

Correcting for relatedness in standard mouse mapping populations; and something about epistasis

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Abstract

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Author summary

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Abstract

It is well known that epigenetic modifications, such as histone modifications, and DNA methylation are a major mode of regulating gene transcription.

It is not well known how variation in epigenetic modifications across genetically distinct individuals contributes to heritable variation in gene expression.

When we map an eQTL, how much of the effect of the eQTL is mediated through epigenetic modifications?

We investigated this question in genetically diverse mice.

local imputed histone modifications matched eQTL extremely well, suggesting that a large portion of variation in gene expression mapped to local genotype is mediated through histone modifications.

In contrast percent DNA methylation is not determined by local genetics, and does not contribute to eQTLs.

Introduction

It is well established that epigenetic modifications, such as histone modifications, and DNA methylation influence gene expression [26704082, 22641018, 22781841]. Across cell types, unique combinatorial patterns of histone modifications mark chromatin states that establish cell type-specific patterns of gene expression [20657582, 21441907]. Similarly, the methylation of CpG sites around gene promoters and enhancers influences transcription in a cell type-specific manner [21701563, 20720541].

These patterns of histone modifications and DNA methylation are established during development. The result is a canonical epigenetic landscape for coordination of major patterns of gene expression for each cell type [sources about development]. As an organism ages and responds to its environment, patterns of both histone modifications [citation] and of DNA methylation change [citation]. Such changes have been linked to senescence [Horvath clock] and cancer [citations].

Epigenetic modifications coordinate the usage of a single genome to be used for many different types of cells with diverse morphology and physiology. This amazing feature of epigenetic modifications has been intensely studied, and the variation in epigenetic landscapes across cell types has been extensively documented [citations]. Less well understood, however, is the role that genetic variation plays in determining epigenetic landscapes.

Across genetically diverse populations of humans or mice, individual cell types, such as hepatocytes, or cardiomyocytes, have globally similar gene expression profiles that define their role within the greater organism. However, it is also true that across individuals, gene expression varies widely within the global constraints of cell type. This variation can increase or decrease an organism's risk of developing disease. Variation in gene expression has been extensively mapped to variation in genetic loci, or expression quantitative trait loci (eQTL). Large, coordinated efforts, such as the Genotype-Tissue Expression (GTEx) Project [32913073, 32913075] have identified and catalogued many such loci in humans, and countless independent studies have identified eQTL in mice and other model organisms.

Although the link between genetic variation and gene expression has been well studied, there is relatively little known about inter-individual variation in epigenetic modifications, and how these variations are related to variations in genotype and gene expression. The generation of a more complete picture of inter-individual variation in epigenetic modifications has the potential to increase our understanding of the mechanisms of gene regulation, provide insights into the mechanisms establishing cell type-specific epigenetic landscapes, and to improve the functional annotation of the genome as it relates to the regulation of gene expression. The vast majority of SNPs associated with human disease traits are located in non-coding regions, suggesting that they influence gene regulation, rather than protein function [citation]. However, annotation of these regions is difficult without additional genomic features, such as histone modifications and DNA methylation. Overlaying a map of variation in epigenetic features has the potential to provide a picture of how genetic variation changes functional elements, like enhancers and insulators, in the genome [citation].

Advances in chromatin immunoprecipitation (ChIP) and sequencing technologies now enable genome-wide surveys of histone modifications with relatively few cells [20077036], thus opening the door to the possibility of cataloguing epigenetic variation across cell types and individuals. Here, we performed a survey of epigenetic variation in hepatocytes across nine inbred mouse strains. We included the eight founders of the Diversity Outbred/Collaborative Cross (DO/CC) [citation] mice, as well as DBA/2J, which, along with C57Bl/6J, is one of the founders of the widely used BxD recombinant inbred panel of mice [citation]. We assayed four histone modifications (H3K4me1, H3K4me3, H3K27me3, and H3K27ac), as well as DNA methylation. We used ChromHMM [citation] to identify 14 chromatin states, classified by unique combinations of the four histone marks, and investigated the association between variation in these states and variation in gene expression across the nine strains. We separately investigated the relationship between DNA methylation and gene expression across strains.

We further investigated the relationship between epigenetic state and gene expression by imputing the 14 chromatin states and DNA methylation into a population of DO mice. We then mapped gene expression to the imputed epigenetic states to assess the extent to which eQTLs are driven by variation in epigenetic modification. We thus linked genetically controlled variation in epigenetic modifications to variation in gene expression in mice, and we provide the first resource documenting epigenetic variation across a wide panel of genetically diverse mice.

Materials and Methods

Inbred Mice

information about housing, animal use, etc.

Hepatocyte acquisition

Samples were taken from 12-week female mice of nine inbred mouse strains: 129S1/SvImJ, A/J, C57BL/6J, CAST/EiJ, DBA/2J NOD/ShiLtJ, NZO/HILtJ, PWK/PhJ, and WSB/EiJ. Eight of these strains are the eight strains that served as founders of the Collaborative Cross/Diversity Outbred mice [REF]. The ninth strain, DBA/2J, will facilitate the interpretation of existing and forthcoming genetic mapping data obtained from the BxD recombinant inbred strain panel [REF]. Mice were aged and processed in groups to maintain a steady sample preparation workflow. Mice were housed, born, and aged in the same mouse room, with uniformity in timing, diet, and all other possible conditions. Female mice were used for all experiments due to potentially confounding effects from variation in testosterone among males that can affect liver gene expression, as well general experience that female expression is less variable than male in multiple tissues. This will also facilitate the analysis of maternal effects on offspring in later studies. Three mice were used from each strain.

Liver perfusion

To purify hepatocytes from the liver cell population, the mouse livers were perfused with collagenase to digest the liver into a single-cell suspension, and then isolated using centrifugation. Mice were harvested at 9:00 AM and sacrificed by cervical dislocation. Mice were placed over a stack of paper towels in preparation to catch excess liquid, and the appendages were pinned out to hold the body in place. to keep the fur from contaminating the liver sample later, the fur was wiped down with 70% ethanol. The mouse skin was then cut open and peeled back to the appendages to allow clear access to the abdominal cavity. The fascia was cut open and back to the ribs, being careful to not nick the liver. Moving the intestines and stomach to the right side, the vena cava and hepatic portal vein should be clearly visible below the liver.

For the perfusion, a 23G x $\frac{3}{4}$ '' BD Vacutainer Safety-Lok needle (REF 367297) was attached to 1.6mm ID BioRad Tygon tubing (R-3603) connected to a Pharmacia peristaltic pump that allows a flow of up to 8 ml/min. The liver will be processed with three solutions: 5mM EGTA in Leffert's buffer, Leffert's buffer wash, and 87 CDU/mL Liberase collagenase with 0.02% CaCl₂ in Leffert's buffer. The three solutions were at 37°C before perfusion.

The needle was placed into the vena cava for the perfusion superior to the kidneys and inferior to the liver. With the peristaltic pump running slowly, the vena cava was pierced at shallow 15° angle and the needle was inserted to a shallow depth (around 2-3mm of the needle tip in the vena cava). Once the needle is inserted into the vena cava, the volume on the peristaltic pump is increased to 5-7mL/min. The liver will immediately blanch, and the hepatic portal vein is immediately severed to allow flushing of the liver.

The 1x EGTA buffer was used to flush the blood out of the liver and start the digestion of the desmosomes connecting the liver cells. To help with the perfusion, pressure was applied to the hepatic portal vein for 5 second intervals causing more solution to be forced through the liver, which can be seen visually by the liver swelling. After 35ml of the 1x EGTA solution is passed through the liver, the solution was switched to the 1x Leffert's buffer. The pump was turned off during the switch to prevent air from being sucked into the tubing while the tubing is transferred to the new

solution. To wash, 7-10ml of the Leffert's buffer was passed through the liver to flush out the EGTA, which otherwise chelates the calcium ions necessary for collagenase activity in the next buffer. The pump was turned off again to switch to the Liberase solution. To digest the liver, 25-50mL of Liberase solution (~ 4.3 wunsch units) was passed through the liver. Throughout the perfusion process, periodic pressure was applied to the hepatic portal vein to help pump the buffers more completely through the liver. As the liver was digested with the Liberase, it will swell and look soggy and limp. Over-digestion leads to increased contamination with non-hepatocyte cell types, and further reduces cell viability.

After perfusion, which takes around 15-20min to complete, the liver was carefully cut out of the abdominal cavity and placed in a petri dish with 35 mL ice-cold Leffert's buffer with 0.02% CaCl_2 . The digested liver was passed through Nitex 80 μm nylon mesh (cat #03-80/37) into a 50mL conical, using additional ice-cold Leffert's buffer with 0.02% CaCl_2 if necessary, and a rubber policeman. After the liver cells from both animals were collected, they were put through two wash and spin cycles to purify the hepatocytes and remove other types of cells. To isolate the hepatocytes, the much larger size of the hepatocyte cells was exploited in very slow 4 min, 50 x g spins that leave smaller other cell types in suspension. After each spin, the solution was decanted as waste, and the enriched cell pellet of hepatocytes was resuspended in 30ml ice-cold Leffert's buffer with 0.02% CaCl_2 . After the second spin, the solution should be almost clear, indicating that other cell types have been removed. The hepatocytes are resuspended in room temperature PBS, counted, and volume adjusted to 1×10^6 cells/mL.

We aliquoted 5×10^6 cells for each RNA-Seq and bisulfite sequencing, and the rest were cross-linked for ChIP assays. Two 5×10^6 aliquots (5mLs) of liver cells were removed into two 15mL conicals. These were spun down at 200 rpm for 5 min, and resuspended in 1200 μL RTL+BME (for RNA-Seq) or frozen as a cell pellet in liquid nitrogen (for bisulfite sequencing). Meanwhile, 37% formaldehyde in methanol (VENDOR) were added to the remaining cells to a final concentration of 1%. The cells were rotated at room temperature for 5 min to cross-link protein complexes to the DNA bound to them. After cross-linking, 10x glycine (VENDOR) is added to a final concentration of 125 mM and rotated for 5 min to quench the formaldehyde and stop cross-linking. The cells were spun down at 2000 rpm for 5 min, decanted, and resuspended in PBS to 5×10^6 cells/mL. The cells were divided into 5×10^6 aliquots in 2mL tubes. The tubes were spun down again at 5000 x g for 5 min, decanted, and the cell pellets frozen in liquid nitrogen. All cell samples were stored at -80°C until used.

Hepatocyte histone binding and gene expression assays

Hepatocyte samples from 30 treatment and control mice were used in the following assays:

1. RNA-seq to quantify mRNA and long non-coding RNA expression, with approximately 30 million reads per sample.
2. Reduced-representation bisulfate sequencing to identify methylation states of approximately two million CpG sites in the genome. The average read depth is 20-30x.
3. Chromatin immunoprecipitation and sequencing to assess binding of the following histone marks:
 - a. H3K4me3 to map active promoters
 - b. H3K4me1 to identify active and poised enhancers
 - c. H3K27me3 to identify closed chromatin

- d. H3K27ac, to identify actively used enhancers 175
- e. A negative control (input chromatin) Samples are sequenced with ~ 40 million 176
reads per sample. 177

The samples for RNA-Seq in RTL+BME buffer were sent to The Jackson Lab Gene Expression Service for RNA extraction and library synthesis. 178
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Histone chromatin immunoprecipitation assays 180

The H3K4me1 and H3K4me3 histone chromatin immunoprecipitation assays were 181
performed on cross-linked hepatocytes using similar protocols. For all histone ChIP 182
assays, the crosslinked chromatin was prepared the same way. First, the aliquot of 183
 5×10^6 hepatocyte cells was lysed to release the nuclei by rotating the sample in 184
hypotonic buffer for 20 min at 4°C . The cells were pelleted by spinning for 10min, 10K 185
 \times G, at 4°C . The cells were resuspended in 130ul MNase buffer with 1mM PMSF 186
(VENDOR) and 1x protease inhibitor cocktail (Roche VENDOR) to prevent histone 187
protein degradation, then digested with 15U of MNase. The micrococcal nuclease 188
digests the exposed DNA, but leaves the nucleosome-bound DNA intact. After 10min of 189
incubation at 37°C , the chromatin was digested into primarily mononucleosomes. This 190
was confirmed by DNA-purification of the MNase-digested chromatin run out on an 191
agarose gel, which yielded mostly 150bp fragments, and few 300bp fragments. The 192
MNase digestion was stopped by adding EDTA to 10mM, and incubating on ice for 5 193
min. The digested chromatin was purified by spinning out insoluble parts at top speed 194
for 10 min at 4°C . The chromatin was transferred to a new tube and spun again to 195
further remove impurities and reduce background in the ChIP assays. The final 196
chromatin was transferred to a fresh tube, and used immediately in the ChIP. 197

To prepare for the ChIP, $20\mu\text{L}/1 \times 10^6$ cells Dynabead Protein G beads were 198
aliquoted into an Eppendorf tube. A magnetic tube holder was used to attract the 199
beads to the wall of the tube, and then the solution was carefully pipetted off, leaving 200
only the beads behind. The beads were washed twice with buffer to prepare them for 201
binding to the antibody. For this binding step and the chromatin binding step, the 202
buffer used was either RIPA buffer for the H3K4me3 and K3K27me3 ChIPs, or ChIP 203
buffer (VENDOR) for the H3K4me1 ChIP. The ChIP buffer was gentler and less 204
stringent than RIPA buffer, which was better for the weaker binding of the H3K4me1 205
antibody that was used. The buffers were supplemented with 50 mg/mL BSA 206
(VENDOR) and 0.5 mg/mL Herring Sperm DNA, both of which are blocking agents 207
that reduce background and non-specific binding. The ChIP assays also varied in the 208
amount of input chromatin and corresponding size of the reaction that was necessary to 209
yield sufficient DNA for sequencing. H3K4me3 ChIP needed only 1.5×10^6 cells, and 210
H3K4me1 and K3K27me3 ChIP used 4×10^6 cells. To perform the ChIP, $20\mu\text{L}$ of 211
Dynabeads per 1×10^6 cells is incubated with $5\mu\text{L}$ of histone antibody for $> 20\text{min}$ in 212
 $50\mu\text{L}/1 \times 10^6$ cells RIPA (or ChIP) buffer supplemented with 50 mg/mL BSA, 0.5 213
mg/mL Herring Sperm DNA, 1xPIC, and 1mM PMSF. The antibodies used were 214
(XXX). Once the antibody was bound to the Dynabeads, the beads were washed twice 215
with $100\mu\text{L}/1 \times 10^6$ cells RIPA buffer with BSA and Herring Sperm DNA. 216

Next, the MNase-digested chromatin were added, which was at a concentration of 217
 1×10^6 cells/ $25\mu\text{L}$. The ChIP reaction was incubated overnight with rotation at 4°C , to 218
allow the histone protein to bind to the antibody, which was bound to the magnetic 219
beads. In order to calculate enrichment for each ChIP sample, a known amount (10 or 220
 $20\mu\text{L}$) of MNase-digested input chromatin was saved. 221

The next morning, the ChIPs underwent a series of washes to remove unbound 222
chromatin. The H3K4me3 and H3K27me3 ChIPs were washed 3x with $100\mu\text{L}/1 \times 10^6$ 223
cells RIPA buffer, and the H3K4me1 ChIP was washed with a low salt wash (0.1% SDS, 224

1% Triton X-100, 2mM EDTA, 20mM Tris-HCl pH 8, 150 mM NaCl), a high salt wash (0.1% SDS, 2% Triton X-100, 2mM EDTA, 20mM, Tris-HCl, pH 8, 500mM NaCl), and a LiCl wash (0.25 M LiCl, 1% IGEPAL-CA630, 1% deoxycholic acid (sodium salt), 1 mM EDTA, 10 mM Tris-HCl pH 8). After three washes, the ChIPs were washed twice with TE buffer and transferred to a new tube during the last TE wash to reduce background. At this point, the histone of interest and the histone-bound DNA fragment had been purified from the MNase-digested, cross-linked chromatin, and was bound by histone-specific antibody to the magnetic Dynabeads. In the next step, a high-salt elution buffer is used to degrade the antibody binding interactions to the beads and the histone, and concurrently, proteinase K is added to digest the protein away from the DNA-protein complexes. The ChIP was incubated with the elution buffer and proteinase K at 68°C for > 6 hours to liberate the DNA. At the same time, the saved input chromatin was also digested in the same buffer. Afterwards, the beads were removed using the magnet, and the DNA was purified using the Qiagen PCR purification kit. Quantification was performed using the Qubit quantification system, which is accurate to $0.02\text{ng}/\mu\text{L}$ and only requires a small amount of sample to measure concentration. The ChIP sample was enriched for only DNA that was bound to the histone of interest. The goal for each ChIP was to yield 10 ng of ChIP DNA for sequencing. Not all samples met this criterion, and the H3K4me1 ChIPs often had a total yield of $\sim 2\text{ng}$ of DNA.

To test the efficiency of the ChIPs, quantitative PCR using QuantiFAST was performed. Two sets of primers were used, one set in a known region of histone binding (positive control), and one set in a region without histone binding (negative control). The qPCR was performed both on the ChIP DNA and the input DNA. Then the relative enrichment of positive vs negative assays was compared between the ChIP and input DNA.

The ChIP DNA was submitted to The Jackson Lab GES service for library preparation and sequencing. Libraries were made using the Kapa Hyper Prep kit with adapters at $0.6\mu\text{M}$. The libraries were amplified by 10 cycles of PCR. These libraries were not size selected, although most fragments were ~ 150 bp due to MNase-digestion. The samples were sequenced with 40 or more million reads per sample, which is almost 2x more reads than the ENCODE project, which sequenced using 20 million reads.

Diversity Outbred mice

We used previously published data from a population of diversity outbred (DO) mice [Svenson et al. 2012] to compare to the data collected from the inbred mice. The DO population included males and females from DO generations four through 11. Mice were randomly assigned to either a chow diet (6% fat by weight, LabDiet 5K52, LabDiet, Scott Distributing, Hudson, NH), or a high-fat, high-sucrose (HF/HS) diet (45% fat, 40% carbohydrates, and 15% protein) (Envigo Teklad TD.08811, Envigo, Madison, WI). Mice were maintained on this diet for 26 weeks.

Genotyping

All DO mice were genotyped as described in Svenson et al. (2012) using the Mouse Universal Genotyping Array (MUGA) (7854 markers), and the MegaMUGA (77,642 markers) (GeneSeek, Lincoln, NE). All animal procedures were approved by the Animal Care and Use Committee at The Jackson Laboratory (Animal Use Summary # 06006).

Founder haplotypes were inferred from SNPs using a Hidden Markov Model as described in Gatti *et al.* 2014. The MUGA and MegaMUGA arrays were merged to create a final set of evenly spaced 64,000 interpolated markers.

Tissue collection and gene expression

At sacrifice, whole livers were collected and gene expression was measured using RNA-Seq as described in (Chick, Munger et al.~2016, and Tyler et al.~2017). Transcript sequences were aligned to strain-specific genomes, and we used an expectation maximization algorithm (EMASE) to estimate read counts (<https://github.com/churchill-lab/emase>).

Data Processing

Sequencing

The raw sequencing data from both RNA-Seq and ChIP-Seq was put through the quality control program FastQC. FastQC identifies problems or biases in either the sequencer run or the starting library material. The FastQC readout includes total number of reads, sequence quality, duplication level, and overrepresented sequences. All of our samples had comparable quality levels and no outstanding flags. However, the ChIP-Seq data was flagged for having a high level of duplicate reads. This can be explained by the use of MNase to shear the DNA into 150 bp fragments. If the binding positions of nucleosomes are fixed, then the MNase enzyme will cleave the DNA in the same place in multiple cells, resulting in duplicate pieces of DNA. Despite evidence that the duplication rate has a biological explanation, duplicates were removed before downstream analysis, as is typical in sequencing workflows, to avoid potential biases caused by starting libraries that have less diversity.

For the sequence analysis, reads from each sample were mapped to strain-specific pseudogenomes that integrate known SNPs from each strain. While the B6 samples were aligned directly to the reference mouse genome, the other samples were from genetically different strains. Strain-specific sequence variation in transcripts can affect alignment quality and result in biased estimates of abundance. To counteract potential strain biases, sequencing data from each strain were aligned to a custom strain pseudogenome, allowing a more precise characterization of gene expression and histone binding. The pseudogenomes were created using the EMASE computational program [REF] designed to construct customized genomes based on known SNP and indel attributes. The resulting custom genomes are called pseudogenomes, because they are based on inserting small known variations into the reference genome, but do not attempt whole genome sequencing for each strain and complete rebuild the entire genomic sequence from the scaffold up. The strain-specific pseudogenomes were then used in the Bowtie mapping algorithm to align and map reads from the RNA-Seq and ChIP-Seq experiments.

Quantifying gene expression

Once the sequencing data was mapped to the custom genomes, edgeR is used to quantify transcripts. The edgeR program uses a Trimmed Mean of M-values (TMM), which adjusts each sample for library size and RNA composition using the assumption that most genes are not differentially expressed. The output is sample read count for each of the ENSMUSG transcript ID's. Next, transcripts with less than 1 CPM in two or more replicates were filtered to remove lowly expressed genes. Also, the data were trimmed to include only protein-coding transcripts.

ChIP-Seq quantification:

After the ChIP-Seq sequencing data were mapped to the custom pseudogenomes, peaks were called in each sample using MACS 1.4.2 [REF], with a significance threshold of

$p \leq 10^{-5}$. In order to compare peaks across strains, the MACS output peak coordinates were converted to common B6 coordinates using g2g tools [REF].
 Annat's stuff to get fastq files to bam files bam to bed binarize bed files

Quantifying DNA methylation

Annat's stuff to get bed files.

Analysis

Filtering transcripts

For all gene expression data, we remove transcripts with extremely low read counts, by filtering out those whose mean read count across all individuals was less than five.

We used the R package sva [REF] to perform a variance stabilizing transformation (vst) on the RNA-Seq read counts from both inbred and outbred mice. In the inbred mice we used a blind transformation, while in the outbred mice, we included DO wave and sex in the model. For eQTL mapping, we performed rank Z normalization on the RNA-Seq read counts across transcripts from the outbred mice.

Analysis of histone modifications

Identification of chromatin states

We used ChromHMM [29120462] to identify *chromatin states*, which are unique combinations of the four chromatin modifications, for example, the presence of both H3K4me3 and H3K4me1, but the absence of the other two modifications. We conducted all subsequent analyses at the level of the chromatin state.

To ensure we were analyzing the most biologically meaningful chromatin states, we calculated chromatin states for all numbers of states between four and 16, which is the maximum number of states possible with four binary chromatin modifications (2^n). We then investigated a number of features of each state in each model: presence/absence of histone modifications, distribution patterns across the genome, and the effect of each state on gene expression. We compared chromatin states from the different models based on these analyses and selected the 14-state model. Each of these analyses, and the model comparison, are described below.

Emission probabilities

Emission probabilities are a primary output of ChromHMM (Figure XXXA). They define the probability that each histone mark is present in each detected state. Low probabilities suggest absence, or low levels of the mark, and high probabilities suggest presence. To compare states to each other and to annotate states, we declared a histone mark to be present in a state if its emission probability was 0.3 or higher.

Genome distribution of chromatin states

We investigated genomic distributions of chromatin states in two ways. First, we used the ChromHMM function OverlapEnrichment to calculate enrichment of each state around known functional elements in the mouse genome. We analyzed the following features:

- **Transcription start sites (TSS)** - Annotations of TSS in the mouse genome were provided by RefSeq [26553804] and included with the release of ChromHMM, which we downloaded on December 9, 2019 [29120462].

- **Transcription end sites (TES)** - Annotations of TES in the mouse genome were provided by RefSeq and included with the release of ChromHMM.
- **Transcription factor binding sites (TFBS)** - We downloaded TFBS coordinates from OregAnno [26578589] using the UCSC genome browser [12045153] on May 4, 2021.
- **Promoters** - We downloaded promoter coordinates provided by the eukaryotic promoter database [27899657,25378343], through the UCSC genome browser on April 26, 2021.
- **Enhancers** - We downloaded annotated enhancers provided by ChromHMM through the UCSC genome browser on April 26, 2021.
- **Candidates of cis regulatory elements in the mouse genome (cCREs)** - We downloaded cCRE annotations provided by ENCODE [22955616] through the UCSC genome browser on April 26, 2021.
- **CpG Islands** - Annotations of CpG islands in the mouse genome were included with the release of ChromHMM.

In addition to these enrichments around individual elements, we also calculated chromatin state abundance relative to the main anatomical features of a gene. For each transcribed gene, we generated a chromatin state matrix with genomic position in rows, and mouse strains in columns. Each cell contained the chromatin state assignment for a 200 base pair (bp) window, defined by ChromHMM, for each strain. We normalized these bp positions for each gene, such that they ran from 0 at the transcription start site (TSS) to 1 at the transcription end site (TES). We also included 1000 bp upstream of the TSS and 1000 bp downstream of the TES, which were converted to values below 0 and above 1 respectively.

To normalize the coordinates, we first centered all coordinates on the TSS of the gene by subtracting off the base pair position of the TSS. Centered positions were then divided by the length of the gene in base pairs from the TSS to the TES. We then binned the relative positions into 41 bins defined by the sequence from -2 to 2 incremented by 0.1. If a bin encompassed multiple positions in the gene, we assigned the mean value of the feature of interest to the bin. To avoid potential contamination from regulatory regions of nearby genes, we only included genes that were at least 2kb from their nearest neighbor, for a final set of 14048 genes.

Chromatin state and gene expression

We calculated the effect of each chromatin state on gene expression. We did this both across genes and across strains. The first analysis identifies states that are associated with high expression and low expression within the hepatocytes, and the second analysis investigates whether variation in chromatin state across strains contributes to variation in gene expression across strains.

For each transcribed gene, we calculated the proportion of the gene body that was assigned to each chromatin state. We then fit a linear model separately for each state to calculate the effect of state proportion with gene expression:

$$y_e = \beta x_s + \epsilon$$

where y_e is the rank Z normalized gene expression of the full transcriptome in a single inbred strain, and x_s is the rank Z normalized proportion of each gene that was assigned to state s . We fit this model for each strain and each state to yield one β coefficient with 95% confidence interval. The effects were not different across strains, so we averaged the effects and confidence intervals across strains to yield one summary effect for each state.

To calculate the effect of each chromatin state across strains, we first standardized transcript abundance across strains for each transcript. We also standardized the proportion of each chromatin state for each gene across strains. We then fit the same linear model, where y_e was a rank Z normalized vector concatenating all standardized expression levels across all strains, and x_s was a rank Z normalized vector concatenating all standardized state proportions across all strains. We fit the model for each state independently yielding a β coefficient and 95% confidence interval for each state.

In addition to calculating the effect of state proportion across the full gene body, we also performed the same calculations in a position-based manner. This second analysis yielded an effect of each state at multiple points along the gene body and a more nuanced view of the effect of each state.

Selecting the most biologically meaningful model

We performed the above analyses on all states from the four-state model to the 16-state model to find the most meaningful clustering of histone modifications. Across all models, the states were remarkably stable (Supplemental Figure XXX). As we increased the number of states detected by the model, new states appeared, but previously detected states were not disrupted. This stability was apparent in all state measures: emissions probability patterns, overall abundance, effect on expression, and localization along the genome. The one exception to this stability was that highly abundant state (present in 65% of transcribed genes) detected first in the four-state model was split into two distinct states in the 10-state model. These states were also highly abundant (appearing in 40% and 41% of transcribed genes), and had distinct genomic distributions and emissions probabilities (Supplemental Figure XXX). These two states remained stable with increasing numbers of clusters through to the 16-state model. States arising after the 10-state model were of lower abundance, appearing in 2% or less of transcribed genes.

All of the higher abundance states were established in the 10-state model. However, as we moved toward higher numbers of clusters, the resolution on the lower-abundance states improved in terms of the emission probability profiles, and strength of the correlation with gene expression. For example, the 14-state model better resolved a state that had appeared in the 10-state model but was not strongly correlated with gene expression. In the 14-state model, the emission patterns were closer to binary, and the strength of the correlation with expression was increased. Beyond 14 clusters, the new states identified were extremely rare (1% of transcripts or less), and were not strongly correlated with gene expression. We thus selected the 14-state model and the model with the most biologically meaningful clusters.

Analysis of DNA methylation

Creation of DNA methylome

We combined the DNA methylation data into a single methylome cataloging the methylated sites across all strains. For each site, we averaged the percent methylation across the three replicates in each strain. The final methylome contained 5,311,670 unique sites across the genome. Because methylated CpG sites can be fully methylated, unmethylated, or hemi-methylated, we rounded the average percent methylation at each site to the nearest 0, 50, or 100.

Distribution of CpG sites

We used the enrichment function in ChromHMM described above to identify enrichment of CpG sites around functional elements in the mouse genome. We further performed a

gene-based analysis of abundance similar to that in the chromatin states. As a function of relative position on the gene body, we calculated the density of CpG sites as the average distance to the next downstream CpG site, as well as the percent methylation at each site.

Effects of DNA methylation on gene expression

As with chromatin state, we assessed the effect of DNA methylation on gene expression both within strains (across genes), and across strains. We used the same linear model described above, except that y_s became the rank Z normalized percent methylation either across genes or across strains. However, unlike with the chromatin states, we only calculated the effects of DNA methylation on gene expression in a position-dependent manner.

Imputation of genomic features in Diversity Outbred mice

To assess the extent to which chromatin state and DNA methylation are responsible for local expression QTLs, we imputed local chromatin state and DNA methylation into a population of diversity outbred (DO) mice described above and in Svensson et al. 2012. We compared the effect of the imputed epigenetic features to imputed SNPs.

All imputations followed the same basic procedure: For each transcript, we identified the haplotype probabilities in the DO mice at the genetic marker nearest the gene transcription start site. This matrix held DO individuals in rows and DO founder haplotypes in columns.

For each transcript, we also generated a three-dimensional array representing the genomic features derived from the DO founders. This array held DO founders in rows, feature state in columns, and genomic position in the third dimension. The feature state for chromatin consisted of states one through 14, for SNPs feature state consisted of the genotypes A,C,G, and T (Fig XXX?).

We then multiplied the haplotype probabilities by each genomic feature array to obtain the imputed genomic feature for each DO mouse. This final array held DO individuals in rows, the genomic feature in the second dimension, and genomic position in the third dimension. This array is analogous to the `genoprobs` object in R/`qtl2` (CITE). The genomic position dimension included all positions from 1 kb upstream of the TSS to 1 kb downstream of the TES. SNP data for the DO founders in mm10 coordinates were downloaded from the Sanger SNP database [1921910, 21921916], on July 6, 2021.

To calculate the effect of each imputed genomic feature on gene expression in the DO population, we fit a linear model. From this linear model, we calculated the variance explained (R^2) by each genomic feature, thereby relating gene expression in the DO to each position of the imputed feature in and around the gene body.

Results

Gene expression varies widely and reproducibly across inbred strains of mice. This is seen as a clustering of individuals from the same strain in a principal component plot of the hepatocyte transcriptome across strains (Figure XXX). The effect of genotype on this variation can be measured in a mapping population as expression quantitative trait loci (eQTL), which associate genetic variation with variation in gene expression. In this study we investigated the extent to which variation in epigenetic modifications, such as histone modifications and DNA methylation, are associated with genetically controlled variation in gene expression.

Chromatin state overview

To investigate this association, we used ChromHMM to identify 14 chromatin states composed of unique combinations of four histone modifications in the hepatocytes of nine inbred strains of mice. Ppanel A in Figure XXX shows the representation of each histone modification across the states.

The states were distributed non-randomly around known functional elements in the mouse genome (Figure XXXB). The majority of the states were enriched around the TSS, and other TSS-related functional elements, such as promoters and CpG islands. Two states (states 2 and 1) were primarily found in intergenic regions, three states (states 9, 13, and 11) were enriched around known enhancers, and one (state 6) was enriched predominantly near the TES. The majority of these states were also associated with variation in gene expression (Figure XXXC). The colored bars in this panel show the effect of each state on gene expression variation across the inbred strains. For reference, the paired tan bars show the effect of each chromatin state on gene expression in hepatocytes. These effects are the same sign as the across-strain effects, for the most part, and tend to be stronger.

By merging histone modification patterns with enrichments near functional elements and the effects of each state on gene expression, we were able to suggest annotations for many of the 14 chromatin states (Figure XXXD). A more detailed description of these annotations is in Table XXX.

The states in Figure XXX are shown in order of their effect on expression, which helps illustrate several patterns in the data. The state with the most negative effect on gene expression, state 1, is the absence of all measured modifications. The next few states all contain the repressive mark H3K27me2, and are all associated with reduced gene expression. The states with the most positive effects on expression all have some combination of the activating marks, H3K4me3, H3K4me1, and H3K27ac, and the repressive mark is less commonly seen in these states with positive effects. Maybe unsurprisingly, we were also better able to annotate the states with strong negative effects and strong positive effects, but the states in the middle with weak effects were more difficult to annotate.

That states 1 and 2 were associated with reduced gene expression both within hepatocytes and across strains suggests that there may be differential epigenetic silencing of genes in hepatocytes across strains. Further, the majority of chromatin states were associated with variation in expression across strains, suggesting that epigenetic regulation of gene expression through histone modification may contribute substantially to variation in gene expression across genetically distinct individuals. That most states have the same effects across genes within a cell type and across strains suggests that the mechanisms that are used to regulate cell type specificity also contribute to variation in genetically distinct individuals.

Spatial distribution of epigenetic modifications around gene bodies

In addition to looking for enrichment of chromatin states near annotated functional elements, we characterized the fine-grained spatial distribution of each state around gene bodies (Figure XXXA-B). We also characterized the distribution of CpG sites and their percent methylation at this gene-level scale (Figure XXXC-D).

The spatial patterns of the individual chromatin states are shown in (Figure XXXA), and an overlay of all states together emphasizes the difference in abundance between the most abundant states (states 14, 12, and 1), and the remaining states, which were relatively rare (Figure XXXB).

Each chromatin state had a characteristic distribution pattern relative to gene bodies. For example, state 1, which was characterized by the absence of all measured histone modifications, was strongly depleted near the TSS, indicating that this region is commonly subject to histone modification. However, its abundance increased steadily to a peak at the TES. In contrast, states 12 and 14 were both concentrated at the TSS. State 12 was very narrowly concentrated right at the TSS, whereas state 14 was more broadly abundant both upstream and downstream of the TSS. Both were associated with increased expression in the inbred mice, suggesting promoter or enhancer functions. The state third state in this group of high-expressing states, state 13, was depleted nere the TSS, but enriched within the gene body, suggesting that this state may mark active intragenic enhancers.

The states with weaker effects on expression, the states with more gray shades, were also of lower abundance, but still had distinct patterns of abundance around the gene body suggesting the possibility of distinct functional roles in the regulation of gene expression.

DNA methylation also showed strong positional effects (Figure XXXC and D). Across all genes, the TSS had densely packed CpG sites relative to the gene body (Figure XXXC). As expected, the median CpG site near the TSS was consistently hypomethylated relative to the median CpG site in intergenic regions. CpG sites within the gene body were slightly hypermethylated compared to intergenic CpGs (Figure XXXD).

Spatially resolved effects on gene expression

The distinct spatial distributions of the chromatin states and methylated CpG sites around the gene body raised the question as to whether the effects of these states on gene expression could also be spatially resolved. To investigate this possibility we tested the association between both chromatin state and DNA methylation and gene expression with spatially resolved models (Methods). We tested the effect of each chromatin state on expression across genes within hepatocytes (Figure XXXA) and the effect of each chromatin state on the variation in gene expression across strains (Figure XXXB).

All chromatin states demonstrated spatially dependent effects on gene expression within hepatocytes. For about half of the states, the effects on expression were concentrated at or near the TSS, while in the other states effects were seen across the whole gene body or predominantly in the intragenic region. The direction of the effects matched the overall effects of each state seen previously. Remarkably, the spatial effects were recapitulated for almost every state when we looked across strains. That is, variation in chromatin state across strains contributed to variation in gene expression in the same manner that cell-type expression was being established. One notable exception was state 9, whose presence upregulated genes within hepatocytes, but did not contribute to expression variation across strains.

We also examined the effect of percent DNA methylation across genes within hepatocytes, and as it contributred to expression variation across the inbred strains (Figure XXX). As expected, hypomethylation at the TSS was associated with lower expression in hepatocytes. However, percent DNA methylation did not contribute at all to expression variation across strains, implying that although DNA methylation is used in gene regulation within a cell type, it it not heritable and does not contribute to variation in gene expression across genetically diverse individuals.

Imputed chromatin state explained variation in expression in diversity outbred mice

Thus far, we have used inbred strains of mice to identify correlations between local chromatin state and gene expression. However, we cannot establish causality in this population. For that we need a mapping population in which we can associate genetic or epigenetic variation at a single locus with changes in gene expression. A mapping population will also allow us to establish the extent to which variation in epigenetic factors contributes to observed expression quantitative trait loci (eQTL).

To compare the contribution of genetic and epigenetic features to eQTLs in a genetically diverse population, we imputed chromatin state, DNA methylation, and SNPs into a population of DO mice described previously [Svenson, Tyler] (Methods). Chromatin state is largely determined by local genotype, especially early in life [REF], and can thus be reliably imputed from local genotype. Further, we have shown here that local chromatin state correlates with variation in gene expression across inbred strains. DNA methylation, on the other hand, is known not to be highly heritable [REF], and thus cannot be reliably imputed from local genotype. We have also shown here that DNA methylation is not correlated with variation in gene expression across inbred strains. The imputation of DNA methylation thus serves as an estimate of a lower bound the ability of a feature imputed from local haplotype to explain gene expression in a new population.

For each transcript in the DO population, we imputed the local chromatin state across the gene body based on the gene's local founder haplotype and the chromatin state at the corresponding position in the inbred mice. We did the same for DNA methylation and SNPs (Figure XXX).

After imputing each genomic feature into the DO population, we mapped gene expression to the imputed features and calculated the variance explained. An example of the data used for the imputation and the results of the mapping are shown for the gene *Pkd2* in Figure XXX. Panels B, C, and D show the chromatin state, SNP genotype, and DNA methylation status at multiple positions along the gene body. The boundaries of the gene body along with the direction of transcription are shown by the arrow under panel A. Panel A shows the variance explained by each genomic feature at each position with a distinct symbol. The variance explained by the local haplotype is shown as the dashed line at the top of this panel. There are two particularly interesting regions in this gene. One is at the TSS and the immediately surrounding area, and the other is just downstream of the TSS. Both regions have high variation in chromatin state that are highly correlated with gene expression in the DO (blue plus signs in top panel). Just upstream of the TSS there are SNPs (red x's in the top panel) that also explain large amounts of variation in expression, although not as much as chromatin state. A similar pattern is seen at the internal region just downstream of the TSS. Percent DNA methylation does not vary across the strains in either of these critical regions, and does not contribute to variation in expression across genetically distinct individuals.

The overall distributions of variance explained by each feature across all transcripts is shown in Figure XXX. These distributions show the haplotype effect for the marker nearest each transcript compared with the maximum effect across the gene body for each of the other imputed features. Overall, local haplotype explained the largest amount of variance of gene expression in the DO ($R^2 = 0.17$). The variance explained by local chromatin state was very highly correlated with that of haplotype (Pearson $r = 0.96$) and explained almost as much variance in gene expression in the DO as local haplotype ($R^2 = 0.15$).

The mean variance explained by SNPs was lower ($R^2 = 0.13$) than that explained by haplotype and was not as highly correlated with local haplotype as chromatin state was (Pearson $r = 0.93$). DNA methylation, the lower bound for variance explained by a

feature imputed from local haplotype, explained the least amount of expression variance in the DO population ($R^2 = 0.09$), and had a much lower correlation to haplotype than either chromatin state or SNPs (Pearson $r = 0.74$).

Discussion

In this study we showed that variation in histone modifications in inbred mice mirrors genetic variation, and we further showed that this variation was highly related to variation in gene expression across strains. These observations suggest that cell type-specific patterns of histone modifications are determined by local genotype, and may be a major mechanism through which expression QTL (eQTL) are generated. This hypothesis was supported by the high concordance between chromatin state, which was imputed from local genotype, and gene expression in an independent outbred population of mice.

The high resolution of the chromatin states combined with spatial patterns of distribution and effect on gene expression offers opportunities for the annotation of functional elements in and around genes. For example, the chromatin state patterns in the gene *Pkd2*, suggest two enhancers—one at the TSS, and the other just downstream of the TSS inside the gene body. The positive effects of these putative enhancer regions in the inbred mice replicated in outbred mice suggesting that these effects are robust and contribute to variation in gene expression seen in diverse populations.

The putative enhancers are not apparent in the SNP patterns or in the patterns or DNA methylation, which suggests that chromatin modification is the mechanism through which gene expression is regulated by these regions. Further, the richness of the information in this chromatin state layer provides data with which to further annotate the effects of SNPs underlying these regions. There are SNPs throughout the gene, as seen in Figure XXX, and many of them are associated with variation in gene expression. However, while the SNPs within the putative enhancer regions may change expression by altering histone modifications placed in those regions, SNPs further downstream, which are also associated with variation in expression may work through another mechanism, such as through directly disrupting transcription, or by altering the transcript such that it is processed differently post transcriptionally. The intermediate resolution of the chromatin state between that of SNPs and haplotype thus provides a highly informative layer of information between genotype and gene expression.

In contrast to chromatin state, percent DNA methylation was not associated with variation in gene expression variation across inbred strains or in the outbred population. This was largely due to a lack of variation in methylation across strains. An example of this observation is shown in panel D of Figure XXX. Despite strain variation in both genotype and chromatin state at the TSS of *Pkd2*, DNA methylation is completely invariant. The CpG island at the TSS is unmethylated in all strains. Thus, although chromatin state appears to be highly influenced by local genotype, percent DNA methylation is not.

Similar observations have been made in human studies [33931130]. Multiple twin studies have estimated the average heritability of individual CpG sites to be roughly 0.19 [27051996, 24183450, 22532803], with about 10% of CpG sites having a heritability greater than 0.5 [24183450, 22532803, 24887635]. Trimodal CpG sites, i.e. those with methylation percent varying among 0, 50, and 100%, have been shown in human brain tissue to be more heritable than unimodal, or bimodal sites ($h^2 = 0.8 \pm 0.18$), and roughly half were associated with local eQTL [20485568]. Here, we did not see an association between trimodal CpG sites and gene expression across strains (Supplemental Figure XXX).

The diversity in the effects observed in the 14 chromatin states highlights the

importance of analyzing combinatorial states as opposed to individual histone modifications. To illustrate this point, consider the three states with the largest positive effects on transcription. Each of these three states had a distinct combination of the three histone marks associated with transcriptional activation: H3K4me1, H3K4me3, and H3K27ac. State 12 was characterized by high levels of H3K4me3 and H3K27ac, and low levels of H3K4me1. State 13 was characterized by high levels of H3K4me1 and H3K27ac, and low levels of H3K4me3. And state 14 was characterized by high levels of all three activating marks (Figure XXX). Although all three states were associated with increased gene expression, each had a completely distinct spatial distribution. State 12 was distributed in a very narrow band centered on the TSS, while state 14 was distributed across a much broader region centered upstream of the TSS. State 13 had a completely different distribution – it was depleted at the TSS, and most abundant within the gene body and near the TES. This variation in spatial distribution was mirrored in the spatial effects on transcription. State 12, which we annotated as an active promoter, was positively associated with transcription when it was present at the TSS. In contrast, states 13 and 14, which we annotated as enhancers, were associated with increased transcription when present anywhere in the gene body (Figure XXX). We would not be able to detect such patterns if analyzing any of the histone modifications in isolation. These results highlight the complexity of the histone code and the importance at analyzing combinatorial states.

Some of the histone mark combinations had counterintuitive effects, which raises the intriguing possibility of identifying new modes of expression regulation through histone modification. State 6 was characterized by high levels of H3K4me3 and low levels of the other three modifications. It appeared in 8% of transcribed genes and had a consistent negative effect on transcription just downstream of the TSS in both inbred and outbred mice.

The modification H3K4me3 is most frequently associated with increased transcriptional activity [citation], so the association with state 6 with reduced transcription is a deviation from the dominant paradigm. This physical distribution of this state is also interesting. It is depleted at the TSS, and enriched just upstream and just downstream of the TSS (Figure XXX). It is also enriched just downstream of the TES, although it does not appear to influence transcription at this location (Figure XXX). The group of genes marked by state 6 were enriched for functions such as stress response, DNA damage repair, and ncRNA processing suggesting that this state may be used to regulate subsets of genes involved in responses to environmental stimuli.

Other states of note included two bivalent states, which are states that combine an activating histone modification and a repressing histone modification [citation]. We detected a poised enhancer state (state 3) and a bivalent promoter state (state 4). Both states were associated with downregulation across inbred strains when present near the TSS. This effect, however, did not replicate in the outbred mice, perhaps because the effect was too weak to detect given the number of animals in the population.

Both bivalent promoters and poised enhancers are dynamic states that change over the course of differentiation and in response to external stimuli [citation]. Bivalent promoters have been studied primarily in the context of development. They are abundant in undifferentiated cells, and are typically resolved either to active promoters or to silenced promoters as the cells differentiate into their final state [23788621, 22513113]. These promoters also been shown to be important in the response to changes in the environment. Their abundance increases in breast cancer cells in response to hypoxia [27800026]. Poised enhancers are also observed during differentiation and in differentiated cells [32432110]. In concordance with these previous observations, the genes marked by states 3 and 4 were enriched for developmental vascular development and morphogenesis. That we identified these states in differentiated hepatocytes may

indicate that a subset of developmental genes retain the ability to be activated under certain circumstances, such as during liver regeneration in response to damage. It is also possible these states were induced in the inbred strains in response to stress, rather than genetically coded. This could explain why the negative effect on gene expression did not replicate in the outbred mice. However, given that we detected this state in all nine inbred strains in relatively equal proportions, this latter hypothesis seems less likely.

for the majority of transcripts we still only explain about 10% of the variance with haplotype and local epigenetics other variation explained by more complex genetic architecture? stochasticity? other genomic features?

There was one state, state 9, that was present more often in highly expressed genes in hepatocytes than in more lowly expressed genes, but had no effect on variation in gene expression across strains. This is another interesting pattern that may indicate a chromatin state that marks genes that need to be highly expressed in a cell type-specific manner, but does not contribute to across-strain variation in expression. The genes marked by state 9 were enriched for

which was characterized by high levels of H3K4me1 and low levels of the other measured modifications.

intragenic enhancer: abundance and positive effect peak inside gene body effect carries all the way through to DO mice enrichments: cellular macromolecule localization, cellular protein localization, intracellular transport, establishment of protein localization, protein transport, RNA processing, cellular response to stress.

intragenic enhancer that has effect in hepatocytes, but not across strains: genes in hepatocytes with this state in the gene body are more highly regulated than genes without this state. The effect is quite strong in hepatocytes, but does not extend to variation across strains, even in the inbred mice. This state marks genes that are important in hepatocytes, but are not differentially regulated across strains.

enrichments: small molecule metabolic process, oxoacid metabolic process, organic acid metabolic process, carboxylic acid metabolic process, lipid metabolic process, organic substance catabolic process, monocarboxylic acid metabolic process

Annotation of chromatin states

Make this section into a table

We identified 14 chromatin states corresponding to 14 distinct combinations of histone modifications (Figure XXXA). To annotate these states to functional elements, we combined previously known annotations with functional enrichments and relationship to gene expression. The characterizations are summarized in Figure XXX. Figure XXXA further shows the relative abundance of each state in and around the gene body. This high-resolution image of abundance helped further refine the annotations of each state. Figure XXXB shows that overall states 1 and 7 were the most abundant states with state 7 being highly enriched at the TSS, and state 1 being strongly depleted at the TSS, but enriched within the gene body and in intragenic spaces. We describe the reasoning behind the annotation of each state below:

State 1 - heterochromatin was characterized by the absence of all measured marks, enrichment in intergenic regions, and strong downregulation of gene expression. This state was strongly depleted at the TSS of expressed genes (Figure XXXA), but the most abundant state in the gene body and outside the gene body. This state may multiple different states that could be resolved with the measurement of more histone modifications. For example, intergenically, state 1 may mark heterochromatin, which is characterized by H3K9 trimethylation [12867029], which was not measured here. However, state 1 was also highly abundant in the gene bodies of expressed genes, but was associated with reduced expression. This could suggest differential distribution of

heterochromatin across strains, or could represent an additional transcriptionally repressive state.

State 2 - repressed chromatin was characterized by the presence of H3K27me3, which has been shown previously to correlate with transcriptional silencing [REF]. This state was not enriched in any particular functional element, but was associated with strong downregulation of transcription.

State 3 - poised enhancer was primarily characterized by the presence of H3K27me3, a mark associated with polycomb silencing [REF], and H3K4me1 a mark associated with enhancers [REF]. The co-occurrence of these opposing marks has previously been associated with a functional element known as a poised enhancer [21160473].

This element has been studied mostly in the context of development. Bivalent promoters are abundant in undifferentiated cells, and are resolved either to active promoters or silenced promoters as the cells differentiate into their final state [REF]. These promoters have also been shown to be important in the response of cancer cells to environmental disturbances such as hypoxia [REF]. The presence of bivalent promoters in adult mouse hepatocytes is interesting. They may mark genes poised for expression during liver regeneration, or for responding to a particular environmental stimulus. There were XXX genes that were marked with this bivalent promoter state at the TSS across all strains. This group of genes was enriched for developmental processes as well as alcohol metabolism (Fig? Table?).

State 4 - intragenic enhancer was characterized by the presence of H3K4me1, which is known to mark cell type-specific enhancers, both active and poised [REF]. The presence of H3K4me1 alone, in the absence of H3K27ac, as it occurs in state 4, has been shown to mark inactive, or poised enhancers [21106759]. The addition of H3K27ac can then activate the enhancer to increase transcription. When present within the gene body, this state acts as an intragenic enhancer, which acts as an alternative promoter, and can be transcribed bidirectionally to produce short RNAs known as eRNA [20393465]. This state was modestly enriched in known enhancers and was associated with slightly increased gene expression. The presence of H3K4me1 in the absence of H3K4me3 has been shown to mark intragenic enhancers and to be associated with increased transcription, as these regions can be transcribed independently of the full gene [Kowalczyk et al. 2012]. We annotated this state as a weak enhancer.

State 5 - active enhancer was characterized by the co-occurrence of H3K4me1, which marks cell type-specific enhancers, and H3K27ac, which specifically marks active enhancers [21106759, 21160473]. This state was strongly enriched in known enhancers, and its presence had a strong positive effect on transcription. We thus annotated this state as a strong enhancer.

state 0100 (just H3K4me3 absence of others) - interesting abundance pattern. Just up and downstream of the TSS, and just downstream of the TES. Associated with slight downregulation of expression. Could be antisense transcription [22768981]? This fits the negative association, but the position of the negative association is just downstream of the TSS. can't really find any other sources on this. The probably with most studies is that they are looking only at a single mark, and not noting the absence of the other three marks.

state 0110 -

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Data and Software Availability

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All data used in this study and the code used to analyze it are available as part of a reproducible workflow located at... (Figshare?, Synapse?).

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Supplemental Figure Legends

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Fig 1

Supplemental Table Descriptions

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Fig 2. Correlations between traits and the first PC of the kinship matrix.

References

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