Integrative analysis of haplotype–resolved epigenomes across human tissues

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Allelic differences between the two homologous chromosomes can affect the propensity of inheritance in humans; however, the extent of such differences in the human genome has yet to be fully explored. Here we delineate allelic chromatin modifications and transcription among a broad set of human tissues, enabled by a chromosome-spanning haplotype reconstruction strategy. The resulting large collection of haplotype-resolved epigenomic maps reveals extensive allelic biases in both chromatin state and transcription, which show considerable variation across tissues and between individuals, and allow us to investigate cis-regulatory relationships between genes and their control sequences. Analyses of histone modification maps also uncover intriguing characteristics of cis-regulatory elements and tissue-restricted activities of repetitive elements. The rich data sets described here will enhance our understanding of the mechanisms by which cis-regulatory elements control gene expression programs.

We performed chromatin immunoprecipitation followed by sequencing (ChiP-seq) to generate extensive data sets profiling 6 histone modifications across 16 human tissue types from four individual donors (181 data sets). In combination with previously published data sets, we conducted in-depth analyses across 28 cell/tissue types, covering a wide spectrum of developmental stages, including embryonic stem cells, early embryonic lineages and somatic primary tissue types representing all three germ layers (Fig. 1a) (protocols received approval from IRB/ESCRo and Mid-American Transplant Services, and research consent was obtained from families). The modifications demarcate active promoters (histone H3 lysine 4 trimethylation (H3K4me3) and H3 lysine 27 acetylation (H3K27ac)), active enhancers (H3 lysine 4 monomethylation (H3K4me1) and H3K27ac), transcribed gene bodies (H3 lysine 36 trimethylation (H3K36me3)) and silenced regions (H3K27 or H3 lysine 9 trimethylation (H3K27me3 and H3K9me3, respectively)), and we systematically identified cis-regulatory elements by employing a random-forest-based algorithm (RFECs), predicting a total of 292,495 enhancers (consisting of 175,912 strong enhancers with high H3K27ac enrichment) across representative samples of all 28 tissues types (Supplementary Table 1). We additionally identified 24,462 highly active promoters with strong H3K4me3 enrichment (see Supplementary Table 2). Subsequently, we defined tissue-restricted promoters (n = 10,396) and enhancers (n = 115,222) (Extended Data Fig. 1a). Consistent with previous studies, enhancers appear more tissue-restricted than promoters and cluster along developmental lineages (Extended Data Fig. 1a, b). Moreover, tissue-restricted enhancers were enriched for putative binding motifs of particular transcription factors known to be important in maintaining the cell/tissue type’s identity and function (Extended Data Fig. 2).

Recent studies showed that particular repetitive elements, such as endogenous retroviruses (ERVs), could participate in transcriptional regulation during mammalian development. Given the representation of samples available, we systematically examined histone modifications at different classes of ERVs. While most are inactive, subsets, especially class I ERVs (ERV-I), are marked by H3K27ac in a tissue-restricted manner (Extended Data Fig. 3a and b). For instance, HERV-H element activities are restricted to human embryonic stem cells (hESCs) (Extended Data Fig. 3c, d). Furthermore, some ERVs carried marks of active promoters or enhancers (Extended Data Fig. 3d, e). We also observed that theLTR12C subfamily had substantial H3K27ac enrichment across different tissues (Extended Data Fig. 3e). Notably, the individual members appeared to be tissue restricted, suggesting that although the subfamily can be classified as non-tissue restrictively active, individual LTR12C elements were active only in distinct tissue/cell types (Extended Data Fig. 3e). Taken together, the data illustrate that human ERVs display precisely controlled patterns of activity in distinct tissues.

Intriguingly, 15.2% (n = 3,717) of strong promoters were also predicted as enhancers in other tissues, analogous to observations in mice, where intragenic enhancers act as promoters to produce cell-type-specific transcripts. These sites possessed histone modification signatures of active enhancers in some tissue/cell types but were enriched with active promoter marks in others. We termed these sequences cis-regulatory elements with dynamic signatures (cREDS). For example, 1,321 cREDS enhancers showed enrichment of H3K27ac and H3K4me1 and a striking depletion of H3K4me3 in lung (Fig. 1b, c and Supplementary Table 3). However, the signature shifted to that of active promoters in other tissues (Fig. 1b, c). cREDS are also found in other cell/tissue types (Extended Data Fig. 4a). To determine whether cREDS have dual functions, we selected a subset of promoter-marked elements and validated their function with a luciferase reporter assay in hESCs. The majority (7 out of 10) showed promoter activity (Extended Data Fig. 4b). Similarly, 10 of 11 selected cREDS with enhancer signatures in hESCs also functioned as enhancers (Extended Data Fig. 4c). Additionally, subsets of enhancers previously validated in transgenic mice also possessed dynamic signatures (Extended Data Fig. 5). Furthermore, we selected two cREDS, predicted as enhancers in the left heart ventricle, with significant cap

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analysis of gene expression (CAGE) signal[^21], typical of active promoters (Extended Data Fig. 6a, b), and found that they possess heart-restricted enhancer activities in an *in vivo* zebrafish reporter assay (Extended Data Fig. 6c). Consistent with reporter activities, transcriptional properties (RNA-seq values based on reads per kilobase per million mapped reads (RPKM) within ±1 kb of the elements) of cREDS enhancers and promoters are similar to non-cREDS enhancers and promoters, respectively (Fig. 1d). Interestingly, when comparing isoform dynamics across H1 and IMR-90 RNA-seq data sets[^22] with cREDS identified between these two cell types we discovered that a subset of cREDS promoters was accompanied by creation of new transcripts and/or alternative exon usage (*n* = 99) (Fig. 1e), revealing a possible function whereby cREDS influence cell/tissue-specific transcript variants. Taken together, these data show that cREDS can potentially function as both promoters and enhancers in distinct cell types and fine-tune transcriptionmes.

Reasoning that global analysis of allelic histone modification and gene expression patterns would elucidate mechanisms of long-range gene regulation by distal cis-regulatory elements, we re-analysed RNA-seq and ChIP-seq data sets by considering haplotype information. For this purpose, we applied HaploSeq[^1], which integrated genome sequencing with high-throughput chromatin conformation capture (Hi-C) data sets to derive chromosome-spanning haplotypes (see Supplementary Information). For four different tissue donors, we generated haplotypes spanning entire chromosomes with 99.5% completeness on average (the coverage of haplotype-associated genomic regions) and average resolution (the coverage of phased heterozygous SNPs) ranging from 78% to 89% (Fig. 2a and Supplementary Tables 4 and 5). The accuracy of haplotype predictions was validated by the concordance with SNPs residing in the same paired-end sequencing runs. The concordance rates were 99.7% and 98.4% for H3K27ac ChIP-seq reads (described below) and RNA-seq reads, respectively, indicating high accuracy. We then re-analysed 36 mRNA-seq data sets from 18 tissues (including 16 tissues noted above with the addition of bladder and adipose tissue) and 187 ChIP-seq data sets for 6 histone modifications (Supplementary Table 6), from up to 4 individual donors, in a haplotype-resolved context. Although widespread allelic imbalances in gene expression had been previously noted[^22-23], it remains unclear whether this phenomenon is consistent across distinct tissues and individuals, and the underlying mechanism remains undefined. To address the first point, we defined genes with allelically biased expression by means of mapping the RNA-seq reads in each tissue sample in a haplotype-resolved manner. We observed extensive allelically biased gene expression, ranging from 4% to 13% of all informative genes (>10 allelic read counts) in each tissue sample (false discovery rate (FDR) = 5%, Extended Data Fig. 7a, b). Comparatively, the proportion of allelically biased genes in individual tissue donors ranged from 6% to 23% of all informative genes, giving a combined total of 2,570 allelically biased genes (Fig. 2b and Supplementary Table 7). As a control, known imprinted genes (*n* = 15) showed common allelic biases across multiple samples (Fig. 2c) and donors (Extended Data Fig. 7c). Our data sets, representing the only collection of haplotype-resolved transcriptomes across an array of tissues from multiple individuals, allowed us to characterize allelic transcription across tissues and donors. While most genes with allelically biased expression demonstrate bias in multiple samples, approximately 75% exhibit statistically significant donor-specific bias (Fig. 2d and Extended Data Fig. 7d). This suggests a connection between sequence differences of individuals and allelically biased gene expression. In support of this model, genes frequently demonstrate consistent direction of allelic bias across multiple tissues of a given donor (Fig. 2e and Extended Data Fig. 7e). Interestingly, allelically biased genes were not restricted to the same tissue type across distinct donors. Rather, they were mostly specific to individual samples derived from different donors (Fig. 2f and Extended Data Fig. 7f), possibly resulting from differential levels of tissue-restricted transcription factors among different tissue samples. As natural genetic variations can affect enhancer selection and function in mammalian cells[^28], we hypothesized that polymorphisms at
cis-regulatory sequences underlie the widespread allelic transcriptional biases. We thus exploited the unique resource of 187 haplotype-resolved ChIP-seq data sets to analyse the state of cis-regulatory elements. We identified allelically biased marks at promoter regions (H3K27ac, H3K4me1, H3K4me3, H3K27me3 and H3K9me3) and transcribed gene bodies (H3K36me3) (see Supplementary Information). In support of our hypothesis, the allelic biases of gene expression strongly agreed with chromatin states of sequences at or near the genes (Fig. 3a, b and Extended Data Fig. 8a).

Furthermore, if allelic imbalances of enhancer activities indeed contribute to allelically biased gene expression, we expect that chromatin states at enhancers will be concordant with the expression of their targets. Therefore, we generated additional H3K27ac ChIP-seq data sets with deeper coverage and longer sequencing reads (for better delineation of alleles) for 14 of the previously analysed tissue samples and an additional 6 samples from independent donors (Supplementary Table 7). Of the informative enhancers (with >10 polymorphism-bearing sequence reads), 11.6% (n = 11,714, FDR = 1%) showed significant allelically biased H3K27ac enrichment in any tissue types (Fig. 3c and Supplementary Table 8). H3K27ac biases were validated by allele-specific ChIP-qPCR (Extended Data Fig. 8b). Interestingly, identical genotypes often yielded the same direction of biases in allelic enhancer activities (Fig. 3d). We further tested whether sequence variations are systematically associated (H3K36me3) among informative genes and the unphased (grey) variants across chromosome 1 is shown. ** (P < 9.9 × 10⁻⁵), Kolmogorov–Smirnov (KS) test. ** Including only tissues with two or three equivalent samples derived from distinct donors (n = 10), genes with allelic imbalances were defined as common between individuals (consistent bias among same tissue type from multiple donors) or as individual-restricted. Random control represents average from randomly selected samples (10,000 iterations). Abbreviations are defined in the legend to Fig. 1. e, Fold change of gene expressions between alleles (parental allele 1 (P1) and P2) in adenral gland from donor 2 (x axis) is compared to all other tissues from donor 2 (y axis). ** Allele bias of enhancer activities among two donors (P1 vs P2) in adenral gland (donors 3) is compared to all other tissues from donor 2 (y axis). A histogram illustrates the proportions of allelically expressed genes in donors 1 (n = 1,375) defined in various numbers of tissues. The fraction of all testable genes or allelically expressed genes (y axis) is calculated for the number of tissues where they are identified as active (x axis) (P value <2.2 × 10⁻¹⁶, KS test).

Figure 2 | Widespread, individual-specific allelic bias in gene expression. a, Genome browser snapshots illustrate completeness and resolution of haplotypes resolved in donor 4. The y axis indicates the number of variants within 100-kb windows. The density of all (blue), phased (orange) and unphased (grey) variants across chromosome 1 is shown. b, Proportion of genes with allelically biased expression among informative genes and the number of tissue samples derived from each donor (n = 11) are described. c, Box-plot illustrates occurrence of imprinted (n = 15) and other allelically biased genes, excluding pseudogenes (n = 2,334) across samples. ** P < 9.9 × 10⁻⁵, Kolmogorov–Smirnov (KS) test. d, Including only tissues with two or three equivalent samples derived from distinct donors (n = 10), genes with allelic imbalances were defined as common between individuals (consistent bias among same tissue type from multiple donors) or as individual-restricted. Random control represents average from randomly selected samples (10,000 iterations). Abbreviations are defined in the legend to Fig. 1. e, Fold change of gene expressions between alleles (parental allele 1 (P1) and P2) in adenral gland from donor 2 (x axis) is compared to all other tissues from donor 2 (y axis). f, A histogram illustrates the proportions of allelically expressed genes in donor 1 (n = 1,375) defined in various numbers of tissues. The fraction of all testable genes or allelically expressed genes (y axis) is calculated for the number of tissues where they are identified as active (x axis) (P value <2.2 × 10⁻¹⁶, KS test).
with allelic H3K27ac, which reflects enhancer activities. Indeed, H3K27ac biases were strongly correlated with specific genotypes, whereby given identical genotypes, this histone modification was biased to the same alleles, both across tissue types and individuals (Fig. 3d–f and Extended Data Fig. 9a). Furthering this finding, we analysed previously generated data sets from lymphoblastoid cell lines and found similar significant correlation of genotype and molecular phenotype of H3K27ac enrichment (Extended Data Fig. 9b). Taken together, these data reveal that extensive allelic imbalance events are associated with sequence variants in cis-regulatory elements.

We discovered that allelic enhancers resided in significantly closer proximity to genes with allelically biased expression, as compared to non-allelic enhancers (Fig. 4a, b). We also observed examples where distinct tissues from the same donor showed similar allelic biases of gene expression.

**Figure 4 | Allelic histone acetylation at enhancers is associated with allelically biased gene expression.** a. Average distance of allelic (5% FDR) and non-allelic enhancers to the closest allelically expressed gene is significantly different ($n = 3,829$, $P < 2.2 \times 10^{-16}$, KS test). b. Genome browser snapshots show an allelic enhancer within the intron of the allelically expressed A4GALT gene (P1, red; P2, blue) on chromosome 22 across three samples. c. Line plot presents the fraction of concordant allelic bias between allelically expressed genes and allelic enhancers in terms of distance. The allelic enhancer–gene pairs were defined with FDR cutoff values of 5% ($n = 14,082$) (black), 1% ($n = 6,057$) (blue) and 0.1% ($n = 2,362$) (orange). Permuted control of a set of enhancer–gene pairs was included ($n = 14,082$) (grey). Distance between allelically biased enhancer–gene pairs and fraction of concordant allelic bias are denoted by $x$ and $y$ axes, respectively ($P < 2.2 \times 10^{-16}$, KS test). d. Fractions of tissue-restricted enhancer–gene pairs ($n = 3,106$) (y axis) that show concordant (blue) or discordant (orange) allelic biases in the same tissue, are presented across a range of Pearson correlation coefficients ($x$ axis) ($P < 2.2 \times 10^{-16}$, KS test, random permuted control). Concordant and discordant allelic enhancer–gene pairs are marked in motif sequence.

**Figure 5 | Motif disruption by genetic variants is concordant with allelic H3K27ac biases at enhancers.** a. Differential GABPA binding motif scores between two alleles ($P1 - P2$ motif scores) in left heart ventricle are correlated with the proportion of H3K27ac reads corresponding to the $P1$ allele (top). Values range from negative to positive, indicating $P1$ and $P2$ motif disruption, respectively. An example on chromosome 12 illustrating that $P1$, with a motif preserving $C$ allele, has higher H3K27ac enrichment and that $P2$, with the motif disrupting $T$ allele, has little H3K27ac enrichment (bottom). b. Three examples ($FL1$ in spleen, $SPDEF$ in sigmoid colon, and $TEAD$ in aorta) of motif-disrupted enhancers demonstrate allelic biased activities. The variant location and genotypes of $P1$ and $P2$ alleles are marked in motif sequence. c. All possible motif disrupted enhancer–gene pairs within the indicated distance window are defined with concordant allelic bias (blue, gene–enhancer pairs with biases towards the same allele) or discordant allelic bias (red, gene–enhancer pairs with biases towards different alleles). Only thymus, left heart ventricle and aorta were considered due to the availability of Hi-C data. Short-range pairs are defined if any allelically expressed genes are located <20 kb away. $***P < 2.5 \times 10^{-5}$, binomial test.
expression and H3K27ac at enhancers (left ventricle and right ventricle from donor 3); however, the same tissue type derived from a different donor (left ventricle from donor 1) yielded no consistent patterns (Fig. 4b), supporting the hypothesis that allelically biased gene expression is driven by individual-specific genetic variation in enhancers. Indeed, within close proximity, the concordance between allelic enhancers and gene expression is significantly higher than permuted control enhancer/gene sets (Fig. 4c). Remarkably, 56% of allelic enhancer–gene pairs are greater than 300 kb apart (Extended Data Fig. 10a, b), the delineation of which was enabled by whole-chromosome-spanning haplotypes.

Similar to genes, many allelically biased enhancers are tissue restricted (Extended Data Fig. 10c). We reasoned that gene expression biases could result from tissue-restricted enhancer activities, evidenced by significant correlation between allelic enhancers and allelically expressed genes (Fig. 4d). Allelic enhancers also significantly overlapped with expression quantitative trait loci (eQTLs) (Fig. 4e), DNase I hypersensitivity QTLs and H3K27ac QTLs (Extended Data Fig. 10d), defined independently28–30, corroborating the functional roles of identified allelic enhancers on gene regulation. Taken together, these observations support a model whereby allelic biases of cis-regulatory element activities could be responsible for allelic gene expression.

Finally, to elucidate further the mechanism by which allelically biased enhancer activities arise, we examined single nucleotide polymorphisms (SNPs) that potentially disrupt or weaken transcription factor binding motifs. We calculated changes in motif score between alleles (motif disruption score) at allelic enhancers and discovered 133 transcription factor motifs showing significant concordance between allelic reduction of enhancer activities and transcription factor motif disruption (Fig. 5a, b) (FDR 10%, Supplementary Table 9). Moreover, genes with allelically biased expression were concordant with enhancer motif disruptions within close proximity (<20 kb) or displaying strong Hi-C interactions at longer distances (>20 kb) (Fig. 5c and Supplementary Information). Our results therefore suggest that genetic variations are probably responsible for allelic enhancer activities and consequently allelically biased gene expression.

By generating chromosome-spanning haplotypes, we carried out a comprehensive survey of allelic chromatin state and gene expression. We found evidence for extensive allelically biased gene expression, which is connected to change in chromatin states at cis-regulatory elements, probably resulting from transcription factor binding disruption by sequence variations. These observations mirror findings in mice where allelic biases of cis-regulatory element activities could be responsible for allelic gene expression29, and demonstrate that such a phenomenon is probably widespread in the human genome. These observations shed light on the importance of considering genetic variants in understanding individual-specific gene regulation. Analyses of haplotype-resolved transcriptomes and epigenomes in additional individuals and tissues should further illuminate the role of sequence variations in defining individual-specific transcriptional programs and phenotypes.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information is available in the online version of the paper.

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Author Contributions D.L., W.X., J.R.E., N.C.C. and B.R. led the data production. L.J., N.R., M.Z. and B.R. led the data analyses. L.J., N.R., S.S.-Y., Y.O., L.E., M.H., A.S. and P.R. conducted analyses. S.L. and Y.L. processed tissue samples. D.L., A.S., A.Y.L., C.A.Y., S.K. and H.Y. produced data. D.L., J.J., N.R. and B.R. wrote the manuscript.

Author Information Chip-seq and RNA-seq data sets were deposited at the Gene Expression Omnibus (GEO) under accession number GSE16256. Hi-C data sets were deposited at GEO under accession number GSE83752. Reprints and permissions information is available at www.nature.com/reprints. The authors declare competing financial interests; details are available in the online version of the paper. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to B.R. (biren@ucsd.edu).

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Extended Data Figure 1 | Active enhancers cluster along developmental lineages. a, Pie charts showing fractions of tissue-restricted and non-tissue-restricted strong enhancers and promoters. b, Hierarchical clustering with optimal leaf ordering based on all H3K27ac-marked highly active enhancers. Four major clusters are represented: early embryonic cell types (blue); a large set of meso/endoderm-derived tissues (dark green); a set consisting of ectoderm-derived brain tissues (red); and a small cluster of mesoderm cell lines (purple), which bridged the early embryonic lineages with the somatic tissues. Although TRO did not fall within any clusters, it shared the highest degree of similarity with the early embryonic cell lines. On a subsequent level, two clusters are seen separating endoderm-derived tissues (grey line) and mesoderm-derived tissues (green line). Heart tissues are denoted by yellow asterisks. c, Clustering of tissues by promoters’ histone acetylation status shows grouping of tissues that are of similar types but are less evident in germ-layer divisions than clustering of enhancers.

**Legend:**
- Blue: Early embryonic cell lines
- Green: Mesoderm derived tissues
- Brown: Endodermal derived tissues
- Red: Ectodermal derived brain tissues
Extended Data Figure 2 | Tissue-restricted enhancers are enriched for transcription factor motifs important for cell identity and/or function. Significantly enriched motifs ($P < 10^{-10}$) across all 28 tissues are divided into 29 clusters (method described in Supplementary Information). An overall $P$ value is generated for the enrichment of each tissue for each cluster. The figure illustrates $-\log(P)$ value of (a) pancreas, (b) anterior caudate and (c) liver-restricted enhancer motif enrichment for the various clusters. For ease of visualization, any cluster with $P$ values greater than 0.05 is denoted 0. Red highlighted text refers to a subset of motifs for transcription factors with literature support (see Supplementary Information) to have function in (a) the pancreas, (b) the brain and (c) the liver.
Extended Data Figure 3 | Endogenous retroviruses (ERVs) are enriched for active cis-regulatory element marks in a tissue-restricted fashion. a, A clustered heat map showing the H3K27ac enrichment (RPKM) of all mappable elements of the three classes of ERVs. b, Distribution of the Shannon entropy of H3K27ac across enhancers, promoters and three classes of ERVs is shown as a density curve, demonstrating that H3K27ac enrichment of ERVs is more tissue-restricted than promoters and slightly less than enhancers. c, Box-plots illustrating the H3K27ac enrichment of 127 mappable members of the HERV-H subfamily across all tissue/cell types. The enrichment in H1 hESCs is significantly higher than all other cell/tissues types ($P < 1.4 \times 10^{-7}$, Wilcoxon test). d, UCSC genome browser snapshots showing example of an HERV-H element harbouring H1-restricted active promoter marks, corresponding RNA-seq signal and H3K36me3 enrichment. Notably, this particular element has been annotated in Refseq as the ES cell related gene (ESRG), a human-specific long non-coding RNA gene. e, UCSC genome browser snapshots showing example of a LTR12C element harbouring TRO-restricted active enhancer chromatin marks. f, A matrix illustrating the average H3K27ac enrichment for subfamilies of class I ERVs across all cell and tissue types. LTR12C subfamily (green arrow) shows enrichment of H3K27ac across many distinct cell types and tissues.
Extended Data Figure 4 | cREDS are enriched with dynamic histone mark signatures in different tissues and have putative cis-regulatory functions. a, Heat maps showing the enrichment (RPKM) of the H3K27ac, H3K4me3 and H3K4me1 at MES-restricted enhancers (n = 650), which are predicted as promoters in other tissues, across all 28 samples. The red box highlights the histone modifications in MES cells. b, A schematic of the pGL3-enhancer vector used in these luciferase-reporter assays (top) and the activity of 10 selected cREDS with promoter signatures and a negative control region cloned 5' to the reporter gene after transfection into H1 hESCs (bottom). Luciferase activity of each region is normalized to the average activity of the negative controls. c, A schematic of the pGL3 promoter vector used in these luciferase-reporter assays (top) and the activity of 11 selected cREDS with enhancer signatures and a negative control region cloned 3’ to the reporter gene after transfection into H1 hESCs (bottom). Luciferase activity of each region is normalized to averaged activity of negative control regions. Error bars reflect standard deviation between three technical replicates.
Extended Data Figure 5 | VISTA validated enhancers also possess dynamic histone modification signatures across tissues. Example screen shots of VISTA validated enhancers and the patterns of activity in vivo are displayed along with histone modification patterns in representative tissues (adapted from VISTA enhancer browser).
Extended Data Figure 6 | cREDS show enrichment of CAGE signal and putative enhancer functions in a zebrafish reporter assay. a, UCSC genome browser screen shots show the two cREDS elements (grey shading) harbouring enhancer and promoter signatures in distinct tissue types. When compared to CAGE data sets from the FANTOM5 project, these elements show substantial overlap with transcript signals (red and blue signals indicate CAGE signal on the forward and reverse orientation, respectively). b, Selected cREDS (same elements as above) with enhancer marks in left ventricle show heart-restricted enhancer activity, as indicated by GFP expression, in 3 days post-fertilization (3 dpf) zebrafish larvae. In parallel, pT2MX negative control did not show any GFP expression. White arrow indicates location of the 3 dpf zebrafish heart. For enhancer 1, 13 out of the 38 surviving embryos showed similar patterns. For enhancer 2, 18 out of the 35 surviving embryos showed similar patterns. None of the 30 surviving embryos, injected with the control vector, showed any appreciable GFP signal in the heart. Scale bar, 50 μm.
Extended Data Figure 7 | Identification of widespread allelically expressed genes. a, Fraction of genes with allelically biased expression in each sample. y axis indicates number of samples and x axis indicates fraction of allelically biased genes among informative genes (more than 10 SNP-containing short reads). b, Distribution of fold change of allelically biased genes between P1 and P2 alleles. c, The occurrence of allelically biased imprinted and other genes is shown. x axis refers to the number of individual donors where corresponding allelically expressed genes are commonly detected. d, A density plot showing the fraction of sample-restricted genes with allelically biased expression (grey). Three tissue samples were randomly selected and sample-restricted allelically expressed genes were defined, which includes random variance effect. The random selection was repeated 10,000 times. The shaded blue box indicates the range of fractions of individual-restricted allele-biased genes in all analysed tissue types (n = 10). The fraction of sample-restricted allelically biased genes is lower than individual-restricted allele-biased genes in Fig. 2e. e, Fold change of allele-biased gene expression between two alleles is shown as scatter plot. x axis is for the fold changes in one randomly selected tissue in each donor and y axis is for the fold changes in all other remaining tissues in the corresponding donor. Allelic bias in one tissue is highly correlated with allelic bias in other tissues in the same individual. f, A histogram illustrates the proportions of allelically expressed genes in donor 2 (left) and 3 (right) defined in various numbers of tissues. The fraction of all testable genes or allelically expressed genes (y axis) is calculated for the number of tissues where they are called as active (x axis). The results indicate that the majority of allelically biased genes, as opposed to testable genes, are restricted to one or two tissue samples. KS test was performed between allele-biased genes and testable genes (P < 2.2 × 10⁻¹⁶).
Extended Data Figure 8 | Allele-biased chromatin states. a, Box-plots illustrating haplotype-resolved ChIP-seq signal enrichment on the two alleles at promoter regions. The P1 or P2 allele-biased promoter regions were defined by H3K27ac signals and then H3K4me1, H3K4me3 and H3K9me3 signals were presented for the corresponding promoter regions. All chromatin states are consistent according to the allele-biased H3K27ac patterns. KS test was performed for $P$ value calculation. b, Allelically biased enhancers were tested in thymus from donor 1 and pancreas from donors 2 and 3. H3K27ac enrichment was tested by allele-specific ChIP-qPCR. Two control enhancers were included and showed to have no allelic biases in thymus or pancreas from donor 2 (top right and bottom left, respectively).
Extended Data Figure 9 | Putative enhancers with identical genotypes in different individuals exhibit similar biases in histone acetylation. a, Scatter plots of P1 allele-biased enhancer activities for pairwise comparison of allele-biased enhancers in donor 1 (n = 85) and donor 2 (n = 4,427). x and y axes indicate P1 allele bias. b, Scatter plot of reference allele bias of enhancer activities for pairwise comparison of allele-biased enhancers in all tissues from all three donors and lymphoblastoid data sets obtained from a previous study (n = 309).
Extended Data Figure 10 | Analyses of concordant allelically biased gene–enhancer pairs. a, Frequency of allelically expressed genes according to the distance between concordantly allele-biased enhancer–gene pairs. Blue bars represent data obtained from whole-chromosome-spanning haplotype blocks while green bars represent data obtained from simulated 300-kb haplotype blocks. 56% of enhancer–gene pairs are more than 300 kb apart. b, Accumulation curve showing fraction of allelically biased genes that have at least one concordantly allelic enhancer within a given distance (x axis). Up to 83% of allelically expressed genes are within 2 Mb of a concordantly biased allelic enhancer. c, The frequency of allele-biased enhancers in donors 1, 2 and 3. y axis indicates fraction of enhancers and x axis indicates frequency of allelically biased enhancers. KS test was performed between allele-biased enhancers and testable enhancers. d, Bar plots presenting the number of enhancers overlapping with DHS QTLs and H3K27ac QTLs for allelic enhancers, testable enhancers, and random control regions. ***P < 10^{-5}.