C library of the two biological replicates was used for these analyses.
the PRC-repressed areas, but not the transcribed gene bodies, were highly enriched in the lincRNA promoter–associated target partners. When comparing the active lincRNA promoters with the inactive ones, we found that the active lincRNA promoter–associated target partners were more enriched in annotated cis elements (promoters, enhancers and CTCF-binding sites) and more depleted of heterochromatin regions in both cell lines (Supplementary Table 18).

We next asked whether lincRNA gene promoters are physically associated with super-enhancers31,32. Strikingly, we found that 180 of 695 known super-enhancers in K562 cells and 62 of 635 in H1 ESCs overlapped with lincRNA promoter–associated target partners (Fig. 5a and Supplementary Table 21). Among the 998 distinct lincRNA promoters, 151 in K562 and 70 in H1 cells were in contact with super-enhancers (Supplementary Table 21). Although most super-enhancers were associated with only a single lincRNA promoter, a few were in contact with multiple lincRNA promoters (up to 5; Fig. 5b). Conversely, some lincRNA promoters were in contact with multiple super-enhancers (up to 5 in K562 cells and up to 2 in H1; Fig. 5b, Supplementary Figs. 13 and 14 and Supplementary Table 21). Interestingly, although the majority of chromatin contacts between super-enhancers and lincRNA promoters were short range (<1 Mb genomic distance), a substantial portion of the contacts spanned >2 Mb of genomic distance (Fig. 5c). It is well established that spatial contacts between cis elements are integral to their functions33,34. Hence, these observations suggest that super-enhancers might be extensively involved in the transcriptional regulation of lincRNA gene expression.

lincRNA gene expression has been shown to be tightly regulated12–14, and the use of super-enhancers has been shown to be highly cell-type specific31. We therefore investigated the connections between lincRNA expression and super-enhancer association. First, we observed in both cell lines that the expression of super-enhancer–associated lincRNA genes was higher than that of lincRNA genes not associated with super-enhancers (Fig. 5d); however, we also observed that in each cell line a substantial number of lincRNA genes with undetectable gene expression were nonetheless in contact with super-enhancers (Supplementary Table 21). Second, we confirmed the cell type–specific usage of super-enhancers. This cell-specific usage of super-enhancers was due mainly to the cell-type specificity of the super-enhancers themselves (data not shown). Third, we observed that super-enhancer association coincided with cell type–specific lincRNA expression (Supplementary Figs. 13–15 and Supplementary Table 21).

To further characterize the lincRNA promoter–centered cis-regulatory networks, we examined the transcription factor (TF)
binding status of the target partners of the 998 lincRNA promoters in both H1 and K562 cells. The ENCODE Project Consortium has mapped the TF-binding sites (TFBSs) of 50 TFs in H1 cells and 100 TFs in K562 cells\textsuperscript{35}, including 36 TFBSs mapped in both cell lines (Supplementary Table 22). We found evidence that expression of lincRNA genes is tightly controlled but regulated by distinct mechanisms in H1 and K562 cells. All of the mapped TFBSs except those of two TFs in K562 cells were enriched in the lincRNA promoter–associated target partners (Supplementary Fig. 16a and Supplementary Table 22), which is consistent with our general observation that lincRNA promoter–associated target partners are enriched in cis-regulatory elements (Figs. 4 and 5a). Surprisingly, two of the three most enriched TFBSs among the 50 mapped in H1 cells represent binding sites for two components of the PRC2 complex, EZH2 and SUZ12 (Supplementary Fig. 16a and Supplementary Table 22). By comparison, EZH2-binding sites were not enriched in K562 cells. PRC2 trimethylates histone H3 on lysine 27 (H3K27me3), a mark of transcriptionally silent chromatin that is required for the initial setting of a genomic region to be silenced. These observations are consistent with the recent findings that Polycomb-group proteins play a role in the genome 3D organization in ESCs\textsuperscript{36}. For example, the promoters of two lincRNAs that are not expressed in H1 ESCs, HOTAIR and TCONS\textsubscript{00018052}, were bound by EZH2 and SUZ12, marked by H3K27me3 and associated with multiple PRC2-repressed sites in H1 cells but not in K562 cells (Fig. 3 and Supplementary Figs. 15 and 16b), indicating that the silent or poised state of these two lincRNAs involves PRC2 in H1 ESCs but not in K562 cells. Unlike in H1 cells, in which 4 of the 11 most enriched TFs were repressors, in K562 cells all of the ten most enriched TFs in the lincRNA promoter–associated target partners were transcriptional activators, including oncogenes TAF1, PML, MYC, ATF1 and IRFI, with TAL1 and JUN ranked at 12th and 14th, respectively (Supplementary Fig. 16a and Supplementary Table 22), coinciding with the status of K562 as a cancer cell line.

**DISCUSSION**

DNase Hi-C and targeted DNase Hi-C represent significant steps toward overcoming the RE digestion–associated resolution limit of existing genome architecture assays. In addition, applications such as chromosome-scale scaffolding of de novo genome assemblies\textsuperscript{37,38} and whole-genome haplotype reconstruction\textsuperscript{39} will also benefit from the improved resolution, reduced sequence bias and higher genome coverage of DNase Hi-C.

Currently, targeted mapping of fine-scale chromatin conformation can be accomplished using methods centered on either protein complexes (ChIA-PET) or genomic loci (4C). Recently, Hughes et al.\textsuperscript{31} published Capture-C, a method similar to targeted DNase Hi-C, for high-throughput mapping of physical contacts among cis-regulatory elements. Capture-C is a combination of 3C with DNA-capture technology; hence, the limitations associated with RE digestion also apply to Capture-C (Supplementary Table 14, Supplementary Fig. 17 and Supplementary Note 6). One of the straightforward applications of targeted DNase Hi-C will be to systematically link disease-associated noncoding single-nucleotide polymorphisms to their target genes in the context of nuclear 3D organization. Targeted DNase Hi-C may also prove valuable for characterizing phenotype-associated chromatin 3D signatures and for probing the relationship between genome architectural defects and disease pathogenesis.

This fine-scale approach should have wide applications for the identification of regulatory elements and targets of specific genes, as we demonstrate for lincRNAs. Our work, for the first time (to our knowledge), reveals a potential link between the transcriptional regulation of lincRNA genes and two master regulators of development: super-enhancers and the PRC2 complex. Polycomb-group proteins are essential for early development, and PRC2 has been shown to control the expression of developmental genes in hESCs\textsuperscript{40}. Together, these observations support an important developmental role for lincRNAs.

**METHODS**

Methods and any associated references are available in the online version of the paper.

**Accession codes.** NCBI Gene Expression Omnibus: sequencing data are available at accession number GSE56869.

*Note:* Any Supplementary Information and Source Data files are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

W.S.N., C.A.B., C.M.D., A.K. and Z.D. conceived of the project. Z.D. developed methods and supervised W.S.N., C.A.B., C.M.D., A.K. and Z.D. Z.D., C.A.B., J.S., A.K., C.M.D., X.D., C.B.W., W.S.N., C.A.B., C.M.D., A.K. and Z.D. wrote the paper.

**COMPETING FINANCIAL INTERESTS**

The authors declare no competing financial interests.

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ONLINE METHODS

Cell culture. K562 (ATCC, CCL-243) cells were grown in RPMI-1640 with 10% FBS and a penicillin/streptomycin mix (100 units/ml and 100 mg/ml, respectively). The H1 (WA-01) ESC line was obtained from WiCell Research Institute and was cultured as previously described. Briefly, the cells were cultured on a feeder layer of irradiated primary mouse embryonic fibroblasts (MEFs) in Dulbecco’s modified Eagle’s medium (DMEM)/F-12 medium supplemented with 20% serum replacer, 1 mM sodium pyruvate, 0.1 mM non-essential amino acids, 50 U/ml penicillin, 50 µg/ml streptomycin, 0.1 mM β-mercaptoethanol (Sigma-Aldrich) and 4 ng/ml basic fibroblast growth factor (bFGF).

Prior to the experiments, the cells were transferred to growth factor-reduced Matrigel (Becton Dickinson) in MEF conditioned medium (CM). All reagents are from Invitrogen unless otherwise specified. All cell cultures were ensured to be mycoplasma contamination-free.

Generation of DNase Hi-C and targeted DNase Hi-C libraries.

**Formaldehyde cross-linking.** 2 x 10^6–5 x 10^6 K562 or H1 cells were cross-linked with 1% or 2.5% formaldehyde, respectively, for 10 min at room temperature. Fixation was quenched with 0.125 M glycine for 10 min at room temperature. Fixed cells were washed with PBS and resuspended in cold lysis buffer 1 (10 mM Tris, 10 mM NaCl, 0.2% Igepal and complete protease inhibitor (Roche)) for 10 min. Cells were then snap frozen to 80 °C.

**DNase I digestion.** 2 x 10^6–5 x 10^6 cells were thawed on ice, resuspended in 1,000 µl TE lysis buffer (50 mM Tris (pH 7.0), 1 mM EDTA, 1% SDS) and incubated at 37 °C for 10 min. Nuclei were then collected, washed once with 0.15% Igepal, resuspended in 0.5 x DNase I digestion buffer (5 mM Tris-HCl (pH 7.5 at 25 °C), 5 mM MnCl_2, 0.05 mM CaCl_2, 0.25 unit/µl RNase A (Roche)), incubated at 37 °C for 10 min and digested with 0.015 unit/µl (for example, 5–6 µl in 400 µl reaction volume) DNase I (Fermentas) for 5 min at 25 °C. Digestion was stopped by adding 1/10 reaction volume of 10× stop solution (10% SDS, 250 mM EDTA). Twice the reaction volume of Ampure XP SPRI magnetic beads (Beckman Coulter) was added to the reaction, mixed well, divided into three Eppendorf tubes, incubated at room temperature for 5 min, collected via a DynaMag-Spin magnet (Invitrogen), washed twice with 80% ethanol and air dried for 2 min.

**End repair, dA-tailing and ligation of biotin-labeled bridge adaptors.** For DNA fragment-end repair, in each of the three tubes, air-dried beads attached with DNase I-digested chromatin complexes were resuspended in 400 µl 1× T4 ligation buffer (Fermentas) containing 0.25 mM dNTPs, 0.075 unit/µl T4 DNA polymerase (Fermentas, 6 µl) and 0.15 unit/µl Klenow fragment (Fermentas, 6 µl) and incubated at room temperature for 1 h. The reaction was stopped by adding 0.75% SDS, and beads were collected as described above.

dA-tailed chromatin complexes were resuspended in 200 µl 1× rapid ligation buffer (Fermentas) containing 25 units of T4 DNA ligase (Fermentas), 10 µM T-tailed, biotin-labeled bridge adaptor (Supplementary Table 23) and 20 µM blunt-ended non-biotin-labeled bridge adaptor (Supplementary Table 23; as not all of the chromatin fragments are dA-tailed after the above dA-tailing step, to reduce spurious ligation products resulting from blunt-ended chromatin fragments, this blunt-ended non-biotin-labeled adaptor is added as a blocker) and incubated at room temperature for 1 h. The reaction was stopped by adding 0.75% SDS, and beads were collected as described above. Note that ligation of the bridge adaptors can also be carried out at 16 °C overnight under the regular T4 DNA ligation conditions following the manufacturer’s instructions (Fermentas).

**Fragment-end phosphorylation and in-gel ligation.** Adaptor-ligated chromatin complexes were resuspended in 200 µl 1× T4 ligation buffer (Fermentas) containing 0.5 unit/µl T4 polynucleotide kinase (New England Biolabs) and incubated at 37 °C for 1 h. The above 200-µl reaction complexes were transferred to a 15-ml tube, and 11.8 ml of prewarmed (37 °C) 1× T4 ligation buffer (Fermentas) containing 0.4% UltraPure low–melting point agarose (Life Technologies) and 200 units of T4 DNA ligase (Fermentas) were added to the tube. The gel mixture was then well mixed and quickly solidified in ice water. Ligation was carried out at 16 °C for 4 h and 25 °C for 1 h.

**Reverse cross-linking and DNA purification.** Following ligation, the agarose gel was melted by incubating at 70 °C for 10 min. The reaction tube was transferred to a 42 °C water bath incubator. After 10 min preincubation, 30 µl of agarase (Fermentas) were added to the tube, and digestion of the agarose gel was carried out overnight. The ligation mixture was then concentrated to about 1 ml final volume by using the Amicon Ultra-15 centrifugal filter units (NMWL, 30 kDa, Millipore) according to the manufacturer’s instruction. Protein digestion was carried out by adding 80 µl of 20 mg/ml proteinase K (Fermentas) and 100 µl of 10% SDS, and the tubes were incubated overnight at 65 °C. The next day, an additional 10 µl of 20 mg/ml proteinase K was added to each tube, and the incubation was continued at 55 °C for another 2 h. DNA was precipitated with 3 µl GlycoBlue (Ambion), 0.3 M Na-acetate (pH 5.2) and equal volume of isopropanol (–80 °C, 2 h). Precipitated DNA was further purified by QIAquick Gel Extraction Kit (Qiagen). Concentration was determined by measurement on a Nanodrop-1000 spectrophotometer (Thermo Scientific).

**DNA dangling end removal, DNA fragmentation, library end repair, dA-tailing and sequencing adaptor ligation.** Approximately 5 µg purified DNA was treated with 50 units T4 DNA polymerase (Fermentas, 5 U/µl) in a 500-µl reaction containing 0.2 mM of each dATP and dGTP at 25 °C for 30 min. The reaction was stopped by adding 25 µl 0.5 M EDTA. The reaction mixture was then concentrated to about 20 µl final volume by using the Amicon Ultra-0.5 centrifugal filter units (NMWL, 30 kDa, Millipore) according to the manufacturer’s instructions. TE lysis buffer (50 mM Tris (pH 7.0), 1 mM EDTA, 1% SDS) was added to bring the volume to 110 µl. The DNA fragments were sheared to a size of 100–300 bp using a Covaris S2 instrument with the following parameters: duty cycle, 2%; intensity, 5; cycles per burst, 200; set mode, frequency sweeping; and number of cycles, 5. The liquid was transferred to a 1.5-ml Eppendorf tube, and the volume was...
brought to 200 μl. The DNA was then attached to 200 μl Ampure XP beads, washed with 80% ethanol twice and air dried.

DNA end repair was carried out by resuspending the beads in 100 μl 1× end repair reaction mix (Fermentas) containing 4 μl end repair enzyme mix and incubated at 16 °C for 10 min. The reaction was stopped by adding 0.5% SDS, and beads were collected as described above.

Beads with end-repaired DNA were then resuspended in 100 μl 1× NEBuffer 2 (New England BioLabs) containing 2 mM dATP and 0.1 unit/μl Klenow (Exo-) (New England BioLabs) and incubated at 37 °C for 1 h. The reaction was stopped by adding 0.5% SDS, and beads were collected as described above.

Beads with dA-tailed DNA were resuspended in 50 μl 1× rapid ligation buffer (Fermentas) containing 3 μM T-tailed Illumina sequencing adaptor (Supplementary Table 23) and incubated at room temperature for 30 min. The reaction was stopped by adding 0.5% SDS, and beads were collected as described above. DNA was then eluted to 200 μl water.

**Biotin pulldown, whole-genome chromatin interaction library amplification and purification.** Biotin-labeled, paired-end adaptor-ligated DNAs were immobilized to Dynabeads MyOne C1 streptavidin beads (Life Technologies) as follows. 25 μl of MyOne C1 beads were washed with 400 μl 1× binding and washing (B&W) buffer (5 mM Tris-HCl, pH 7.5, 0.5 mM EDTA, 1 M NaCl). The beads were isolated from bulk solution via a DynaMag-SPIN magnet (Invitrogen). The beads were then washed once with 400 μl 1× BSA (NEB) and once with 400 μl 1× B&W buffer. Isolated beads were resuspended in 200 μl 2x B&W buffer and combined with the above 200 μl labeled library DNA. The reaction was incubated at 25 °C for 20 min and constant rotation. The supernatant was removed, and the DNA-bound beads resuspended in 300 μl 1× B&W plus 300 μl 1× TE lysis buffer and transferred to a new tube. The beads were further washed twice with 600 μl 1× B&W buffer, once with 600 μl 1× NEBuffer 2 and once with 600 μl 1× EB buffer (Qiagen). The beads were resuspended in 80 μl 1× EB buffer.

For DNase Hi-C library generation, the above enriched whole-genome chromatin-interaction library was PCR amplified for a variable number of cycles (between 9 and 11) with KAPA HiFi HotStart DNA polymerase (Kapa Biosystems) and the standard Illumina paired-end indexed primer pairs (Supplementary Table 23). Approximately ten 100-μl PCR reactions were carried out for each library. PCR products for each library were pooled and size fractionated with the Ampure XP beads. DNA concentration was determined by measurement on a Nanodrop-1000 spectrophotometer. DNase Hi-C libraries were submitted for paired-end sequencing, the enrichment efficiency of each targeted DNase Hi-C library was semiquantitatively assessed via quantitative PCR (ABI 7500) with SYBR green. The identities of qPCR products were confirmed by both DNA gel electrophoresis and melting-curve analysis. Primers used in qPCR assays are listed in Supplementary Table 23.

**Design of the targeted DNase Hi-C probe (bait) libraries using the SeqCap custom library design technology.** The targeted DNase Hi-C probe libraries were designed using the online tool NimbleDesign (http://www.nimblegen.com/products/nimbledesign/index.html). The 220-kb promoter-enhancer library (design ID: 120924 HG19 200kb CRE EZ), which targets 110 known cis-regulatory elements (Supplementary Table 5), was initially designed to test the feasibility of targeted DNase Hi-C. All of the target elements were manually selected, and the positions of the gene promoters were based on the TSSs annotated in RefSeq and the UCSC “known gene” data sets for human genome build hg19. The genomic position of Vista enhancers was extracted from the VISTA Enhancer Browser (http://enhancer.lbl.gov/). The enhancer activities of these Vista enhancers were validated by transgenic reporter assays according to the VISTA Enhancer Browser. In general, this library was designed according to three considerations. (i) To provide a convenient way for validating the results of targeted DNase Hi-C assays, all of the selected cis elements were experimentally annotated, and most of them are well characterized. (ii) The total size of the target regions is relatively small (220 kb) to avoid the enormous sequencing efforts required for validating the results and to allow quick turn-around time of optimization. (iii) The size of each target region is small (2 kb, except that the target covering the HS2-HS3 sites of the β-globin LCR is about 7 kb) to test the sensitivity of targeted DNase Hi-C.

The lincRNA promoter library (design ID:130408 HG19 lincRNA P2 EZ HX3) targets the 5-kb promoter regions (from the TSSs to the 5 kb upstream) of 1,030 distinct (nonoverlapping) lincRNA genes (Supplementary Table 10). These lincRNA genes were independently annotated by two research groups, and their expression in H1 ESCs or K562 cells has been validated. The three lincRNAs described in ref. 42—lncRNA ESI1, lncRNA...
ES2 and lncRNA ES3—were also included in the library. Plus, the 7-kb region covering the HS2 and HS3 sites of the β-globin LCR and the 5-kb proximal promoter regions of the NANOG and SOX2 genes, which were covered in the 220-kb probe library, were included in this lincRNA library as positive controls for evaluating the performance of targeted DNase Hi-C assays with this probe library.

**Computational analysis of DNase Hi-C libraries.**

**Mapping and processing of sequence reads.** We sequenced both the DNase Hi-C and the targeted DNase Hi-C libraries using paired-end reads of length of 150 bp (raw sequencing files are available at http://noble.gs.washington.edu/proj/dnase-hi-c/). Because the DNase Hi-C data were multiplexed, we demultiplexed the reads using the exact 8-bp barcodes. Then we performed an exhaustive search and cleaning of the Illumina primer and adaptor sequences in the 150-bp full-length reads and extracted the remaining read fragments of various lengths from 25 to 80 bp. We then mapped each end of these cleaned paired-end reads separately to the human genome (GRCh37/hg19 assembly, obtained from the UCSC Genome Browser) using BWA. We retained only the reads that mapped uniquely, allowing at most three mismatches and requiring mapping quality MAPQ ≥30. For subsequent analysis, we only used read pairs for which both ends were successfully mapped according to the above criteria. Finally, to eliminate the bias due to the PCR duplication step, we removed redundant paired-end reads. We define two reads as redundant if both ends of the reads are mapped to identical locations in the genome.

After PCR duplicate removal, we generated whole-genome contact maps at both 1-Mb and 40-kb resolutions. To do so, we partitioned the whole genome into nonoverlapping bins and counted the number of contacts (i.e., mapped paired-end reads) observed between each pair of bins. The dimension of the resulting contact map is the total number of bins in the genome, and entry \((i, j)\) is the contact count between bin \(i\) and \(j\).

**Normalization.** We normalized the whole-genome contact maps obtained from DNase Hi-C data using an iterative correction method. We first preprocessed the contact maps at 1-Mb or 40-kb resolution by setting the entries that may be dominated by self-ligation products to 0. These entries are the diagonal, super-diagonal (+1 off-diagonal) and sub-diagonal (−1 off-diagonal) contact counts. In addition, we excluded from the correction process, as previously suggested, bins with the lowest 2% read coverage. Last we applied the iterative correction procedure on this preprocessed contact map to obtain a normalized contact map with near-equal row and column sums.

**Topological domain calling.** We identified topological domains using a previously described hidden Markov model–based software tool. To facilitate direct comparison with the previously published topological domains in H1 cells, we carried out the domain calling for DNase Hi-C data in H1 cells using human GRCh36/hg18 assembly. We applied the topological domain calling on normalized contact maps of our DNase Hi-C data at 40-kb resolution. In total, we obtained 2,528 and 2,040 domains in the H1 and K562 DNase Hi-C data sets, respectively. As in previous work, we classified the regions between the topological domains either as “domain boundaries” (≤400 kb) or “unorganized chromatin” (>400 kb).

To measure the consistency between the topological domains inferred from DNase Hi-C and those from published Hi-C data in H1 ESC cells, we calculated the overlaps of domain boundaries obtained between these two assays. We deemed two boundaries, one from each assay, as overlapping if they overlap by at least 1 bp or are adjacent to each other, as described in ref. 16. We performed 1,000 domain shufflings to calculate the expected domain boundary overlaps, similarly as described in the “Enrichment of intradomain targeted DNase Hi-C contacts” section. We used Fisher’s exact test to determine the statistical significance of the observed domain boundary overlaps.

**Eigenvalue decomposition and chromatin compartments.** We carried out eigenvalue decomposition on the normalized contact maps of H1 and K562 DNase Hi-C data sets as described in Lieberman-Aiden et al. For each chromosome, we used the intrachromosomal contact matrices at 1-Mb resolution. We calculated the Pearson correlation between each pair of rows of the contact matrix and applied eigenvalue decomposition (using the eig function in MATLAB) to the correlation matrix. The sign of either first or the second eigenvector defined chromosome compartments for each chromosome. Similar to Lieberman-Aiden et al., we used the second eigenvector in cases where the first eigenvector values are either all positive or all negative. We then compared the percentage of 1-Mb bins that were assigned the same compartment label from our DNase Hi-C libraries as from previously published Hi-C data in the same cell line (H1 (ref. 16), K562 (ref. 9)).

**Coverage comparison between DNase Hi-C and Hi-C.** We compared the percentage of the genome covered by the reads from the DNase Hi-C assay and that from the original Hi-C assay. We used the high-quality mapped paired-end reads after the PCR duplication removal step from two cell lines from each assay. For DNase Hi-C, we used H1 and K562 libraries generated in this work. To control for differences in sequencing depth and read length, we subsampled each library/replicate to the same number of reads and enforced a uniform read length of 50 bp per read end. We performed subsampling at two different read depths: ~15M and ~30M paired-end reads, corresponding to expected genome coverage percentages of 50% and 100%, respectively. We computed the genome coverage as the percentage of base pairs covered by at least one read over the entire genome length. We repeated the subsampling 20 times for each library/replicate. The variance of coverage across different samplings was negligible and was therefore not reported. In addition to the two subsampling comparisons, we also computed the percentage of the genome covered using all the reads available for each library/replicate. For RE-based Hi-C libraries, the specific (i.e., valid) reads should always be within the vicinity (e.g., ±500 bp) of a site of the RE (e.g., HindIII) that was used to generate the Hi-C library. Given the RE of choice and the reference genome that is used to map the reads, we can compute the theoretical upper bound on the coverage of an RE Hi-C library using only specific (±500 bp) and long-range (>1 kb) read pairs. For libraries created from the human genome (hg19) using HindIII, this upper bound on the coverage is 23.1% (dashed line), which has not been reached by published RE Hi-C data sets.

**Computational analysis of targeted DNase Hi-C libraries.**

**Identification of target captured contacts.** We trimmed, mapped and filtered the paired-end sequencing reads from targeted
DNase Hi-C data in similar fashion to the trimming, mapping and filtering of the DNase Hi-C data as described in the “Mapping and processing of sequence reads” section. However, for targeted DNase Hi-C data, we performed an additional filtering step to keep only the paired-end reads for which at least one end mapped to or within 150 bp from one of the captured target regions. We used these target-captured reads to define contact maps at 1-kb and 10-kb resolutions.

We next measured the capture efficiency of each target region as its captured read coverage (number of captured read pairs per kilobase of target region length). We identified 4 out of 113 targets in the 220kb P-E library, and 32 out of 1,033 targets in the lincRNA-P library with very low captured read coverage (lower than 25% of the average captured read coverage among all targets) in both H1 and K562 cells. These low-coverage targets are located mainly in unmappable genomic regions. We excluded them in further analyses.

Visualization of targeted DNase Hi-C contact profiles. To visualize intrachromosomal contact profiles, we plotted the domainograms near each target locus using the Cseqpipe software. The upper panel of the domainogram shows the main trend (at 5-kb resolution) of the contact profile. The lower panel reflects multiple-scale contact profiles at 2-kb to 50-kb resolution.

Normalization of targeted DNase Hi-C data. To correct biases in targeted DNase Hi-C data, we estimated a bias factor for each bin (either 1 kb or 10 kb). To do so, we first set the bias of unmappable bins (mappability score <0.5) to be 1. A mappability score of 0.5 for a bin means that half of the bases in that bin are not uniquely mappable for 50-bp reads. We calculated mappability scores using GEM46. Then we assessed the biases in target bins (those overlapping with the designed target regions on the DNA-capture array) and nontarget bins, separately, using the following strategies.

For bins that overlap with target regions, we approximated their biases by measuring their coverage in the targeted DNase Hi-C data. The bias factor is calculated as the number of captured read pairs at each target region per kilobase of the target region length. Bins overlapping with the same target region share the same bias factor. We then normalized the bias factors at target bins so that their average equals to 1.

For bins that do not overlap with any target, we estimated the biases from corresponding whole-genome DNase Hi-C data in the same cell type. First, assuming, as in the ICE method, that all nontarget bins have equal ‘visibility’, we took the bin coverage (i.e., the row margins of the contact matrix) at either 1-kb or 10-kb resolution, normalized by dividing by the average among all mappable bins in the genome. We then truncated the normalized bin coverage at 5 and 95 percentiles and performed smoothing by taking the average of ten neighboring bins. Our normalization method is similar to ICE in the sense that taking the contact margins is equivalent to the ICE correction with only one iteration. Accordingly, we have observed that contact margins and ICE iteratively learned bias factors are highly correlated at 40-kb resolution (Supplementary Fig. 6a,b). In our case, we chose not to perform iterative corrections at 1-kb resolution because the contact matrix becomes very large and sparse at 1-kb resolution and at the current sequencing depth. Consequently, the normalization procedure is computationally expensive and is also unstable.

Spline fitting and statistical confidence estimation. We applied the Fit-Hi-C method to our targeted DNase Hi-C data sets to identify statistically significant contacts associated with the target regions. The Fit-Hi-C approach uses an iterative spline-fitting procedure to estimate the null distribution of intrachromosomal contact probability at any given genomic distance and calculates statistical significance of observed contact counts using a binomial model. To apply the Fit-Hi-C method to the targeted DNase Hi-C data, we made two modifications to the original method. First, because in targeted DNase Hi-C experiments, DNase I was used instead of restriction enzymes, we aggregated chromatin contacts using fixed-size bins (either 1 kb or 10 kb) instead of aggregating within restriction enzyme fragments. Second, we estimated the null contact probability using only the pairs of loci with at least one end overlapping one of the captured target regions.

The Fit-Hi-C method also combines the genomic distance effect together with the normalization biases learned in the “Normalization of targeted DNase Hi-C data” section to calculate the contact probability between each pair of contact bins. Fit-Hi-C first performs the spline fitting on the raw contact counts to estimate the genomic distance effect. Then for any given pair of contact bins with genomic distance d, the contact probability (i.e., the binomial parameter) is calculated by multiplying the prior contact probability obtained from the spline $p_{raw} = spline(d)$ with the corresponding bias factors at the two contacting bins.

For short-range (<10 Mb) intrachromosomal chromatin contacts, we first parsed the target-captured reads at 1-kb resolution and then applied the modified Fit-Hi-C to estimate the null distribution of contacts within the genomic distance range of 5 kb–10 Mb. We discarded short-range contacts (<5 kb) because they are mainly self-ligation products (Supplementary Table 24). We used one round of refinement (i.e., two rounds of spline fitting) to estimate this null distribution and then identified the significant contacts at false discovery rate (FDR) <0.05. For long-range (≥10 Mb) intrachromosomal chromatin contacts, we parsed the target-captured reads at 10-kb resolution and then used the modified Fit-Hi-C to estimate the null distribution of contacts within the full range of full chromosome length using one round of refinement. Similarly to short-range contacts, we identified the significant contacts at FDR <0.05. For interchromosomal chromatin contacts, we identified significant target-captured contacts at 10-kb resolution using a simple binomial model, as described in Duan et al. and used an FDR <0.05. To eliminate contacts that are introduced by mapping biases at low-mappability regions, we further discarded contacts that were associated with genomic bins that have low mappability score (<0.5). A mappability score of 0.5 for a bin means that half of the base pairs in that bin are not uniquely mappable for reads of the length at which mappability is calculated. We calculated mappability scores using GEM46 using a 50-bp read length.

Neighborhood effect filtering. In the event of a bona fide chromatin looping contact, we expect the immediately flanking bins around the contacting regions to be also within relatively close proximity (Supplementary Fig. 6c). Thus, we applied a neighborhood filter using the contact significance we computed from Fit-Hi-C as follows. For each chromatin contact between target i and nontarget genomic bin j, we call it a high-confidence contact if the contact itself meets the stringent FDR cutoff of 0.05, and at least three out of ten neighboring bins (five on
each side) of bin $i$ contact the target $t$ at a permissive FDR cutoff of 0.1. After the neighborhood filtering, we merged adjacent and nearby (<3 bins apart) high-confidence contact bins associated with the same target.

**Comparison with ChIA-PET data.** To validate the targeted DNase Hi-C method, we compared the high-confidence contacts identified by targeted DNase Hi-C in K562 cells to those that were identified by RNAP II–mediated and CTCF-mediated ChIA-PET data generated by the ENCODE Consortium. We extracted intrachromosomal ChIA-PET contacts for which one end falls within the designed target regions and the other end falls in mappable regions (mappability score ≥0.5) within 10-Mb genomic distance. For each target region, we calculated the overlaps of the intrachromosomal high-confidence contacts identified by targeted DNase Hi-C and by ChIA-PET. Statistical significance of the observed overlap was calculated using a hypergeometric test.

**Comparison with 5C data.** To validate the targeted DNase Hi-C method, we compared the high-confidence contacts identified by targeted DNase Hi-C in H1 and K562 cells to those that were identified by 5C studies by the ENCODE consortium. The 5C experiments assess interactions among 44 ENCODE pilot regions using two separate 5C primer pools. We extracted intrachromosomal 5C contact peaks for which one end falls within the designed target regions and the other end falls in mappable regions (mappability score ≥0.5) within 10-Mb genomic distance. For each target region that overlaps with a 5C primer, we calculated the overlaps of the intrachromosomal high-confidence contacts identified by targeted DNase Hi-C and by 5C. Statistical significance of the observed overlap was calculated using a hypergeometric test.

**Enrichment of intradomain targeted DNase Hi-C contacts.** For targeted DNase Hi-C, we label a contact as an intradomain contact if both the captured target region and the partner locus lie within the same topological domain. To test whether the short-range (<10 Mb) high-confidence intrachromosomal contacts identified by targeted DNase Hi-C are enriched for intradomain contacts, we computed the ratio $R$ between the number of high-confidence contacts that have both ends within one topological domain (intradomain) to the number of contacts that occur across two different domains (interdomain). To estimate the significance of the ratio $R$, we randomly shuffled topological domains by preserving the distribution of the domain lengths for each chromosome. We achieved this as follows. Excluding chromosome ends, if there are $n$ domains for a chromosome then there will be $m = n - 1$ boundaries. Note that boundaries are gapped regions between adjacent domains, not single points. To construct our null model, for each chromosome, we separately shuffled the domains and the boundaries and then interleaved the two shuffled lists to build the permuted domain structures on that chromosome. We did this randomization for each chromosome and repeated the process 1,000 times to create a null model. We then computed the average and s.d. of $R$ over all randomizations. We assumed the test statistic $R$ followed normal distribution and assessed the statistical significance of the observed intradomain enrichment over the enrichment gathered from 1,000 randomizations.

**Enrichment of genomic features.** To evaluate the association between a set of regions with a given genomic feature $f$ and the high-confidence contacts identified by targeted DNase Hi-C, we applied the Genome Structure Correction (GSC) test. The tested features include DNase I open chromatin regions, super-enhancers and more. Each feature was compared to the set of loci that participate in a significant contact with a target region (i.e., target partners). The GSC method estimates the statistical significance of the observed overlap between target partners $g$ and the given genomic feature $f$ using a block subsampling approach. More specifically, GSC iteratively subsamples two large, equal-sized blocks $A$ and $B$ from the genome and identifies the subsets of feature $f$ in the two blocks ($f_A$ and $f_B$, respectively) as well as the subsets of contact partners in these two blocks ($g_A$ and $g_B$, respectively). To estimate the empirical null, GSC swaps feature subsets $f_A$ and $f_B$ in two blocks and calculates the expected overlaps. In other words, the expected overlaps are estimated by calculating the overlaps between $f_A$ and $g_B$, and between $f_B$ and $g_A$. This block subsampling procedure is repeated multiple reads to build the empirical null. The GSC method has been shown to be suitable for genome-scale feature enrichment tests. Other randomization experiments, such as random shuffling or shifting, do not preserve local genome properties or spatial relationships of the features and therefore tend to underestimate the null and overestimate the significances of the overlaps.

In our comparisons, we focused only on target partners that fell within 10-Mb distance from the corresponding target regions. Thus, we performed the GSC test on these restricted regions to avoid underestimation of the null. The details of the GSC test procedure is described as follows.

1. For a given targeted DNase Hi-C library, we defined the eligible genomic bins (at 1-kb resolution) as those that are within 10-Mb distance from at least one target lincRNA promoter region and have a mappability score ≥0.5.
2. For each chromosome, we concatenated all eligible bins on the chromosome to generate an artificial chromosome.
3. We transformed the original genomic coordinates of the high-confidence target partners to those in the artificial genome.
4. Similarly, we transformed the coordinates of the genomic feature to the new ones in the artificial genome.
5. We ran GSC on the artificial genome with the following parameters: region fraction (-r) 0.3, subregion fraction (-s) 0.3, subsampling number (-n) 10,000, statistical test (-t) region overlap marginal.

For each genomic feature of interest, we performed two reciprocal GSC enrichment tests separately. The first tests whether the given feature is enriched within the target partners. The second is to test whether the target partners are enriched within the genomic feature. The GSC software reports a Z score and P value for each enrichment test.

We applied the GSC procedure to five types of genomic features: ENCODE genome-wide segmentations, DNase I open chromatin regions, FAIRE-seq open chromatin regions, super-enhancers and transcription factor binding peaks. Details for each of these genomic features are as follows.

1. We used the 25-state whole-genome segmentation generated by Segway. First, we aggregated the 25 segmentation labels into eight groups on the basis of their associated genomic and epigenomic properties. The eight groups are...
promoters, poised promoters (for H1 cells only), enhancers, transcribed regions, FAIRE-only open chromatin (FAIRE-only regions are FAIRE peak regions excluding promoters, enhancers, CTCF binding sites and DHSs), CTCF distal elements (CTCF), repressed regions and quiescent chromatin (Dead). We then ran GSC to estimate the statistical significance of the observed overlaps between the target partners and each label group.

2. For DNase I open chromatin regions, we downloaded the DNase I peak regions generated using DNase-seq by the ENCODE consortium29 (uniform peak calls, Jan/2011 freeze, http://ftp.ebi.ac.uk/pub/databases/ensembl/encode/integration_data_jan2011/byDataType/openchrom/jan2011/idrPeaks).

3. For FAIRE-seq open chromatin regions, we downloaded the FAIRE-seq peak regions generated by the ENCODE consortium29 (uniform peak calls, Jan/2011 freeze, http://ftp.ebi.ac.uk/pub/databases/ensembl/encode/integration_data_jan2011/byDataType/openchrom/jan2011/faire_fseq_peaks).

4. We downloaded the coordinates of super-enhancers in H1 ESC cells and in K562 cells from Hnisz et al.31.

5. We obtained regions of TF binding peaks identified by ChIP-seq from the ENCODE consortium29. We downloaded the file named “wgEncodeRegTfbsClusteredWithCellsV3.bed.gz”, from http://hgdownload-test.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeRegTfbsClustered, which contains the lists of TF binding peak clusters identified by combining data from 91 cell types and 189 TF targeting antibodies. From this full list, we extracted the peaks for H1 ESC and K562 cells that have 50 and 100 TFs in H1 and K562 cells, respectively, including 36 TFs whose binding sites have been mapped in both cell lines. We performed the GSC enrichment test for each TF in each cell type in which its binding sites were mapped.

**lincRNA expression analysis.** To estimate expression profiles of lincRNAs, we analyzed the poly(A)++ long RNA-seq data sets generated by the ENCODE consortium29. We used GENCODE v7 data27 to annotate the transcriptome, which included both protein-coding genes and noncoding RNAs. We mapped RNA-seq reads using TopHat v2.0.0 (ref. 49) to both the human genome and transcriptome with default parameters. Then we measured the expression of genes and lincRNAs as fragments per kilobase of exon length per million mappable reads (FPKM) using Cufflinks v2.0.2 (ref. 50). We analyzed biological replicates separately and averaged FPKMs from two biological replicates to obtain the final expression level. We labeled a lincRNA as “expressed” if its expression level is greater than the median expression of all lincRNAs.

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