Physical interactions between distal regulatory elements have a key role in regulating gene expression, but the extent to which these interactions vary between cell types and contribute to cell-type-specific gene expression remains unclear. Here, to address these questions as part of phase III of the Encyclopedia of DNA Elements (ENCODE), we mapped cohesin-mediated chromatin loops, using chromatin interaction analysis by paired-end tag sequencing (ChIA-PET), and analysed gene expression in 24 diverse human cell types, including core ENCODE cell lines. Twenty-eight per cent of all chromatin loops vary across cell types; these variations modestly correlate with changes in gene expression and are effective at grouping cell types according to their tissue of origin. The connectivity of genes corresponds to different functional classes, with housekeeping genes having few contacts, and dosage-sensitive genes being more connected to enhancer elements. This atlas of chromatin loops complements the diverse maps of regulatory architecture that comprise the ENCODE Encyclopedia, and will help to support emerging analyses of genome structure and function.
according to their tissue group of origin. The groupings are concordant across gene regulatory phenotypes, suggesting that loop variation recapitulates cell-type identity in a similar manner to enhancer activity and gene expression, and is mainly driven by epigenetic factors.

- We found that approximately one-quarter of cohesin-mediated chromatin loops varied across cell types, showing substantial variability in interactions at the sub-TAD scale. Variable loops tend to span shorter distances and are depleted in housekeeping genes.

- Approximately one-quarter of cohesin-mediated loops are anchored by enhancers across diverse cell types, representing the most enriched loop-associated chromatin state. Enhancer anchors participate in more interactions than promoter anchors and are enriched for interactions with other enhancers and transcription start sites (TSSs), consistent with groups of enhancer-associated loops regulating promoters.

- Cell-type-specific loops coincide with different chromatin states. For example, stem cell loops show reduced active promoter and transcribed states, and increased bivalent states, which may point to a role of these loops in maintaining pluripotency.

- In our interaction map, genes that have more interactions are depleted for housekeeping functions and enriched for pathogenic variants and haploinsufficiency, suggesting that the connectivity of a gene is linked to its function and role in disease.

- Loop variation modestly correlates with gene expression variation, especially for loops that link an enhancer directly to a promoter; a weaker positive correlation is observed for genes internal to loops and for neighboring genes within the same loop.

- Group-specific (blood and embryonic) loops show enrichment of cell-type-specific transcription factor (TF) motifs at loop ends and are enriched in genes with group-specific functions. Genome-wide association study (GWAS) variants for autoimmunity are enriched in blood-specific loops, but not in embryonic loops, pointing to the importance of cohesin-mediated loops for understanding the mechanisms of human disease variants.

**Genome-wide map of chromatin interactions**

To identify cell-type-specific chromatin loops on a genome-wide scale, we generated 3D chromatin interaction maps at single-cohesin peak resolution (about 2 kb) using a modified ChIA-PET assay (Extended Data Fig. 1a, Methods). In brief, this chromatin conformation capture assay incorporates an immunoprecipitation step followed by proximity ligation to measure the frequency of interactions between pairs of genomic regions bound by a protein of interest. We chose the RAD21 subunit of the cohesin complex, which facilitates physical contacts between genes and enhancers, and is essential for chromatin loop assembly and subsequent TAD formation. Henceforth we refer to these cohesin-mediated chromatin loops as loops or interactions. We generated a median of about 200 million paired-end reads (2 × 101 bp) per experiment (Extended Data Fig. 1b, Supplementary Table 2). To study the interplay between loops, regulatory elements, and gene expression, we also generated chromatin immunoprecipitation with sequencing (ChiP–seq) data for the histone mark H3K27ac, which demarcates active promoters and enhancers, and paired-end RNA sequencing (RNA-seq) data (Supplementary Table 2). The ChiP efficiency for RAD21 and H3K27ac passed ENCODE ChiP–seq quality standards (Extended Data Fig. 1c, d). All experiments were performed in biological replicates.

To define a comprehensive, high-resolution set of chromatin loops, we pooled ChiA-PET data sets across all 24 cell types in our study (representing about 10 billion reads) and called a unified set of interactions using different 3C-based assays, such as in-situ Hi-C, ChIA-PET and Hi-ChIP (Methods). We generated a median of about 200 million paired-end reads (2 × 101 bp) per experiment (Extended Data Fig. 1b, Supplementary Table 2). To study the interplay between loops, regulatory elements, and gene expression, we also pooled ChiA-PET data sets across all 24 cell types in our study (representing about 10 billion reads) and called a unified set of interactions using different 3C-based assays, such as in-situ Hi-C, ChIA-PET and Hi-ChIP (Methods).

**Loop variability and cell type**

Inspection of our data set revealed two broad classes of loops—those in which the normalized interaction frequencies (the number of paired-end tags (PETs) that link the two ends of a loop) varied across cell types and those that were relatively non-variable (Fig. 1a). For example, both DPPA4 and DPPA2 were entirely contained within two loops in the stem cell lines that we used (H1-hESC, H9-hESC, and MSiPS); however, these loops were either absent (for example, in GM12878 and MSLCL cells) or displayed reduced interaction frequency in a number of cancer cell lines (for example, Jurkat and K562 cells). Consistent with this observation, both...
Fig. 2 | Chromatin loop variation across 24 cell types. a, Examples of variable (left) and non-variable loops (right) across cell types. Chromatin loops are displayed above the corresponding RAD21 signal tracks. The colour density of loops corresponds to normalized interaction frequency (darker blue indicates higher frequency). b, PCA of normalized chromatin loop interaction frequencies (n = 85,294 loops versus n = 48 samples (24 cell types × 2 replicates each)). Colours denote the germ layer origin of each sample (Supplementary Table 5). c, Correlation of interaction frequencies between pairs of cell types (all types, n = 1,104 pairs; isogenic, n = 15; germ layer, n = 316; tissue, n = 160; biological replicates, n = 24; P values calculated using a two-sided Wilcoxon rank-sum test). Centre line represents the median, box extent ranges from 25th to 75th percentile, and whiskers extend to most to 1.5× the interquartile range. Summary statistics for the boxplots can be found in Supplementary Table 9. d, Size distribution of variable chromatin loops versus two different sets of non-variable control loops (n = 35,698, significance assessed using two-sided t test). Centre line represents the median, box extent ranges from 25th to 75th percentile, and whiskers extend to most to 1.5× the interquartile range. Summary statistics for the boxplots can be found in Supplementary Table 9. e, Overlap of variable and non-variable chromatin loops with contact domains. f, Enrichment of cell-type-specific genes and depletion of housekeeping genes (n = 2,220) in variable versus non-variable loops (n = 35,698). P values calculated using a two-sided Fisher’s exact test. Summary statistics for the figure can be found in Supplementary Table 9.

DPPA2 and DPPA4 are active during development29 and have been implicated in cancer30.

We sought to use our measurement of interaction frequencies to systematically identify variable loops across different cell types. First, we subjected normalized interaction frequencies across all cell types to principal component analysis (PCA) (Fig. 2b). All cell types fell into one of three main clusters—blood, stem-cell like (embryonic), and mesoderm—whereas isogenic cell types (i.e., cell lines derived from the same donor) were more dispersed across the clusters (Extended Data Fig. 2c). Principal component analysis (PCA) of the normalized interaction frequencies showed a greater degree of enrichment for variable loops in the embryonic and mesoderm-like clusters (Extended Data Table 2).

Having established that normalized interaction frequencies could reliably group related cell types, we next sought to quantify loop variability. We used a linear mixed effects model to identify loops that varied in interaction frequency across our set of 24 cell types (Methods). To test for variability, we filtered loops to include only those with four or more PEs in at least one sample, which yielded 85,294 loops (Supplementary Table 5). At FDR < 10%, we identified 35,698 variable loops, or 41% of all tested loops (28% of the pan-cell line loop set) (Extended Data Table 5, Methods). Variable loops spanned significantly shorter distances than non-variable loops (130 kb versus 178 kb) (Fig. 2d, Extended Data Fig. 2g, h). Variable loops also tended to overlap contact domain boundaries to a lesser degree than non-variable loops (Fig. 2e, Methods).

Finally, we examined whether loop variability is associated with specific types of genes. Using our RNA-seq data, we defined genes as broadly expressed or cell-type-specific (Methods). Overall, variable loops showed enrichment for cell-type-specific gene expression relative to non-variable loops, whereas they were depleted in genes that are expressed across all cell types (Fig. 2f, Extended Data Fig. 2i). In agreement with this observation, non-variable loops were enriched in genes that are broadly expressed across a larger set of tissue types24,34 (Extended Data Fig. 2j).

Cell-type-specific loops and chromatin states

Enhancers often exert their influence on gene expression over large distances through direct 3D chromatin contacts with multiple distal promoters29–31. To study the subset of cohesin loops that mediate enhancer contacts, we profiled the enhancer mark H3K27ac using ChIP–seq and quantified signal at 288,711 genomic regions that were enriched for enhancer activity in at least two cell types (‘enhancers’) (Supplementary Table 5, Methods). Loop ends from our pan-cell-type data set showed increasing overlap with enhancer regions the more interactions they were involved in; the same was observed for contact domain boundaries (Fig. 3a). On the other hand, loop ends with few interactions tended to coincide with promoters (this result was robust to the threshold used to define the pan-cell-line loop set (Methods)). Together, these data are consistent with a ‘hub and spoke’ model in which groups of enhancers work together through cohesin-mediated looping to target and regulate multiple promoters34.

Manual inspection of our data indicated that cell-type-specific loops tended to overlap with cell-type-specific regulatory elements, such as enhancers (Fig. 3b). Thus, we investigated which chromatin states overlapped cell-type-specific loop ends. To this end, we obtained chromatin state calls from the Roadmap Epigenomics Mapping Consortium30 for 12 cell types (Supplementary Table 1). Next, for each cell type, we identified a set of cell-type-specific interactions—loops with high interaction frequencies in the cell of interest, but reduced frequencies in all other cell types (Methods). Finally, we tabulated the number of chromatin state elements across eight categories that overlapped each set of cell-type-specific loop ends (Methods). As in the pan-cell-type data set, in nearly all cell types, genomic elements in the enhancer state (ENH) represented a large proportion (about 23%; Fig. 3c, Extended Data Table 3).
Next, we investigated whether cell-type-specific cohesin-mediated chromatin loops might specifically connect cell-type-specific enhancers and expressed genes. Similar to past studies that have approached this question using a promoter-centric view, we observed strong enrichment for enhancer–promoter (ENH–TSS) interactions (Fig. 3e). In addition, interactions between enhancers (ENH–ENH), but not between promoters (TSS–TSS), were enriched. Studies in different systems have shown that the number of enhancers linked to a given promoter is associated with the RNA expression level. We looked for this effect in promoters that are linked to enhancers by cohesin-mediated chromatin loops, by binning genes on the basis of the number of linked enhancers; the number of enhancers was modestly but significantly correlated with expression level (Fig. 3f), suggesting that the recruitment of additional physically linked enhancers may help to regulate gene expression. Again, these results were robust to the choice of threshold used to define cell-type-specific interactions (Methods).

**Gene connectivity corresponds to function**

We next investigated whether the number of physically interacting enhancers could be related to the basic properties of a gene. We hypothesized that genes that encode products with effects that depend strongly on their levels of expression (‘dosage-sensitive’ genes) would have more enhancer contacts to support a more robust regulatory architecture than other genes. To test this idea, we obtained a list of genes that were annotated as haploinsufficient (such that loss of one copy leads to pathogenicity) or haploinsufficient genes were enriched among genes with a higher number of loops, enhancers, and cell-type-specific enhancers connected to them (Fig. 3g, Extended Data Fig. 4a, b, Supplementary Table 9), suggesting a link between the dosage pathogenicity of a gene and increased regulatory contacts. We next tested whether other categories of human disease-related genes also tended to have more distal contacts. Analogous to the case for haploinsufficient genes, we find that genes identified as being disease-associated in GWASs tended to be more highly connected to distal regions, including enhancers (Extended Data Fig. 4b, f). The same was true for genes with a reported pathogenic or likely pathogenic variant in ClinVar (Fig. 3h, Extended Data Fig. 4c, g), indicating that genes associated with both common and rare human diseases possess more extensive regulatory wiring than other genes. By contrast, housekeeping genes, which we defined as being broadly expressed among our 24 cell types (Methods), were depleted from genes with higher numbers of loops (Fig. 3i, Extended Data Fig. 4d, h). This is consistent with the finding that housekeeping genes active during mouse development have a median of zero enhancers. Together, these results indicate that genes for which misregulation makes an organism particularly vulnerable have a complex regulatory architecture that may ensure correct expression through the redundancy or fine-tuning of regulatory interactions.

**Loop interaction frequency and gene expression**

We next investigated the extent to which changes in loop interaction frequency correspond to changes in gene expression. For example, we observed physical interaction between a distal enhancer and the promoter region of the gene MTDH, a known oncogene that activates the NFkB pathway. This interaction was frequent in blood cell lines (for example, GM12878, MSLCL, and SU-DHL-2), and accompanied by higher expression of MTDH RNA. Conversely, in cell types where looping with the promoter region was reduced or absent (for example,
Chromatin loops regulate alternative splicing

A recent report\textsuperscript{31} found a link between intragenic CTCF-mediated chromatin loops and alternative splicing within the same cell type across individuals. To assess whether a similar mechanism might drive cell-type-specific isoform use, we identified 1,372 loops associated with 1,074 genes that linked the promoter and the gene body (Extended Data Fig. 5a, b). Loop strength showed the highest correlation with differentially used exons (DUEs) (Methods) that were next to the loop anchor when comparing the normalized signal across all cell types (Fig. 4h) and also when using fold change for a specific lineage (Fig. 4i). These results suggest that the presence of an intragenic loop can affect the inclusion of the exon next to it, as exemplified by ARHGEF7, which selectively includes exon 6 in cell types where the loop is present (Fig. 4j).

Consequently, we observed a high correlation between loop strength and exon abundance ($R = 0.49$) (Extended Data Fig. 5c, d).

Group-specific loops

Clustering of interaction frequencies across the genome revealed three distinct cell type clusters—blood, embryonic, and solid-tissue-derived (Fig. 2b). We next sought to identify and characterize loops that were present in each group. We rank-ordered all loops tested for variability according to the extent to which their interaction frequency was elevated in cell types in one group compared to the other two (Methods). We hypothesized that group-specific loops were relevant to the determination and maintenance of cell fate, which are governed by cell-type-specific TFs acting on proximal and distal regulatory elements. To examine enrichment of TFs at loop ends, we intersected each set of group-specific loops with motif positions for 598 TFs\textsuperscript{11,13}.

Among the most significantly enriched TFs at blood-loop anchors were haematopoietic TFs involved in lymphoid B- and T-cell development, such as SPIB\textsuperscript{55}, SPIJ/PJU.\textsuperscript{11}, TCF3 (which is mutated in most Burkitt lymphomas\textsuperscript{50}), and ZBTB7A (which is recurrently mutated in acute myeloid leukaemia\textsuperscript{50}) (FDR < 5\%)(Fig. 5a, Methods). At the embryonic-specific loop anchors, we found modest enrichment for a smaller set of TFs, among them PKNOX1 (which regulates haematopoietic stem and progenitor cell activity\textsuperscript{19}) and PKNOX2 (which is essential for limb development\textsuperscript{59}) (FDR < 5\%)(Extended Data Fig. 6a). Our results were fairly robust to the choice of threshold used to define group-specific loops (Methods).

To confirm the motif analysis, we reasoned that functional motifs are more likely to be present in open chromatin and thus we expected to see enrichment of chromatin accessibility signal. Indeed, the motif instances for the haematopoietic TFs SPIB, SPIJ/PJU.1, and TCF3 coincided with higher chromatin accessibility at blood-specific loop anchors than in the all-loops, as measured by assay for transposase-accessible chromatin using sequencing (ATAC-seq) in the blood lineage cells (Fig. 5b–d, Methods). This supports the notion that haematopoietic TFs are probably more active in blood-specific loops.

Next, we investigated whether cell-type-specific loops are associated with cell-type-specific biological processes, which would indicate that cohesin loops are integral to cell-type-specific transcriptional programs. Loops that were present more frequently in blood cell types than
in other cell types were enriched for genes involved in leukocyte activation ($P < 10^{-15}$), mature B cell activation ($P < 10^{-20}$), and numerous other immune-related categories (Fig. S5, Supplementary Table 6, Methods). Loops that were gained in the embryonic group were enriched for genes with a more complex pattern of functional categories, including differentiation and morphogenesis (Extended Data Fig. 6b). These results suggest that the cell-type-specific differences in chromatin looping are likely to be functionally important.

Finally, we investigated to what extent disease-associated variants identified by GWASs tend to occur at loop ends. To this end, we intersected GWAS single-nucleotide polymorphisms (SNPs) from 86 traits with each set of group-specific loop ends (Methods). Similar to the GO enrichment analysis, we identified enrichment of GWAS SNPs for different sets of diseases in each set of group-specific loops (Fig. S5, Extended Data Fig. 6c). The blood-specific loops were enriched for SNP associations with autoimmune diseases, including multiple sclerosis, coeliac disease, and Crohn’s disease (Fig. S5). Lipid-associated traits (for example, LDL, HDL, and total cholesterol) were also enriched. These enrichments were not significant in the embryonic-specific loops (Extended Data Fig. 6c); we confirmed this result through a relative enrichment test, which directly compared each set of group-specific loops (Supplementary Table 7, Methods). The most significantly associated traits identified for embryonic-specific loops were fasting insulin, serum creatinine, and height (Extended Data Fig. 6c), the latter of which has previously been associated with stem-cell-specific chromatin patterns.

We confirmed the GWAS enrichments we observed using GRASP overlaps by a complementary approach, partitioned linkage disequilibrium (LD) score regression, using a common set of 47 traits. Similar traits were enriched in the blood-specific group of loop anchors, including HDL cholesterol and autoimmune-related diseases, along with a number of blood cell traits (Fig. S5g). In embryonic lineages, the only significantly enriched trait at FDR = 1% was ‘years of education’, which is thought to be driven by brain associations and was also consistent with the embryonic and neural progenitor populations in this group (Extended Data Fig. 6d). We also examined the sensitivity of the LD score regression approach to additional corrections for underlying genomic features, for example, super enhancer annotations, and observed that the overall trends of enrichment remained consistent (Extended Data Fig. 7, Supplementary Table 8, Methods).

Together, our results suggest that distinct sets of TFs may help to facilitate cell-type-specific loops, which in turn contain functionally related genes that are critical to the function of each particular cell type, emphasizing the important role of chromatin loops in human traits and suggesting that analysis of the effects of genetic variants in these regions may provide mechanistic insights into disease.

**Discussion**

We have generated one of the most comprehensive 3D chromatin interaction data sets to date, spanning 24 cell types. Owing to the high reproducibility of our data, we were able to identify loops whose interaction frequencies varied across our panel of cell types. About 28% of loops genome-wide varied significantly among cell types and were associated with cell-specific differences in gene expression. The differences in gene expression associated with loop variation are relatively modest, perhaps suggesting that not all varying chromatin loops have functional consequences or that a subset of variable loops may be poised to alter gene expression in specific developmental or physiological contexts. Notably, we found that neighbouring genes, which tend to be on average more co-expressed than non-neighbouring genes, showed more strongly correlated gene expression when contained within the same chromatin loop than when they were located next to each other but did not share a loop. Together, these results indicate that chromatin looping has a role in regulating gene expression, and point to the ability of loops to coordinate the expression of functionally related sets of genes, such as pathways or protein complexes.

The diverse cell types we studied clustered into three main groups—blood, embryonic, and derived from solid tissues—based on shared commonalities in cohesin-mediated chromatin looping. Analyses from both the GTEx Consortium and Roadmap Epigenomics Project—which profiled dozens of tissues and cell types for gene expression and histone modifications, respectively—showed very similar grouping, indicating that both blood and embryonic cell types are likely to have gene regulation programs that differ strongly from those of solid tissues. To our knowledge, this is the first time that quantitative measurements of cohesin-mediated chromatin loops have also been shown to correspond to cell-type identity.

Our results build on previous ENCODE work that has shown that gene architecture is highly variable throughout the human genome. Notably, we have shown that the extent of long-range contacts of a gene correlates with its function and role in human disease. Genes with few contacts are enriched in housekeeping genes, which could reflect simpler circuitry for constitutive, steady expression. Highly connected genes are more strongly associated with both common and rare classes of human diseases, as demonstrated by their enrichment in ClinVar and GWAS genes and in genes that cause disease when haploinsufficient. These observations may indicate that one function of a more extensive regulatory architecture is to safeguard the expression of dosage-sensitive genes.

Maps of 3D chromatin interactions have become increasingly useful in explaining how distal regulatory elements can exert their influence. Here, we have demonstrated how knowledge of cell-type-specific interactions can further expand the utility of such maps. For example, we found that GWAS SNPs were enriched in loops observed in cell types that have been shown to be relevant to a particular disease.
These findings suggest that our data set could have multiple future applications. GWAS and cancer genomics studies continue to deposit disease-related sequence variations into public databases, and most of these variants fall into non-coding regions. As we have demonstrated here, intersecting these variants with cell-type-specific chromatin loops may help to explain how such sequence variation leads to disease.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-020-2151-x.

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**Methods**

**Cell lines**

The cell types and lines in this study were either obtained from cell repositories or established or differentiated in the Snyder and Dalton laboratories at Stanford University and the University of Georgia, respectively (Supplementary Tables 1, 2). All tissue culture was done according to the manufacturer’s recommendations. One of the commercially available cell lines, K1 (thyroid, papillary carcinoma), is on the list of commonly misidentified cell lines (ICLAC). The relevant cell line (CVCf-9918) was also derived from a thyroid papillary carcinoma. In the event of misidentification, the conclusions of our study would not be affected because both cell lines represent papillary thyroid carcinoma.

**ChIA-PET experiments**

We performed ChIA-PET experiments with modifications to previously published protocols. These modifications have also been independently described. We used Illumina’s Nextera tagmentation to generate sequencing libraries. In brief, cells were crosslinked and subjected to nuclear lysis followed by chromatin shearing (no restriction enzyme was used). Immunoprecipitation was performed overnight at 4°C with antibodies against the cohesin subunit RAD21 (Abcam Anti-RAD21 antibody (ab992) https://www.encodeproject.org/antibodies/ENCAB5297YRC/). The immuno-complexes were pulled down with Protein-G dynabeads (Life Technologies #11205D, New York). Biotinylated linkers were ligated to the enriched fragments, followed by proximity ligation overnight at 16°C.

Crosslinking was reversed at 65°C with the use of Proteinase K followed by DNA purification. We used Illumina Nextera Transposase to add sequencing adapters to ChIA-PET libraries. Biotinylated fragments were enriched by pull-down with Streptavidin Dynabeads (M-280; Lifetechnologies #11205D, New York). Biotinylated linkers were ligated to the enriched fragments, followed by proximity ligation overnight at 16°C.

**ChIP–seq experiments**

Chromatin immunoprecipitation followed by massively parallel sequencing was carried out as previously described. Cells were crosslinked with formaldehyde at a final concentration of 1% for 10 min at room temperature. The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature. The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature). The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature). The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature). The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature). The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature). The reaction was quenched with glycine at a final concentration of 125 mM and nuclear lysates were sonicated using a Branson 250 Sonifier (power setting 2, 100% duty cycle for 7 min at room temperature).

**RNA-seq experiments**

RNA samples were extracted using the Qiagen All-Prep kit, following the manufacturer’s instructions. Libraries were prepared from total RNA using the TruSeq Stranded Total RNA Library Prep Kit, following the manufacturer’s instructions. All libraries were sequenced on the Illumina HiSeq 4000.

**ATAC–seq experiments**

ATAC–seq was carried out as previously described and sequencing was carried out on an Illumina HiSeq 2000 with 2 × 100 paired-end sequencing.

**ChIA-PET processing pipeline**

ChIA-PET data were generated in replicate for all 24 cell lines; all libraries were sequenced to an average depth of 214 ± 5.5 (mean ± s.d.) million paired-end reads (referred to as paired-end tags or PETs) (Supplementary Table 2). Data were processed in a similar way to the workflow used in the Mango toolkit, as follows.

**Trim adaptor sequences.** Illumina Nextera adaptor sequences (CT- GTCTCTTATA and TATAAGAGACAG) were trimmed from all PETs using cutadapt in paired-end mode (version 1.11; non-default parameters: -q 15 -O 4 -m 20).

**Trim linker sequence.** All PETs were scanned to identify and remove the linker sequence (GTTGGATAAG), as well as any sequences downstream of the linker sequence. PETs less than 20 bp in length after linker removal were discarded.

**Align paired-end sequences.** Each set of paired-end reads was aligned to the hg19 genome separately using bowtie (version 0.12.8; non-default parameters: -n 2 -l 50 -k 1 --mapq 40 --best -m 1). Paired-end reads that mapped to multiple locations were discarded.

**Remove duplicate paired-end sequences.** PETs that mapped to identical locations were filtered to retain only a single PET.

**Generate a set of unified peak calls.** For each sample, the two sets of uniquely mapped paired-end reads were merged and peaks were called using MACS2 (version 2.1.1.20160309; parameters: -g hs -f BED -q 0.01). Peak calls across all samples were combined and then extended by 500 bp in either direction. Overlapping peaks were merged to form a single interval that spanned all overlapping peaks, after which peaks in ENCODE-defined blacklist regions were filtered. In total, we obtained 286,620 RAD21 peaks (Supplementary Table 3). These merged peak regions were used as our ‘anchor regions’ for all subsequent analysis.

**Generate a set of linked paired peaks.** For all pairs of peaks that were >10,000 bp and <5,000,000 bp apart on chr1-22 and chrX, the total number of PETs that linked each pair was tabulated. For samples with >2,250,000 unique PETs, the total number of PETs was down-sampled to 2,250,000 before any further analysis.

Our final data set consisted of a matrix, M_{ij}, in which each row (i) represents a single paired peak, and each column (j) represents a single sample. Element m_{ij} indicates the number of PETs linking the two anchor regions. We normalized the data by standardizing each row in M_{ij}, and then quantile-normalizing the columns. The range of values in each column was then re-scaled to between 0 and 1000.

**Generating the pan-cell line loop-call data set**

Unique PET data (that is, data from ‘Remove duplicate paired-end sequences’ in the ChIA-PET processing pipeline above) from all cell lines and all replicates were pooled together. Next, we tabulated the number of PETs that connected all pairs of anchor regions >10 kb and <5 Mb apart in our unified peak set (Supplementary Table 3). Finally, the Mango scoring methodology was used to assign each peak pair a P value; Mango uses a Bayesian scoring methodology to determine the expected number of PETs connecting any two regions on the basis of the distance between the two regions and the local ChIP-efficiency. We used a threshold of \( P < 2.3 \times 10^{-7} \) to arrive at our pan-cell line loop set (Supplementary Table 4). We used a relatively stringent cutoff due to the large number of PETs being analysed. At this cutoff our FDR was \( 2.7 \times 10^{-10} \) using the Benjamini–Hochberg procedure and 0.013 using the Bonferroni approach. For all subsequent analysis described below, we used the FDR estimate from the Benjamini–Hochberg procedure.
RNA-seq data were generated in replicate for 22 out of 24 cell lines (Supplementary Table 2); we obtained on average 43 ± 9 million paired-end reads per sample (mean ± s.d.). For samples with >60 million reads, FASTQ files were down-sampled to 60 million reads before further analysis. Transcript abundances were quantified using kallisto (version 0.43.0; non-default parameters: -bias). Transcript sequences (that is, target sequences) were obtained from Gencode (release 25; lifted to GRCh37 coordinates). Duplicate transcripts were removed, as well as transcripts not classified as ‘protein coding’ or ‘lncRNA’, yielding a final list of 93,430 transcripts. For all analyses, we considered only 69,598 transcripts with a maximum abundance of >1 transcripts per million (TPM) across all 23 cell lines. To produce gene-level estimates of expression, we summed the TPM values for all transcripts that belonged to the same gene. For all analyses, we considered only 22,197 genes with a maximum abundance of >1 TPM across all 23 cell lines. For GM12878 cells, we used data from a previous study. To normalize RNA-seq data, we first standardized (that is, z-score scaled) TPM values for each transcript or gene across all cell lines and then quantile-normalized all transcript or gene abundance levels between samples.

To visualize RNA-seq data as signal tracks, down-sampled FASTQ files were aligned to the hg19 genome using Hisat2 (version 2.0.5; non-default parameters: -q 15 -O 4 -m 20). Reads were aligned using bowtie (version 0.12.8; non-default parameters: -X 2000 -m 1 -fr -no-mixed -no-discordant) after which genome-wide coverage tracks were produced using bedtools (bedtools genomewc -bga -split -ibam). Coverage values were scaled by a constant factor (107/total number of reads) to account for differences in sequencing depth.

**H3K27ac ChIP–seq data processing**

ChIP–seq data were generated in replicate for 22 out of 24 cell lines (Supplementary Table 2); we obtained on average 43 ± 9 million paired-end reads per sample (mean ± s.d.). Illumina TruSeq adaptor sequences were trimmed using cutadapt in paired-end mode (non-default parameters: -q 15 -O 4 -m 20). Reads were aligned using bowtie (version 0.12.8; non-default parameters: -X 2000 -m 1 -fr -no-mixed -no-discordant) after which genome-wide coverage tracks were produced using Picard MarkDuplicates. Finally, peaks were called using MACS2 (default parameters).

**ATAC–seq data processing**

ATAC–seq data were generated in 18/24 cell lines; we obtained on average 13 ± 7 million paired-end reads. Adapter sequences were trimmed using cutadapt in paired-end mode (non-default parameters: -q 15 -O 5 -m 30). Reads were aligned to hg19 using bowtie (version 0.12.8; non-default parameters: -X 2000 -m 1) after which duplicate reads were removed using Picard MarkDuplicates. Genome-wide signal tracks for each sample were generated using align2rawsignal (https://code.google.com/archive/p/align2rawsignal/wiki/Method.wiki).

**Overlap between cohesion-mediated chromatin loops and high-resolution Hi-C loops, contact domains and TADs**

We obtained the coordinates for Hi-C loops from seven cell lines (including GM12878) and contact domains in GM12878 to calculate the overlap with our pan-cell line loops (Fig. 1d). We also obtained the coordinates for TADs across 21 human tissues and cell types and compared the size of these TADs to our pan-cell line loops (Fig. 1c).

**Assessing CTCF motif orientation**

A list of CTCF motif positions and orientations was downloaded from the ENCODE project. We used the CTCF_known1 motif for all analysis; this motif most closely matched the one used in a previous analysis. Next, for all motifs that contained exactly one instance of the CTCF motif at both ends (that is, in both anchor regions), we calculated the percentage of loops that had each of four possible orientations (+/−, −/+; +/+ and −/−). This result was relatively robust to the choice of threshold used to define the pan-cell line loop set (FDR<10−5: 69%, FDR<10−4: 68%, FDR<0.01: 66%, FDR<0.05: 64%).

**Characterizing ‘hub’ anchor regions**

Promoter regions were defined as a 500-bp region immediately upstream of a gene; gene coordinates were taken from Gencode Release 25. Enhancer regions were defined as the set of 288,711 H3K27ac peaks defined from our ChIP–seq data set (see ‘H3K27ac ChIP–seq data processing’ for more information). All anchor regions were binned by the number of interactions they had in the ‘merged loop-call’ data set (Supplementary Table 4). We assessed whether anchor regions in a particular bin were enriched for overlap with functional elements such as enhancers, promoters, or contact domain boundaries (taken from a previous publication) using Fisher’s exact test. For each bin, we tabulated the number of anchor regions that overlapped or did not overlap a functional element; we then tabulated the number of anchor regions in all other bins that overlapped or did not overlap a given functional element. These four values were used to populate a 2 × 2 contingency table and to compute a significance of enrichment. To test the robustness of our results with respect to the threshold used to define the set of merged loop-calls, we repeated this analysis using an FDR<1% (summary statistics for the fold-enrichment and P values can be found in Supplementary Table 9). Qualitatively, we observe very similar results to Fig. 3a—regions with many interactions are enriched for enhancers and contact domain boundaries, whereas promoters tend to overlap regions with fewer interactions.

**PCA**

We performed PCA on the matrix of normalized interaction frequencies of 85,294 loops by 48 samples using the pcomp function in R (default options). The 85,294 loops were derived from the set of pan-cell line loops (Supplementary Table 4) after filtering for interactions that had >4 PETs in at least one sample. We repeated the analysis using the entire set of pan-cell line loops at various FDR cutoffs and observed high correlation in PC1 and PC2 values (FDR < 10−4: rpc1 = 0.999, rpc2 = 0.995; FDR < 0.05: rpc1 = 0.983, rpc2 = 0.981). We also observed similar results when using different PET cutoffs to filter loops (>2 PETs: rpc1 = 0.999, rpc2 = 0.997; >10 PETs: rpc1 = 0.993, rpc2 = 0.985).
Testing for similarity in interaction profiles between similar cell types

For a pair of samples, we calculated the Spearman rank correlation between the raw PET counts across the set of pan-cell line loops identified (124,830 loops) for which there were at least four PETs in at least one sample (85,294 loops). For Fig. 2c, we plotted the distribution of correlation coefficients for the following groups: ‘all’ (all pairs of samples excluding replicates); ‘same germline layer’ (the assignment of individual cell lines to germline layers is provided in Supplementary Table 1; note that replicate pairs are included in this grouping); ‘same tissue’ (the assignment of individual cell lines to tissue is provided in Supplementary Table 1; note that replicate pairs are included in this grouping); ‘biological replicates’ (replicate samples); and ‘isogenic cell types’ (these include cell lines derived from a single male individual (MSCL, MSFIB, and MSIPB); note that replicate pairs are included in this grouping).

Differences in the distribution of correlation coefficients were assessed using a two-sided Wilcoxon rank-sum test. P-values were corrected for multiple hypothesis testing using the Bonferroni approach. We repeated the analysis including replicate pairs in the ‘all’ distribution and observed similar results ($P_{all\ vs\ isogenic\ cell\ types} = 0.4$, $P_{all\ vs\ biological\ replicates} = 4.38 \times 10^{-3}$, $P_{all\ vs\ same\ tissue} = 2.52 \times 10^{-35}$, $P_{all\ vs\ same\ germline\ layer} = 1.23 \times 10^{-25}$). The results were also robust to the particular PET threshold used (we examined thresholds of 1–20 PETs in at least one sample; Extended Fig. 2e). Finally, qualitatively similar results were observed when we used normalized PET interaction frequencies instead of raw PET counts ($P_{all\ vs\ isogenic\ cell\ types} = 0.79$, $P_{all\ vs\ biological\ replicates} = 2.6 \times 10^{-15}$, $P_{all\ vs\ same\ tissue} = 9.2 \times 10^{-25}$, $P_{all\ vs\ same\ germline\ layer} = 3.2 \times 10^{-3}$).

Assessing the effect of technical confounders on loop interaction frequency

For each ChIA-PET sample, we recorded the following potential confounding variables: batch (the set of samples which were processed at the same time and pooled together for sequencing); normalized strand cross-correlation coefficient (NSC; a metric of ChIP efficiency/quality); number of peaks called; and number of uniquely mapped PETs between 10 kb and 5 Mb.

We tested for an association between principal components 1–10 (see ‘PCA’) and each covariate described above using a linear model and assessed significance using the ANOVA implementation in R. P-values were corrected for multiple hypothesis testing using the Benjamini–Hochberg procedure. At an FDR <10%, we detected no significant associations. Thus, we chose not to correct for any of these technical confounders when testing for variable loops (see below).

Identifying variable loops

We began with the set of 124,830 merged loop calls and filtered loops to include only those that had ≥4 PETs in at least one sample yielding 85,294 loops. Next, we estimated the mean to variance relationship in the data using the voom method and used the inverse variance weights at the same time and pooled together for sequencing; normalized strand cross-correlation coefficient (NSC; a metric of ChIP efficiency/quality); number of peaks called; and number of uniquely mapped PETs between 10 kb and 5 Mb.

We tested for an association between principal components 1–10 (see ‘PCA’) and each covariate described above using a linear model and assessed significance using the ANOVA implementation in R. P-values were corrected for multiple hypothesis testing using the Benjamini–Hochberg procedure. At an FDR <10%, we detected no significant associations. Thus, we chose not to correct for any of these technical confounders when testing for variable loops (see below).

Linear model. For each loop, we fitted a linear model [log(normalized interaction frequency) - cell type] and assessed its significance using the ANOVA implementation in R. P-values were corrected for multiple hypothesis testing using the Benjamini–Hochberg procedure. At an FDR <10%, we found 21,353 loops; 20,926 of these were also found using the approach described above (98%; 2.34 fold-enriched compared to hypergeometric expectation).

Non-parametric approach. For each loop, we tested for differences in the normalized interaction frequency using a Kruskal–Wallis test. As a non-parametric approach is likely to be under-powered, we ranked ordered all interactions according to P-values and examined the overlap for the top 35,698 interactions (that is, the same number as found using the mixed effects linear model). A total of 23,117 overlapping hits were found (64% of the set found using the mixed effects linear model; 1.35 fold-enriched compared to hypergeometric expectation).

Defining a set of non-variable loops (static loops)

To compare various attributes of our differential loops, we defined two sets of invariant or static loops as follows.

Static (null set) 1. For each of the 85,294 loops we tested for differential interaction frequency, we computed an ad hoc metric as follows:

$$\text{Score}_{\text{static}} = \frac{1}{\text{relative entropy}} \times \text{mean PET frequency}$$

in which relative entropy is defined as follows:

$$\text{Relative entropy} = \sum_j f_j \log_2 \frac{f_j}{q_j}$$

j sums across all samples (that is, cell lines) and $f_j$ represents the fractional PET count in sample $j$ (that is, the ratio of the number of PETs in sample $j$ divided by the total number of PETs for this particular loop). $q_j$ represents the fractional PET count under a null model assuming an equal number of PETs in each sample. In essence, a high static score would indicate a strongly interacting loop with uniform interaction frequencies across all cell lines. All loops were ranked in descending order by their static score and we selected the same number of high-scoring interactions as differential interactions identified (FDR <10%).

Static (null set) 2. From the set of 85,294 loops tested for differential activity, we selected a set of interactions found to not have differential activity (FDR >50%), but matched for the following properties to the set of differential interactions (FDR <10%): number of loops; distribution of loop sizes; and distribution of P-values assigned by Mango (from the merged loop data set).

The last criterion helps to ensure that the static set of interactions is roughly comparable in quality to the differential interaction set.

Defining housekeeping and cell-type-specific genes

For all 22,197 genes, we computed a relative entropy score as defined in ‘Defining a set of non-variable loops (static loops)’ above. We then removed genes with low expression (minimum expression across all samples had to be >1 TPM). Genes in the top and bottom 10% as ranked by the relative entropy score were designated as ‘cell-type-specific’ and ‘housekeeping’ genes, respectively. Finally, we assessed whether variable or non-variable loops were enriched for housekeeping genes or cell-type-specific genes as follows. For the set of variable or non-variable loops (both null set 1 and null set 2), we tabulated the number that contained or overlapped more than one housekeeping or cell-type-specific gene. Similarly, we tabulated the number of variable or non-variable loops that contained or overlapped no genes in either the housekeeping
Chromatin state analysis with cell-type-specific loop ends

Chromatin state calls using a 15-state model for 12 cell lines were obtained from the Roadmap Epigenomics Mapping Consortium (Supplementary Table 1). We merged chromatin states calls into eight categories as follows: (1) TSS–1 TssA, 2 TssAFink; (2) BiV: 10 TssBiV, 11 BiVfink; (3) TX: 3 TxFink, 4 Tx, 5 TxWk; (4) REPRESS: 13 RepPC, 14 RepPCWk; (5) REPEAT: 8 ZNF/Rpts; (6) ENH: 12 EnhBiV, 6 EnhG, 7 Enh; (7) HET: 9 Het; and (8) QUIES: 15 QuiEs.

Next, for each cell line, we identified a set of loops that were present only in the cell line of interest (CellLine QuerY) and not in all other cell lines (CellLine othEr) as follows: 1. Calculate a t-statistic based on the comparison of interaction frequencies (raw PET count) for all samples in CellLine QuerY and CellLine othEr. 2. Rank order each vector of t-statistics in descending order. 3. Define the set of cell-type-specific loops as the top 10% of loops identified in Step 2.

To assess the enrichment of various chromatin states at cell-type specific loop ends, we generated a 2 x 2 contingency table populated with the following four values: 1. Number of loop-ends that participated in a cell-type-specific interaction that overlapped a chromatin element. 2. Number of loop-ends that participated in a cell-type-specific interaction that did not overlap a particular chromatin element. 3. Number of loop-ends that did not participate in a cell-type-specific interaction that overlapped a particular chromatin element. 4. Number of loop-ends that did not participate in a cell-type-specific interaction that did not overlap a particular chromatin element.

Significance was assessed using the Fisher’s exact test. P values were corrected for multiple hypothesis testing (12 cell lines x 8 chromatin states) using the Benjamin–Hochberg procedure. We repeated our analysis using different rank thresholds to define the set of cell-type specific interactions by repeating this analysis using different thresholds (5%, and 1%) and assessed the robustness of our results, by comparing the overlap in enriched/under-enriched chromatin states. At a 5% rank threshold cutoff, eight cell lines had perfect agreement (Hi-hESC, NCI-H1437, H9-hESC, HepG2, K562, LX, MSIP5, MSiFIB). Three agreed for 7/8 chromatin states (HPAEC, GM12878, NP) and one agreed for only 5/8 (Jurkat). At a 5% rank threshold cutoff, eight cell lines had perfect agreement (HPAEC, NCI-H1437, H9-hESC, HepG2, K562, LX, MSIP5, MSFIB). Three agreed for 7/8 chromatin states (MSIP5, Jurkat, HepG2, and Hi-hESC).

In cases of disagreement, except for Hi-hESC, the typical change in result was the BIVALENT state going from over-enriched to no enrichment. For Hi-hESC, the REPEAT state went from under-enriched to no-enrichment. Nevertheless, the vast majority of results were similar across all thresholds.

To assess whether cell-type-specific loops were enriched for TSS–TSS, TSS–ENH, or ENH–ENH, we first identified cell-type-specific loops, genes, and enhancer peaks as described above. To have adequate numbers, we defined the set of cell-type-specific genes as the top 20% of genes identified using the procedure above.

Next, we counted the number of cell-type-specific loops whose ends overlapped one of the three chromatin state combinations described above. Similar to above, we counted the number of non-cell-type-specific loops whose ends overlapped one of the three chromatin state combinations described above. An enrichment test was then performed using Fisher’s exact test.

Testing for an association between gene expression level and number of linked enhancers

For each cell line, we identified a set of (i) cell-type specific loops (that is, high interaction frequency in cell line of interest and not in others), (ii) enhancers, and (iii) genes (that is, high normalized expression levels in cell line of interest and not in others) using the procedure outlined above (see ‘Chromatin state analysis with cell-type-specific loop ends’). Next, for each gene that was expressed in a single cell type of interest, we tabulated the number of cell-type-specific enhancers that were linked to its promoter. To generate Fig. 3f we aggregated results across all cell lines. To test for differences in the distribution of normalized expression levels between numbers of linked enhancers, we used the Wilcoxon rank-sum test. We repeated the analysis using different cutoffs to define cell-type specific loops, including 1% (P = 2.2 x 10-15), 5% (P = 2.2 x 10-15), and 10% (P = 2.2 x 10-15).

Loop architecture in disease-associated genes

We downloaded the lists of disease-associated genes from ClinVar47, the GWAS catalogue46 and haploinsufficient genes45. The set of housekeeping genes was defined as above (Defining housekeeping and cell-type-specific genes). For each list of genes, we tested the association of the gene being part of the specific category (ClinVar, GWAS or haploinsufficient) and having at least X loops connected to its promoter where X was a number from 1 to 10. We repeated the same test, filtering the loops for only enhancer loops (with a H3K27ac signal at the other end), and cell-type-specific enhancer loops (a H3K27ac mark in a given cell type). P values were calculated using Fisher’s exact test and corrected for multiple testing using the Benjamin–Hochberg approach. A list of all enrichments and P values is provided in Supplementary Table 9.

Mapping genes to loops

To integrate gene expression and histone data, we generated a map of genes to loops as follows: ‘All’ (a gene was assigned to any loop within 1 kb of its start or end coordinates, as defined in Genencode version 25 lifted to hg19, or if the ORF overlapped partially with the loop); ‘Promoter’ (a gene was assigned to any loop for which its TSS was within 1 kb of either anchor region); ‘Contained’ (a gene was assigned to any loop it was entirely contained within (that is, start and end coordinates of the gene fell entirely within a loop) and its promoter was more than 1 kb from either anchor region); and ‘Promoter–enhancer’ (one loop end overlapped a promoter, the other end overlaps an H3K27ac peak).

Linking gene expression changes to changes in loop interaction frequency

For each loop, we correlated the normalized interaction frequencies across all cell types (Spearman rank correlation; n = 23 cell types with RNA-seq and ChiA-PET data) with the normalized gene expression levels across all cell types. If a loop mapped to multiple genes, we computed all possible loop–gene correlations. As a control, we shuffled the mapping between loops and genes, while maintaining the total number of genes mapped to a single loop, and re-examined the correlation between loop interaction frequency and gene expression values. This procedure was repeated 100 times and we recorded the mean correlation coefficient for each loop–gene pairing.

In Fig. 4c, we have restricted our analysis to the set of variable loops (FDR < 10%) and plotted the distribution of actual versus randomized correlation coefficients (absolute value) for all loop–gene pairs (n = 90,657). We compared the distribution of actual correlation coefficients to ‘null’ correlation coefficients using the Mann-Whitney U test (P < 2.2 x 10-15). We repeated the analysis using the set of all loops tested for variable interaction frequencies (n = 251,678 loop–gene pairs) and observed significant results (P = 2.2 x 10-15), albeit with a lower mean correlation (0.17 versus 0.19 for the set of variable loops).
To assess what effect the mapping between loop and gene might have, we compared the distribution of correlation coefficients (absolute value) for all loop–gene pairings for all four maps described above (All, Promoter, Promoter–enhancer and Contained). Significance was assessed using a two-sided t-test and P-values were adjusted for multiple hypothesis testing using the Bonferroni approach. We performed three versions of this analysis: (i) using all loops tested for variability (n = 85,294) and all histone peaks (n = 288,711) \( P_{\text{All vs Contained}} = 6.5 \times 10^{-212}, P_{\text{All vs Promoter}} = 2.1 \times 10^{-260}, P_{\text{All vs Promoter-enhancer}} = 1.9 \times 10^{-265}, P_{\text{Promoter vs Promoter-enhancer}} = 1.0 \). (ii) using all loops tested for variability and histone peaks with variable activity. Variability in H3K27ac was assessed using the procedure outlined in ‘Identifying variable loops’. We set a threshold of FDR < 1% to define the set of variable histone peaks \( P_{\text{All vs Contained}} = 6.5 \times 10^{-212}, P_{\text{All vs Promoter}} = 2.1 \times 10^{-260}, P_{\text{All vs Promoter-enhancer}} = 0, P_{\text{Promoter vs Promoter-enhancer}} = 4.9 \times 10^{-29} \). (iii) using all variable loops (FDR < 10%) and all histone peak with variability activity \( P_{\text{All vs Contained}} = 2.2 \times 10^{-119}, P_{\text{All vs Promoter}} = 1.9 \times 10^{-146}, P_{\text{All vs Promoter-enhancer}} = 3.4 \times 10^{-133}, P_{\text{Promoter vs Promoter-enhancer}} = 2.7 \times 10^{-209} \). Taken together, these analysis indicate a stronger link between loop interaction frequency and gene expression when the loop is making direct contact with the gene’s promoter or when linking and enhancer to the promoter. Subsetting either loops or enhancers based on variability does not appear to improve the results.

Finally, we analysed if there was an enrichment for positive loop–gene correlation coefficients for the four maps described above. We tabulated the number of positive and negative coefficients for actual and randomized loop–gene pairs and assessed significance using Fisher’s exact test.

Identifying group-specific loops
All analysis was performed on the set of loops tested for variability (n = 85,294). For each group (blood, embryonic, and solid-tissue-derived), we identified a set of loops that were present only in their member cell lines (Group<sub>pure</sub>) and that did not differ between the other two groups (Group<sub>other1</sub>, Group<sub>other2</sub>) as follows: 1. Compute three sets of t-statistics based on the following three pairwise comparisons: interaction frequencies (normalized interaction frequency) for all cell lines in Group<sub>query</sub> versus Group<sub>other1</sub> \( (t_q) \), interaction frequencies for all cell lines in Group<sub>query</sub> versus Group<sub>other2</sub> \( (t_{q-f}) \), and interaction frequencies for all cell lines in Group<sub>other1</sub> versus Group<sub>other2</sub> \( (t_{f}) \). 2. Rank order each vector of t-statistics in descending order. 3. Define three sets of loops \( (T_q, T_{f}, T_{q-f}) \) such that their respective t-statistics are in the top 10% of \( t_q \), \( t_{f} \) and \( t_{q-f} \) respectively. 4. Define the final set of group-specific loops as \( (T_q \cap T_{f} \cap T_{q-f}) \).

In this way, we specifically identified loops with a high interaction frequency in the group of interest compared to the other two groups and no difference between the other two groups.

Annotating different DUEs
We used bioconductor’s package DEXSeq<sup>77</sup> to identify DUEs. In brief, we flattened the Gencode (release 25; lifted to GRCh37 coordinates) file with parameters ‘-r no’ and used a modified script to extract counts with subRead (parameters ‘-O -s 2 -p T -40’) as described in the vignette<sup>78</sup>. We classified the RNA-seq libraries either according to the three clusters identified with the PCA as described above, or by cell line \( (n = 22) \). Next we normalized for library size and dispersion, tested for DUEs, and estimated the exon log<sub>2</sub>-fold changes between (a) solid vs blood and stem cell-like vs blood, or (b) by cell type vs the median exon abundance. In this way, we identified (a) 95,137 and (b) 39,832 DUEs (FDR = 10%).

Defining intragenic loops
As a way to identify intragenic loops that go from promoters to gene bodies, we followed the methods described previously<sup>13</sup>. Starting from the Gencode annotation (release 25; lifted to GRCh37 coordinates), we only kept protein-coding genes with at least one middle exon. We also removed all exons that overlapped previously defined CAGE peaks<sup>79</sup>. Based on visual inspection, we defined the promoter window as ±1 kb from the TSS and the upstream window as ±5 kb from the 5’ exon boundary. We then identified intragenic loops as those loops for which one anchor fell in the promoter and the second in the upstream window of the same gene. In this way, we identified 1,372 loops within 1,074 genes. From this set, we identified exon–loop pairs (real pairs) by associating an exon with an anchor of an intragenic loop within 5 kb of its 5’ boundaries.

Correlation of exon and loop anchors
We kept unique exon–loop pairs and correlated the normalized counts of exon and anchor strength across the 22 cell lines. As a control, we permuted all exons 100 times, creating new exon–loop pairs. We also accounted for gene expression by correlating all other exons within the same ‘looping’ gene and removed any exons within 20 kb of the centre of the anchor (all pairs). Then we performed a Pearson correlation for all complete observations and depicted only the DUEs across the 22 cell lines. For the scatterplot, we used the three-group classification specified above and we tested for correlation between real pairs and all pairs of the DUEs.

TF enrichment analysis
We obtained the genomic coordinates for motif matches for 598 TFs from a previously published study<sup>83</sup>. For each TF, we tabulated the following four numbers: (i) the number of group-specific loop-ends overlapping a motif location, (ii) the number of group-specific loop-ends not overlapping a motif location, (iii) the number of non group-specific loop-ends overlapping a motif location, and (iv) the number of non group-specific loop-ends not overlapping a motif location. We assessed the significance of enrichment using a two-sided Fisher’s exact test. In cases in which any of values (1)–(4) were less than 5, we excluded this TF from further analysis. P-values were corrected for multiple hypothesis testing using the Benjamin–Hochberg procedure. We repeated the analysis using different rank thresholds used to define the set of group-specific loops. Using a 5% threshold, we observed high correlation of fold-enrichment values \( r_{\text{Blood}} = 0.89, r_{\text{Embryonic}} = 0.88 \). Moreover, out of the 120 significant TF enrichments for the blood-specific loops (FDR < 0.1), 74 were significant at this new threshold (3.74 fold-enrichment, \( P = 5.5 \times 10^{-38} \) via hypergeometric test). For the 89 significant TF enrichment (FDR < 0.1) from embryonic-specific loops, 39 were significant at this new threshold (5.6 fold-enrichment, \( P = 1.1 \times 10^{-28} \) via hypergeometric test). Using a 25% threshold, we again observed high correlation of fold-enrichment values \( r_{\text{Blood}} = 0.83, r_{\text{Embryonic}} = 0.86 \). Moreover, out of the 120 significant TF enrichments (FDR < 0.1) for blood-specific loops, 95 were significant at this new threshold (2.72 fold-enrichment, \( P = 1.5 \times 10^{-10} \) via hypergeometric test).

Transcription factor footprinting in ATAC–seq data
ATAC–seq data were processed (Methods) for signal tracks. Motifs for each TF were intersected with the loop annotations and ATAC–seq data were averaged across all motif instances using a custom Python script. Averaged signal was compared between blood-specific, embryonic-specific, and all loops, and the relevant ratios were computed and plotted for a given TF.

GO biological process enrichment of group-specific loops
Using the procedure outlined in ‘Identifying group-specific loops’ above, we defined 3,384 blood-specific loops, 2,894 embryonic-specific loops, and 2,215 ‘misc’-specific loops. For each loop, we defined its
peaks>–w-ld-chr < weights_hm3_no_hla>–overlap-annot–out < output
filtered for SNPs with minor allele counts of at least five. The following
regions, and so on62 and the full set of Rad21-bound looped regions
we ran LDSC with the standard 1000G Phase III derived LD scores and
the same trait. We only kept GWAS with at least 1,000 SNPs after pruning
in the analysis for sufficient quality to calculate an enrichment (n = 86).
The set of pruned SNPs was then expanded to all linked variants with
European r² ≥ 0.8 for all further analysis.

We performed a rank-based enrichment of GWAS variants in each set of
group-specific loops. We segmented each GWAS study into bins that
represented decreasing tiers of significance. We set a minimum bin size of
50 and filled the first bin with the 50 most significantly associated
variants for each study. We then filled the next bins with 2 × 50, 4 × 50
and 8 × 50 variants and then segmented the remaining variants into bins
at the four quartiles of the remaining P value distribution. We used the
pruned set of SNPs to set the bin thresholds. We then computed the
rank fold change enrichment of peaks across the segmented GWAS80.
For each bin we computed the fraction of GWAS variants that were
less than or equal to the bin’s P value threshold that overlapped the
loop regions. We calculated the fold change enrichment by dividing
this fraction by the fraction of all GWAS variants of any significance
level that overlapped our regions. Baseline enrichment is 1, which indi-
cates no change from the base rate of overlap of all significant and
non-significant variants in the study. An enrichment less than 1 means
the most significant variants are depleted relative to the baseline and
any value greater than 1 indicates that significant variants are enriched.
To compute the significance of these enrichments, we permuted the
P value associated with each GWAS SNP in the study 200 times and
re-computed the enrichment relative to baseline. The empirical P value
indicates the number of permuted studies for which the true study has
a greater enrichment for the most significant bin of GWAS hits.

To compare the enrichment of each given GWAS study between sets of
regions, we computed the total number of pruned genome-wide
significant (P < 10 × 10⁻⁸) SNPs that overlapped each set of peaks and the
total number that did not. An overlap was counted if any SNP in LD with
the pruned SNP overlapped the regions of interest. This is important
as we do not know which is the causal SNP. We then used Fisher’s exact
test to statistically compare the rate of overlap between the two studies
and to determine whether a set of regions was statistically enriched
relative to another (Supplementary Table 7).

LD score regression
Partitioned LD score regression (LDSC) is a method to determine
whether there is an enrichment of GWAS effect sizes in a given por-
tion of the genome65. We used LDSC to test whether our loop anchors,
called loops, and DNase peaks within called loops that changed between
cell types were associated with GWAS signal of complex traits. Using
publicly available summary statistics of GWAS for complex traits86,
we ran LDSC with the standard 1000G Phase III derived LD scores and
weights, correcting for the baseline annotations (which contain the
union of H3K27ac marked regions in the genome, H3K4me3 marked
regions, and so on85 and the full set of Rad21-bound looped regions
genome-wide. Regression coefficients were estimated using the
overlap-annot option to partition effects across overlapping regions63
and with frequency files derived from 1000G Phase III Europeans and
filtered for SNPs with minor allele counts of at least five. The following
command was used: lscd.py–h2<input summary statistics>–ref-l-d-chn<1000G EUR Phase3 baseline>,<tested anchor regions>,<all rad21 peaks>–w-l-d-chn<weights_hm3_no_hla>–overlap-annot–out < output estimates>–frqfile-chr <1000G.mac5eur>. Results were parsed for the
enrichment of the tested anchor region and the reported statistics are
taken directly from the command output.

Correction for super-enhancers and cell type effects in LDSC
Super-enhancers are associated with increased chromatin looping
and also with GWAS enrichment, so we wanted to test whether our
signal was due to a super enhancer signal. As such, we excluded called
super-enhancers from any cell type from the tested anchor and loop
annotations and re-ran the enrichment. In addition, after filtering out
anchors from any loops that overlapped with super-enhancers we still
see enrichment for the same traits (Extended Data Fig. 7, Supplemen-
tary Table 8). To assess whether the signal we observed might be just
attributable to active chromatin in the cell types of interest, we added
in all ten cell-type group annotations as covariates to the regression, along
with the Roadmap control signal for per-mark accounting as previously
described62 (Extended Data Fig. 7, Supplementary Table 8). The result-
ing regression was: lscd.py–h2<input path>–ref-l-d-chr <1000G EUR-
Phase3 baseline>,<tested anchor regions>,<all rad21 peaks>,<roadmap
control>,<cell_type_group 1>,<cell_type_group 2>,<cell_type_group
3>,<cell_type_group 4>,<cell_type_group 5>,<cell_type_group 6>,<cell
_type_group 7>,<cell_type_group 8>,<cell_type_group 9>,<cell_type
_group 10>–w-l-d-chr < weights_hm3_no_hla>–overlap-annot–out < output estimates>–frqfile-chr <1000G.mac5eur>.

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

Data availability
The ChIA–PET data have been deposited in the ENCODE webpor-
tal and can be accessed here: https://www.encodeproject.org/
publications/8d853642-45b4-47cf-ad66-f3ce3058a39d/. The remaining
data have been deposited in the GEO database under accession number
GSE134745. There are no restrictions on data availability. Supplemen-
tary Table 10 lists all available data sets.

Code availability
Scripts for processing the raw data can be found here: https://github.
com/rothii-srivasa/ChiaPET. Custom scripts for generating figures can
be obtained upon request.

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Author contributions F.G., R.S., D.V.S., M. Kasowski and M.S. conceived the study. F.G. and D.V.S. performed the experiments with assistance from R.S. Data analysis was performed by R.S. with assistance from F.G., D.V.S., M. Kasowski and J.B.Z. D.V.S., A.N., Q.L., B.G., A.S., M. Kulik, and S.S. performed the cell culture. M.R.-V., N.S.-A. and F.G. provided expertise and assistance in data analysis. M.R., A.K. and S.D. provided insights and support for the design and execution of the project. F.G., R.S., D.V.S., M. Kasowski, J.B.Z. and M.S. prepared the manuscript with assistance from M.R.V., N.S.A. and P.G.

Competing interests The authors declare no competing interests.

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Extended Data Fig. 1 | Quality metrics and convergence rates. 

a, Flowchart of study design. b, Total number of reads obtained for each ChIA-PET sample. c, Relative strand correlation (RSC) score for RAD21 ChIA-PET data. d, RSC for H3K27ac ChIP-seq data. e–g, Comparison of CTCF motif presence and orientation at chromatin loops identified in this study and other published data sets. e, Fraction of chromatin loops with exactly one CTCF motif at both loop ends. f, Fraction of chromatin loops with at least one CTCF motif at both loop ends. g, Fraction of chromatin loops with convergent CTCF motif orientation.
Extended Data Fig. 2 | Variability in chromatin loops. **a**, PCA of H3K27ac ChIP-seq data (a: n = 288,711 peaks versus n = 44 samples (22 cell types × 2 replicates each)) and RNA-seq data (b: n = 22,197 genes versus n = 46 samples (23 cell types × 2 replicates each)); samples are coloured according to the germ layer from which they originated. **c**, PCA of chromatin loop interaction frequencies (n = 85,294 loops versus n = 48 samples (24 cell types × 2 replicates each)). Colours denote the experimental batch of each sample. **d**, GC content in anchor regions of different sets of chromatin loops. Centre line represents the median, box extent ranges from 25th to 75th percentile and whiskers extend at most to 1.5 × the interquartile range. Summary statistics for the boxplots can be found in Supplementary Table 9. **e**, Correlation of chromatin loop interaction frequencies (Spearman rank correlation; y-axis) between pairs of cell types at varying PET thresholds (x-axis). **f**, Number of variable loops found at different FDR thresholds. **g**, Size distribution of variable chromatin loops versus non-variable loops at different FDR cutoffs. ***P < 0.04. Significance was assessed using a two-sided t-test. Centre line represents the median, box extent ranges from 25th to 75th percentile, whiskers extend at most to 1.5 × the interquartile range. Summary statistics for the boxplots can be found in Supplementary Table 9. **h**, Variability of loops of different sizes. Summary statistics for all box plots can be found in Supplementary Table 9. **i**, Enrichment of cell-type-specific genes and depletion of housekeeping genes (n = 2,220) in variable versus non-variable loops (n = 35,698) for null sets 1 and 2. P-values by Fisher’s exact test. **j**, Enrichment of broadly expressed genes at variable and non-variable chromatin loops. The set of broadly expressed genes was obtained from the GTEx project14,15.
Extended Data Fig. 3 | Cell-type-specific loops exhibit enrichment for specific chromatin states. a, Proportion of chromatin states in cell-type-specific loop ends for various cell types from the blood group (red), the stem cell/embryonic group (purple) and the group derived from solid tissue (black). b, Fold-enrichment of chromatin states in cell-type-specific loop ends for the cell types in a. Number of interactions assessed (top 10%) = 8,529; *P < 0.05; **P < 0.005; ***P < 2.2 × 10^{-16}, n.s. = non-significant; P values assessed by two-sided Fisher's exact test and corrected for multiple hypothesis testing using the Benjamini–Hochberg procedure. See Supplementary Table 9 for a complete list of enrichments and P values.
Extended Data Fig. 4 | Connectivity of genes corresponds to gene function.

a–d, log₂ odds ratios for different groups of genes with a certain number of loops linked to their promoters (adjusted *P* < 0.05 by two-sided Fisher’s exact test; *n* = 19,353 loops). See Supplementary Table 9 for a complete list of *P* values.

- **a**, Haploinsufficient genes;
- **b**, genes in GWAS catalogue;
- **c**, disease genes in ClinVar;
- **d**, housekeeping genes.

E–h, log₂ odds ratios for each cell type shown for genes identified as haploinsufficient (e), gene in GWAS catalogue (f), disease genes in ClinVar (g) or housekeeping genes (h) and having at least a given number of loops ending at its promoter (adjusted *P* < 0.05 by two-sided Fisher’s exact test). See Supplementary Table 9 for a complete list of enrichments and *P* values.
Extended Data Fig. 5 | Chromatin loops are associated with alternative splicing across cell types. a, b, Distribution of the distance (bp) between the centre of the loop anchors and the TSS (a) or the exon 5′ boundary (b). c, DEXSeq plot showing the differential exon usage of all exons for gene ARHGEF7, highlighting exon 6, which is affected by an intragenic loop in the blood cell types. d, Scatterplot of the normalized counts of exon 6 in ARHGEF7 with respect to the log2 transformed fold change in loop strength for all cell types (*n* = 44 (22 cell types × 2 biological replicates); Pearson correlation, 0.49).
Extended Data Fig. 6 | Transcription factor analysis, GO enrichments and GWAS for embryonic-specific loops. a, Fold-enrichments of 598 TF motifs in chromatin loop ends that are embryonic-specific (n = 2,894). Significance was assessed using two-sided Fisher’s exact test. P values were adjusted for multiple hypothesis testing using the Benjamini–Hochberg procedure. **b**, Biological processes associated with embryonic-specific chromatin loops (n = 2,894). Enrichment was performed using GREAT66. **c**, Enrichment of disease-specific GWAS SNPs in embryonic-specific loop ends. MDD, major depressive disorder; BPD, bipolar disorder; HOMAIR, homeostatic model assessment for insulin resistance. **d**, Association of embryonic-specific chromatin loop anchors (n = 2,894) with GWAS traits observed by partitioned LDSC62 using a common set of 47 traits63; (n = 1,100,000 HapMap3 SNPs, block jackknife t-test, mean ± s.d.).
Extended Data Fig. 7 | Association of chromatin loops with GWAS traits. 

**a–h.** Association of blood-specific (a–d) and embryonic-specific (e–h) chromatin loop anchors with GWAS traits observed by partitioned LDSC using a common set of 47 traits (n = 1,100,000 HapMap3 SNPs, block jackknife t-test, centre values indicate the mean ± s.d.). Within each panel: left, all blood-specific loops (a–d) or embryonic-specific loops (e–h); right, set of loops that does not overlap with super enhancers. All panels adjusted for the set of baseline line traits as previously described; in addition, b, f are adjusted for all RAD21 loops; c, g are adjusted for super-enhancers across all cell types, within blood-specific and embryonic-specific loops; d, h are adjusted for cell-group-specific signal and global Roadmap annotation.
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

All analysis was done in R (version 3.3.1) using custom scripts which can be found here: https://github.com/rohith-srivas/ChiaPET

Data analysis

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The ChIA-PET data has been deposited on the ENCODE webportal and can be accessed here: https://www.encodeproject.org/publications/8d853642-45b4-47cf-ad6f-32c3058a39d/
The remaining data have been deposited in the GEO database under accession number GSE134745. There are no restrictions on data availability.
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Life sciences study design

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

| Sample size | No sample-size calculation was performed. |
|-------------|------------------------------------------|
| Data exclusions | For two cell lines we were unable to produce RNA-seq and H3K27ac ChIP-seq data due to lack of material. This exclusion was not pre-established. For integrative analysis we have utilized only 22/24 cell lines. |
| Replication | We have performed all experiments in biological replicates to ensure reproducibility of the data. The data passes ENCODE QC metrics; these analysis have been detailed in our Supplementary figures. |
| Randomization | There are no experimental groups in this study. We have assigned random identifiers to samples during the experimental procedures to minimize batch effects. |
| Blinding | Investigators were not blinded, but samples were stripped of their ID and assigned a random ID at the beginning of the respective experiments. |

Reporting for specific materials, systems and methods

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

| Materials & experimental systems | Involved in the study | Methods |
|----------------------------------|-----------------------|---------|
| n/a | Antibodies | n/a | Involved in the study |
| | Eukaryotic cell lines | | ChIP-seq |
| | Palaeontology | | Flow cytometry |
| | Animals and other organisms | | MRI-based neuroimaging |
| | Human research participants | | |
| | Clinical data | | |

Antibodies

Antibodies used
Abcam Anti-RAD21 antibody (ab992, lot ID:GR184716) and H3K27ac (Abcam #4729, lot ID: GR104852)

Validation
Abcam Anti-RAD21 antibody (ab992, lot ID:GR184716) https://www.encodeproject.org/antibodies/ENCAB529YRC/ and H3K27ac (Abcam #4729, lot ID: GR104852) https://www.encodeproject.org/antibodies/ENCAB000BSK/ |

Eukaryotic cell lines

Policy information about [cell lines](#)

Cell line source(s)

- K1 Sigma-Aldrich
- HepG2 ENCODE
- ARPE-19 ATCC
- JURKAT ATCC
- MCF7 ENCODE
- DU145 ATCC
- NCI-H1437 ATCC
- HT-1376 ATCC
- MSLCL established in the Snyder lab, Stanford University
- SU-DHL-2 ATCC
- SU-DHL-4 ATCC
- GM12878 ENCODE
| Cell lines | Notes |
|-----------|-------|
| C1300     |      |
| ATCC      |      |
| HT-1197   |      |
| ATCC      |      |
| H9-hESC   |      |
| Dalton lab, UGA |
| KU-19 DSMZ |      |
| MSFiB     |      |
| established in the Snyder lab, Stanford University |
| K562 ENCODE |      |
| LX        |      |
| differentiated from H9-hESC in the Dalton lab, UGA |
| K1        |      |
| (Thyroid, papillary carcinoma) is commonly misidentified with CVCL_9918 another Thyroid, papillary carcinoma. Since both cell are papillary carcinoma the conclusions of our study would not be affected |

**ChIP-seq**

**Data deposition**

- Confirm that both raw and final processed data have been deposited in a public database such as GEO.
- Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

**Data access links**

The ChIP-Seq data from this study have been deposited in the GEO database under accession number GSE134745.

**Files in database submission**

A full list of file can be found under the accession numbers provided above.

**Genome browser session**

(e.g. UCSC)

no longer applicable

**Methodology**

- Replicates: two biological replicates per cell type
- Sequencing depth: we obtained on average 43 ±9 million paired-end reads 2*101bp) per sample.
- Antibodies: Abcam Anti-RAD21 antibody (ab992, lot ID:GR184716) and H3K27ac (Abcam #4729, lot ID: GR104852)
- Peak calling parameters: peaks were called using MACS2 (non-default parameters: -q 0.01).
- Data quality: Two replicates were performed per cell line. ChIP data has been validated according to ENCODE standards, e.g. Relative strand correlation (RSC). Quality control metrics and concordance between replicates has been detailed in Extended data figure 1 and 2.
- Software: All analysis was done in R (version 3.3.1) using custom scripts which can be found here: https://github.com/rohith-srivas/ChiaPET. Picard MarkDuplicates was used to remove duplicates. Peaks were called using MACS2.