Genetic and Epigenetic Fine Mapping of Complex Trait Associated Loci in the Human Liver

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Deciphering the impact of genetic variation on gene regulation is fundamental to understanding common, complex human diseases. Although histone modifications are important markers of gene regulatory elements of the genome, any specific histone modification has not been assayed in more than a few individuals in the human liver. As a result, the effects of genetic variation on histone modification states in the liver are poorly understood. Here, we generate the most comprehensive genome-wide dataset of two epigenetic marks, H3K4me3 and H3K27ac, and annotate thousands of putative regulatory elements in the human liver. We integrate these findings with genome-wide gene expression data collected from the same human liver tissues and high-resolution promoter-focused chromatin interaction maps collected from human liver-derived HepG2 cells. We demonstrate widespread functional consequences of natural genetic variation on putative regulatory element activity and gene expression levels. Leveraging these extensive datasets, we fine-map a total of 74 GWAS loci that have been associated with at least one complex phenotype. Our results reveal a repertoire of genes and regulatory mechanisms governing complex disease development and further the basic understanding of genetic and epigenetic regulation of gene expression in the human liver tissue.

Introduction

The liver has a central role in detoxification of endogenous and exogenous toxins, synthesis of essential proteins, and regulation of carbohydrate, lipid, and drug metabolism. As such, the liver is associated with a diverse range of clinically important human traits and was recently reported as one of the most critical tissues for explaining cellular mechanisms at loci revealed by genome-wide association studies (GWASs). GWASs have been effective at providing robust, but imprecise, information about genetic risk factors of complex human diseases. These studies have revealed that most variation associated with complex human phenotypes do not alter protein-coding sequences, making causal variant and trait-relevant gene identification a considerable challenge. Characterization of the regulatory functions of non-coding regions is the first key step toward linking non-coding regions to disease biology. Large-scale efforts such as the Encyclopedia of DNA Elements (ENCODE) and the NIH Roadmap Epigenomics consortia have made major contributions to this end. It remains an important priority to obtain such data across many individuals, to characterize the extent of between-individual variation in the activity of regulatory elements, and to identify the genetic determinants of such differential activity. Discovering genotype-dependent non-coding functional activity can help to fine map and reveal mechanisms underlying complex trait associations. Performing such studies at genome-wide scale in large numbers of human tissues is challenging and therefore has been limited to those performed in easily accessible lymphoblastoid cell lines or blood cell types. Here, we quantify regulatory element activity in the human liver across multiple individuals and integrate these findings with genome-wide gene expression data collected from the same human liver tissues, high-resolution promoter-focused chromatin interaction maps collected from human liver-derived HepG2 cells, and GWAS summary statistics for 20 commonly studied phenotypes with variable levels of suggested causality manifesting in the liver. We identify 2,625 genes and 972 regulatory elements with genotype-dependent activity in the human liver and fine-map a total of 74 GWAS loci that have been associated with at least one complex phenotype. Overall, we provide a unique resource that contributes to basic understanding of genetic and epigenetic regulation of gene expression in the human liver tissue and highlight the benefits of integrating multiple cellular traits for the identification and characterization of disease-relevant genes, regulatory elements, and variants.

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Material and Methods

Study Subjects

Penn Cohort 1

Samples in this cohort were prospectively collected between August 2014 and February 2015 at the Penn Transplant Institute. The cohort was comprised of 50 liver donors (49 deceased, 1 living). Sex and age of the donors were reported as 19 females and 31 males aged between 6 and 77 years old.

Penn Cohort 2

Between January 2012 and August 2014, ~25 mg of liver needle biopsy samples were collected from deceased donors prior to transplantation surgery at the Penn Transplant Institute. All samples were stored in RNAlater. For this study, 96 samples were chosen based on cold ischemic time (138–320 min) and reported sex (48 females, 48 males). Age of the donors ranged between 7 and 75 years old.

GTEx Cohort

Complete description of the Genotype-Tissue expression (GTEx) cohort was published previously.23,24 In this study, 96 individuals (33 females, 63 males; age range 21–68) with genotype and liver gene expression data were included. 37 of the subjects were organ donors and 59 were postmortem. Liver needle biopsy samples from each subject were obtained in two centers. Samples were preserved in PAXgene tissue kits and shipped to the GTEx Laboratory Data Analysis and Coordinating Center LDACC at the Broad Institute for processing.25 All GTEx datasets used in this study were from GTEx Analysis Releasev6p.25

ChIP-Seq Experimental Protocol

Penn Cohort 1

Between 40 and 900 mg of liver wedge biopsies were obtained from each donor prior to transplantation surgery at the Penn Transplant Institute. Flash frozen liver wedge biopsies were processed in a total of 8 batches (six/eight randomized samples per day). On each tissue preparation day, 20 mg of tissue from each liver sample was cut, placed in 1 mL of RNAlater, and flash frozen for isolation of DNA and RNA at a later date. When available, 120 mg of tissue from each subject was processed for the chromatin immunoprecipitation (ChIP) experiment. From 33 subjects, 120 mg of tissue could be used. Tissue from the remaining 17 subjects was limited, so the largest amount available was used (ranging between 20 and 110 mg). The tissue was cut into small pieces (~1 mm3), washed with PBS, and fixed with 1% formaldehyde for 5 min at room temperature. Nuclei were prepared with the Covaris truChIP Tissue Chromatin Shearing Kit with SDS Shearing Buffer according to manufacturer’s recommendations. Chromatin was sheared for 14 min at 5% duty cycle, 140 Watts peak incident power, and 200 cycles per burst using a Covaris S220 Focused-ultrasonicator. Shearing efficiency was assessed using the Agilent High Sensitivity DNA kit and chromatin concentration was determined using a NanoDrop Spectrophotometer. From each subject, a 0.5 μg aliquot of sheared chromatin was kept aside to be used as input chromatin. Each immunoprecipitation was performed using 9 μg of sheared chromatin and 5 μg of antibody (H3K27ac:ab4729, H3K4me3:ab8580) with an overnight incubation at 4°C following the Magna ChIP A/G Chromatin Immunoprecipitation Kit protocol. After elution and reverse-cross-linking of protein-DNA complexes, DNA was cleaned with a QIAGEN QIAquick PCR Purification Kit and quantified using the Agilent High Sensitivity DNA kit. 40 H3K27ac and 45 H3K4me3 samples yielded sufficient DNA (≥2 ng) to generate sequencing libraries. 2 or 5 ng of immunoprecipitated and input DNA was used to generate sequencing libraries using the NEBNext ChIP-Seq Library Prep Master Mix Set for Illumina. Libraries were sequenced to generate 100 bp single-end reads on Illumina HiSeq2500 instruments at the Penn Next-Generation Sequencing Core.

RNA-Seq and Genotyping Experimental Protocol

Penn Cohort 1

RNA and DNA were extracted in a total of 4 batches (12 or 14 randomized samples per day) using QIAGEN’s AllPrep DNA/RNA/ miRNA Universal Kit. Barcoded, strand-specific, polyA+ selected RNA-seq libraries were generated using the Illumina TruSeq Stranded mRNA kit. Quality of each library was assessed using the Agilent Bioanalyzer High Sensitivity DNA Kit. Libraries were then pooled into one group and sequenced to generate 125 bp paired-end reads on Illumina HiSeq2500 instruments at the Penn Next-Generation Sequencing Core. DNA was genotyped using Illumina HumanCoreExome arrays at the Center for Applied Genomics Core at the Children’s Hospital of Pennsylvania.

Penn Cohort 2

RNA and DNA extraction method and library preparation was identical to that of Penn Cohort 1. Libraries were pooled into two groups of 48 samples and sequenced to generate 125 bp paired-end reads on Illumina HiSeq2500 instruments at the Penn Next-Generation Sequencing Core. DNA was genotyped using Illumina HumanCoreExome arrays at the Center for Applied Genomics Core at the Children’s Hospital of Pennsylvania.

GTEx

RNA was extracted from 96 human liver samples as described previously.26 Non-strand specific, polyA+ selected RNA-seq libraries were generated using the Illumina TruSeq protocol. Libraries were sequenced to generate 76 bp paired-end reads. DNA was extracted from whole blood using the QIAGEN Gentra Puregene method and genotyped using the Illumina Human Omni 2.5M and 5M-Quad BeadChip as described previously.26

Genome-wide Promoter-Focused Capture-C Experimental Protocol

Cell Fixation for Chromatin Capture

The protocol used for cell fixation was similar to previously published methods.27 HepG2 cells were collected and single-cell suspension was made with aliquots of 107 cells in 10 mL media (RPMI + 10% FCS). 540 µL 37% formaldehyde was added and incubation was carried out for 10 min at room temperature in a tumbler. The reaction was quenched by adding 1.5 mL, 1 M cold glycine (4°C). Fixed cells were centrifuged for 5 min at 1,000 x g at 4°C, and supernatant was removed. The pellets were washed in 10 mL cold PBS (4°C) by centrifugation for 5 min at 1,000 x g at 4°C. Supernatant was removed and cell pellets were resuspended in 5 mL of cold lysis buffer (10 mM Tris [pH 8], 10 mM NaCl, 0.2% NP-40 [Igepal] supplemented with protease inhibitor cocktails). Resuspended cells were incubated for 20 min on ice and centrifuged to remove the lysis buffer. Finally, the pellets were resuspended in 1 mL lysis buffer and transferred to 1.5 mL Eppendorf tubes prior to snap freezing (ethanol/dry ice or liquid nitrogen). Cells were stored at ~80°C until they were thawed again for digestion.

3C Library Generation

For preparation of initial 3C libraries, 10 million cells were harvested and fixed. Cells were thawed on ice and spun down, and
the lysis buffer was removed. The pellet was resuspended in water and incubated on ice for 10 min, followed by centrifugation and removal of supernatant. The pellet was then resuspended with 20% SDS and 1 x NEBuffer DpnII and incubated at 37°C for 1 h at 1,000 rpm on a MultiTherm (Sigma-Aldrich). Triton X-100 (at 20% concentration) was added and the pellet was incubated for another 1 h. After the incubation, 10 μL 50 U/μL DpnII (NEB) was added and left to digest for 8 h. An additional 10 μL DpnII was added and digestion was left overnight at 37°C. The next day, another 10 μL of DpnII was added and incubated for an additional 3 h. The chromatin was then ligated overnight (8 μL T4 DNA Ligase, HC ThermoFisher [30 U/μL]; with final concentration, 10 U/μL) and shaken at 16°C at 1,000 rpm on the MultiTherm. The next day, an additional 2 μL T4 DNA ligase was spiked in to each sample and incubated for 3 more hours. The ligated samples were de-crosslinked overnight at 65°C with Proteinase K (Invitrogen) and the following morning incubated for 30 min at 37°C with RNase A (Millipore). Phenol-chloroform extraction was performed, followed by an ethanol precipitation overnight at −20°C and then washed with 70% ethanol. Digestion efficiencies of 3C libraries were assessed by gel electrophoresis on a 0.9% agarose gel and quantitative PCR (SYBR green, Thermo Fisher).

**Capture-C**

Custom capture baits were designed using Agilent SureSelect library design targeting both ends of DpnII restriction fragments encompassing promoters (including alternative promoters) of all human coding genes and non-coding RNAs (antisense RNA, snRNA, miRNA, snoRNA, and lincRNA), totaling 36,691 RNA baited fragments. The capture library design successfully covered 95% of the coding gene promoters and 88% of the non-coding RNA promoters. Custom capture bait design failed for 5% of the coding genes, which were either duplicated genes or contained highly repetitive DNA in their promoter regions. The isolated DNA of the 3C libraries generated by DpnII digestion and ligation was quantified and competitive DNA in their promoter regions. The isolated DNA of the 3C libraries generated by DpnII digestion and ligation was quantified and competitive DNA in their promoter regions. The isolated DNA of the 3C libraries generated by DpnII digestion and ligation was quantified and competitive DNA in their promoter regions.

The library was sheared in dH2O using a QSonica Q800R to an average DNA fragment size of 350 bp. QSonica settings used were 60% amplitude, 30 s on, 30 s off, 2 min intervals, for a total of 5 intervals at 4°C. After shearing, DNA was purified using AMPureXP beads (Agencourt), the concentration was checked via Qubit and DNA size was assessed on a Bioanalyzer 2100 using a 1000 DNA Chip. SureSelect XT Library Prep Kit (Agilent) was used to repair DNA ends and for adaptor ligation following the standard protocol. Excess adaptors were removed using AMPureXP beads. Size and concentration were checked again before hybridization. 1 μg of ligated library was used and the standard protocol of the SureSelect XT capture kit was followed to obtain the custom designed Capture-C library. The quality of the captured library was assessed using both Qubit fluorometer and Bioanalyzer's high sensitivity DNA chip. Each SureSelect XT library was initially sequenced on one lane of HiSeq 4000 machine to generate 100 bp paired end reads for QC purposes. All Capture-C libraries were then sequenced three at a time on an S2 flow cell on an Illumina NovaSeq machine, generating ~1.6 billion paired-end reads per sample.

**ChIP-Seq Data Processing**

**Penn Cohort 1**

Quality of the raw sequence data was assessed using FastQC. Low-quality base calls and sequencing adapters were trimmed using Trim Galore! with the following parameters: -stringency 5 -length 50 -q 20. Reads were then aligned to the reference human genome (hg19) using the BWA-MEM algorithm.28 Aligned reads were sorted and filtered based on a minimum mapping quality of 10 using SAMtools-1.3.1.29 MACS230 was used to call peaks for each individual ChIP data using the following parameters:–nomodel–extsize 147 -q 0.01 and the corresponding input data as control. Samples that both had fraction of reads in peaks (FRIP) ≥ 1% as suggested previously31 and that displayed a significant overlap (i.e., right tailed Fisher’s p < 10−6 and at least 2-fold enrichment) with ENCODE DNase hypersensitive sites (the ENCODE DNase hypersensitive site master list generated by the ENCODE Analysis Working Group downloaded in October 2016) were retained for the downstream analyses (Table S1). See Figure S1 for the heatmap plot of Spearman’s correlation of normalized and averaged ChIP-seq read counts for 27 samples that passed the ChIP-Seq QC thresholds. To generate Figure S1, deepTools32 was used to normalize the ChIP-Seq read counts to 1× depth of coverage while excluding chromosome X and average scores were calculated based on 10 kb bins that consecutively cover the entire genome.

To define the final set of ChIP-Seq peaks, ChIP-Seq data from biological replicates (n = 9 for H3K4me3 and n = 18 for H3K27ac) as well as their corresponding input data (n = 9 for Input of H3K4me3 and n = 18 for Input of H3K27ac) were pooled into separate groups. See Figure S2 for heatmap and profile plots of read density signal around TSS (based on GENCODE v19 annotations) and Figure S3 for correlation between read density signal around TSS and gene expression levels. MACS230 was used to call the peaks on the pooled ChIP-seq data of 9 and 18 individuals respectively while using the corresponding input data as control. Among peaks that were called, 68,600 H3K4me3 and 131,293 H3K27ac peaks that have a mean read count of at least 20 were included in further analyses. See Table S2 for chromosomal positions of the peaks, Figure 2A for distribution of peak lengths, and Figure S4 for genomic annotation of ChIP-Seq peaks.

Genomic annotations were obtained using Bioconductor’s GenomicFeatures package33 and based on GENCODE v.19 annotations. H3K4me3 peaks were significantly enriched near promoter regions (≤ 3 kb to TSSs) relative to 1,000 sets of randomly selected size-matching regions of the genome (one-side Fisher’s exact test p < 2.2 × 10−15 when the observed overlap was compared with the mean overlap of 1,000 permutations). H3K4me3 and H3K27ac peaks identified also displayed significant enrichment for ENCODE DNase and FAIRE open chromatin regions as well as ENCODE H3K4me3 and H3K27ac sites in HepG2 cells (p < 2.2 × 10−15; Figure S5). Links to ENCODE datasets used are included in Table S5. ENCODE datasets were intersected with liver histone peaks as well as 1,000 sets of randomly selected size-matching regions. Fisher’s exact test was used to compare the observed number of overlap with the mean overlap of 1,000 sets of randomly selected size-matching regions.

**RNA-Seq Data Processing**

**Penn Cohort 1**

One outlier sample with fewer than one million reads was excluded from the analysis. Quality of the raw sequence data was assessed using FastQC. Low-quality base calls and sequencing adapters were trimmed using Trim Galore! with the following parameters: -stringency 5 -length 50 -q 20 –paired. Trimmed reads were aligned to the reference human genome (hg19) as implemented in STAR aligner34 using (1) genome indexes based on GENCODE v.19 annotations and (2) genome indexes based on discovered as well as expressed and annotated splice junctions. Specifically, STAR v.2.5.35 was run in two-pass...
mode using the following parameters: --outFilterMultimapNmax 10 --outFilterMismatchNmax 10 --outFilterMismatchNoverNmax 0.3 --alignIntronMin 21 --alignIntronMax 0 --alignMatesGapMax 0 --alignSJoverhangMin 5 --twopassMode Basic --twoopass1ReadsN 500000000 --sjdbOverhang 124. Aligned reads were sorted and filtered to retain only primary aligned reads using SAMtools-1.3.1.29 See Figure S6 for a histogram of number of primary aligned reads. RSEM35 was used to estimate gene-level expression as transcripts per million (TPM). 19,133 genes with RSEM expected read count of >6 and TPM of >0.1 in at least 10% of the subjects were defined as expressed. TPM values of the expressed genes were natural log transformed after adding a pseudocount of 1. After log transformation, expression values were quantile normalized between individuals across all expressed genes. For each gene, expression values were then inverse quantile normalized to a standard normal distribution across individuals.

**Penn Cohort 2**

RNA-seq data of Penn Cohort 2 was processed the same way as in Penn Cohort 1. 19,537 genes with RSEM expected read count of >6 and TPM of >0.1 in at least 10% of the subjects were defined as expressed in this cohort.

**GTEx**

Low-quality base calls and sequencing adapters were trimmed using Trim Galore!. Trimmed reads were aligned to the reference human genome (hg19) as implemented in STAR aligner34 using (1) genome indexes based on GTEx's GENCODE v.19 gene level annotations; gencode.v19.genes.v6p_model.patched_contigs.gtf and (2) genome indexes based on discovered as well as expressed and annotated splice junctions. STAR v.2.5.44 was run in 2-pass mode using the same parameters as in Penn Cohorts 1 and 2 except the parameter-sjdbOverhang 75 to correspond to the RNA-seq read length of this cohort. Similarly, aligned reads were sorted and filtered to retain only primary aligned reads using SAMTools-1.3.1.29. RSEM45 was used to estimate gene-level expression as TPM. 22,415 genes with RSEM expected read count of >6 and TPM of >0.1 in at least 10% of the subjects were defined as expressed.

**Genotype Data Processing**

**Penn Cohort 1**

Genotype data were subjected to standard QC checks using whole-genome association analysis toolset PLINK.36 First, genetic sex of individuals was compared to the self-reported sex. Out of 50 subjects, 1 had inconsistency between self-reported (male) and genotyped (female) sex, but the subject was retained in the study because genotype data were concordant when genotypes based on genotyping array and RNA-seq data were compared. Next, variants with HWE p < 10^-6 and variants with more than 5% missing rate were excluded. QC’ed genotype data were phased and imputed with SHAPEIT227 and IMPUTE2,38 respectively, using multi-ethnic panel reference from 1000 Genomes Project Phase 3.37 Following imputation, variants with HWE p < 10^-6, missing rate > 5%, minor allele frequency (MAF) < 5%, and imputation info score < 0.4 were excluded. This yielded in a total of 4,584,583 imputed variants.

**Penn Cohort 2**

Genotype data of Penn Cohort 2 was processed the same way as in Penn Cohort 1. Out of 96 subjects, 2 had inconsistencies between their self-reported (female) and genotyped (male) sex. Both subjects were retained in the study after making sure the genotypes based on genotyping array and RNA-seq data were concordant. After imputation and QC checks, 4,541,981 variants were retained.

**GTEx**

Genotype data was phased and imputed as described previously.25 Variants with HWE p < 10^-6, missing rate < 5%, MAF < 5%, and imputation info score < 0.4 were excluded. 5,598,884 variants were retained after imputation and QC filtering.

**Genome-wide Promoter-Focused Capture-C Data Processing**

Paired-end reads were pre-processed with the HICUP pipeline,40 with bowtie2 as aligner and hg19 as reference genome. Significant interactions at 1-DpnII fragment resolution were called using CHiCAGO,41 an open-source package that is commonly used for detection of robust chromatin-chromatin interactions. For this analysis, CHiCAGO was run using default parameters except for binsize, which was set to 2,500. The 4-cutter restriction enzyme, DpnII, yields high-resolution fragments (median fragment size = 264 bp, mean size = 433 bp) compared to the HindIII 6-cutter (median fragment size = 2,274 bp, mean fragment size = 3,697 bp), which is commonly used in comparable Hi-C-based approaches, but sequencing reads are distributed across many fragments, leaving fewer reads available per fragment to call significant promoter contacts, especially when further from the bait. In order to identify additional distal contacts, we also called interactions at the lower 4-DpnII fragment resolution (median fragment size = 1,440 bp, mean fragment size = 1,736 bp), which is still substantially higher than the HindIII resolution. To this end, we proceeded as described in Chesi et al.,42 and according to recommendations in the CHiCAGO vignette.43 Namely, we generated artificial .baitmap and .rmap files where DpnII fragments were grouped into four consecutively and used these files to run CHiCAGO with default parameters, except for binsize, which was set to 10,000 and removeAdjacent, which was set to False. Results from the two resolutions were merged by taking the union of the interaction calls at either resolution and removing any 4-fragment interaction which contained a 1-fragment interaction.

**Estimating Population Structure**

Principal component analysis (PCA) as implemented in EIGENSOFT37 was performed using the genotype data of each cohort in aggregate with HapMap Phase 3 genotype data from 1,184 individuals from 11 populations (ASW, African ancestry in Southwest USA; CEU, Utah residents with Northern and Western European ancestry from the CEPH collection; CHB, Han Chinese in Beijing, China; CHD, Chinese in Metropolitan Denver, Colorado; GIH, Gujarati Indians in Houston, Texas; JPT, Japanese in Tokyo, Japan; LWK, Luhya in Webuye, Kenya; MEX, Mexican ancestry in Los Angeles, California; MKK, Maasai in Kinyawa, Kenya; TSI, Toscani in Italia; YRI, Yoruba in Ibadan, Nigeria).44 In Penn Cohort 1, 34 of 50 individuals clustered with the HapMap European populations, 12 of them clustered with the HapMap African populations, and the remaining 4 individuals displayed mixed genetic ancestry (Figure S7). In Penn Cohort 2, 62 of 96 individuals clustered with the HapMap European populations, 24 of them clustered with the HapMap African populations, and the remaining 10 individuals displayed mixed genetic ancestry (Figure S8). In GTex, 81 of 96 individuals clustered with the HapMap European populations, 12 of them clustered with the HapMap African populations, 1 individual clustered with the HapMap Asian populations, and the remaining 2 individuals displayed mixed genetic ancestry (Figure S9).
Mapping cis-Expression Quantitative Trait Loci (cis-eQTLs)

cis-eQTLs were mapped by linear regression as implemented in FastQTL v2.184. Associations between total expression level (normalized TPM values) of each autosomal gene and variants within 1 Mb of the transcription start site (TSS) were tested within each cohort while adjusting for sex, first three genotype-based PCs, and PEER factors. In the GTEx cohort, genotyping platform was additionally included as a covariate in eQTL mapping. The most suitable effective number of PEER factor was determined to be 5, 22, and 16 for Penn Cohort 1, Penn Cohort 2, and GTEx cohorts, respectively (Figure S10). Nominal p values between each variant and gene pair within 1 Mb of the TSS were calculated by measuring the Pearson product-moment correlation coefficients and using standard significance tests for Pearson correlation. To identify the most significantly associated variant per gene, adjusted p values were estimated by beta approximation method using the parameter “--permute 10000.” Genome-wide significance was determined by correcting the adjusted p values for multiple testing across genes using Benjamini-Hochberg method (FDR < 0.05 were considered significant).

METAL47 was used for meta-analysis of cis-eQTL mapping by combining nominal p values across three cohorts while taking the sample size and direction of effect into account. For each gene, the most significantly associated variant per gene (i.e., the one with the smallest meta p value) was recorded to form the empirical, true meta p value distribution. To assess the significance of meta p values, eQTL mapping within each cohort was repeated using permuted gene expression data. METAL was successively run on the permuted eQTL results across three cohorts and permuted meta p values were obtained. For each gene, the most significantly associated variant per gene (i.e., the one with the smallest meta p value) was recorded to form the empirical, null meta p value distribution. Next, FDR of 0.05 was estimated such that Probability(p value0 < z)/Probability(p value1 < z) = 0.05, where Probability(p value0 < z) is the fraction of p values from the null meta p value distribution that fall below the p value threshold z and Probability(p value1 < z) is the corresponding fraction in the true meta p value distribution (See Figure S11 for QQ-plots of meta cis-eQTL associations, Figure S12 for relative distance of meta cis-eQTLs to their target gene TSS, and Table S4 for significant meta cis-eQTL results).

Identification of Shared and Liver-Specific cis-eQTLs

Among 2,625 lead cis-eQTLs, 2,552 of them were tested in GTEx Analysis Release6p. For each of these 2,552 lead cis-eQTL-gene expression pairs, posterior probability of association in 43 non-liver GTEx tissues were calculated using METASOFT. cis-eQTLs with a posterior probability of 0.9 in at least 38 non-liver tissues were defined as “shared-eQTLs” and cis-eQTLs with a posterior probability of 0.9 in fewer than 5 non-liver tissues were defined as “liver-specific eQTLs.” Overlap between these three sets of cis-eQTLs (total, shared, and liver-specific) and H3K4me3 and H3K27ac peaks were identified using bedtools intersect --u function. Liver H3K4me3 and H3K27ac peaks identified in this study as well as those of ENCODE consortium (links to ENCODE datasets are in Table S5) were included in this analysis. For each cis-eQTL set, 1,000 matching SNP sets (LD of r^2 >= 0.5, MAF of at least 5%, gene density of at least 5%, distance to nearest gene of at least 50%, LD buddies of at least 50% in European 1000G Phase3) were obtained using SNPSnaps. Odds ratios of observed/expected overlaps were plotted in Figures S13 and S8.

Mapping trans-Expression Quantitative Trait Loci (trans-eQTLs)

Associations between each expressed autosomal protein coding genes and variants that are more than 5 Mb apart were considered as trans. trans-eQTLs were mapped using MatrixeQTL while adjusting for the same covariates that were used in cis-eQTL mapping (i.e., sex, ancestry, PEER factors in each cohort as well as genotyping platform in GTEx cohort). trans-eQTL mapping was performed within each cohort using (1) all linkage disequilibrium pruned variants (r^2 > 0.5, plink parameters –indep 50 5 2 across cohorts), (2) variants that were identified as cis-eQTLs in this study (2,625 variants based on meta cis-eQTL results), and (3) variants that are likely to affect transcription factor activity. In approach 1, our goal was to perform a hypothesis-free genome-wide trans-eQTL scan. In approach 2, we hypothesized that cis-eQTLs through regulation of gene expression can alter protein levels of their cis-eQTLs and coding variants in protein levels can affect expression levels of other genes downstream. In approach 3, we hypothesized that cis-eQTLs and coding variants of transcription factors are likely to alter transcription factor activity (either through altering protein level or protein function) and that differences in transcription factor activity can affect expression levels of other genes downstream. For approach 3, we obtained the curated list of 1,988 transcription factors from T. Ravasi et al.52 We included the significant meta cis-eQTLs for this set of genes and protein coding variants of this set of genes (based on gnomAD release-170228) that passed our initial genotyping QC threshold within each cohort (4,997, 4,860, and 5,516 variants for Penn Cohort 1, Penn Cohort 2, and GTEx, respectively).

METAL was used for meta-analysis of trans-eQTL mapping by combining nominal p values across three cohorts while taking the sample size and direction of effect into account for each type of trans-eQTL approach. Similarly, METAL was run on the permuted trans-eQTL results across three cohorts and permuted meta p values were obtained. Permutation-based FDR was calculated as explained in the cis-eQTL mapping section above. See Figure S14 for QQ-plots of meta trans-eQTL associations. There were two statistically significant trans-eQTL findings when genome-wide approach was used. However, both of these results were filtered due to presence of genes near trans-eQTL (within 100 kb) with evidence of RNA-seq read cross-mapping due to sequence similarity (Table S12). There were no significant findings when only cis-eQTLs or only variants likely to affect transcription factor activity were tested as trans-eQTLs.

Mapping cis-Histone Quantitative Trait Loci (cis-hQTLs)

Picard Tools’ MarkDuplicates function was used with the “REMOVE_DUPLICATES = true” parameter set to remove duplicate reads from aligned and q10 filtered ChIP-seq data (initial ChiP-seq data processing and QC steps were explained above in the ChiP-Seq Data Processing section). Genotype data of Penn cohort 1 were further processed to include only single-nucleotide substitutions in hQTL mapping. Allele-specific read counts were obtained with GATK’s ASEReadCounter function. Total feature counts and GC% values of each feature were used to generate sample specific offset values for each feature. To generate PEER Factors, FPM values (equivalent of TPM for ChIP-seq data) were calculated, quantile normalized between individuals, and inverse quantile normalized to a standard normal distribution within each peak. Sex, first three genotype-based PCs, and five PEER factors were included as covariates. RASQUAL was used to map hQTL.
associations. Variants within 10 kb of each end of a histone peak were considered as cis. All autosomal peaks were included in hQTL mapping. This corresponded to a total of 65,649 and 128,822 tested peaks for H3K4me3 and H3K27ac, respectively. For each peak, the most significant p value was selected to form the empirical, true hQTL p value distribution. To assess genome-wide significance, RASQUAL was successively run using the -r/-random-permutation option. For each peak, the most significant p value from the permutation run was selected to form the empirical, null hQTL p value distribution. Next, FDR of 0.05 was estimated such that Probability(p value0 < z)/ Probability(p value1 < z) = 0.05, where Probability(p value0 < z) is the fraction of p values from the null hQTL p value distribution that fall below the p value threshold z and Probability(p value1 < z) is the corresponding fraction in the true hQTL p value distribution. Peaks with significant hQTLs were excluded if they had potential reference mapping bias (φ < 0.25 and φ > 0.75). See Table S7 for significant hQTL results.

Integrative Analyses of Liver cis-hQTLs with Other Functional Datasets
We used ENCODE Regulation “Txn Factor” track and downloaded the wgEncodeRegTifsClusteredWithCellsV3.bed.gz file, which includes transcription factor binding site clusters together with the input cell sources. Using this file, we extracted the binding sites of 61 transcription factors that were obtained in HepG2 cell line. Link to the ENCODE dataset used is included in Table S5. Peaks with cis-hQTLs were overlapped with 61 different transcription factors’ binding sites in HepG2 cells using bedtools intersect -u function. Enrichment of overlap was calculated relative to 1,000 sets of randomly chosen matching numbers of H3K4me3 and H3K27ac autosomal peaks in our data. One-sided Fisher’s exact test was used to assess the significance of the enrichment.

For 151 of the hQTL-peaks, target interacting gene promoters could be identified using the Capture-C data. For each variant that was located within such hQTL-peak, cis-eQTL p value for the interacting gene was pulled to form the distribution of observed cis-eQTL p values. This observed p value distribution was then compared to the expected distribution of p values that was observed when 151 histone peaks with no hQTLs were chosen randomly from the set of autosomal histone peaks that do not have significant hQTLs. In a complementary analysis, the proportion of eQTL-genes among the 210 genes that interact with hQTL-peaks was compared with the proportion that was observed among 1,000 sets of randomly chosen 210 genes that were baited and expressed but that do not interact with hQTL-peaks. Significance was assessed based on the permutation p value.

Identification of Trait-Related Genes and Regulatory Elements in GWAS Loci
GWAS summary statistics of 20 phenotypes including coronary artery disease, HDL cholesterol, LDL cholesterol, total cholesterol, triglycerides, diastolic blood pressure, systolic blood pressure, mean arterial pressure, rheumatoid arthritis, type 2 diabetes, multiple sclerosis, asthma, psoriasis, Parkinson disease, Alzheimer disease, schizophrenia, Crohn disease, ulcerative colitis, inflammatory bowel disease, and age-related macular degeneration were obtained (see Table S9 for links to datasets). Among GWAS variants with p < 10^{-6}, most significant variant was chosen to represent each 2 Mb region and to define each significant GWAS locus, 1 Mb upstream and 1 Mb downstream of the lead variant. Approximate Bayes Factor colocalization analysis of the coloc package was performed between each eQTL-gene whose TSS was within each GWAS locus and the corresponding GWAS phenotype. To assess the significance of colocalization analysis, we used a previously published approach and assessed whether there was sufficient power to test for colocalization (PP3 + PP4 > 0.8), and for the colocalization pairs that pass the power threshold, we defined PP4/(PP3 + PP4) > 0.9 as the significant colocalization threshold. Similarly, significant hQTL-peaks that were located within each GWAS locus were tested for evidence of underlying GWAS signals. When lead hQTLs and lead GWAS variants were in high linkage disequilibrium (r^2 > 0.8), we considered such hQTL-peaks as the likely trait-relevant regulatory elements in GWAS loci. r^2 was calculated in the 1000 Genomes, Phase 3, European population. All significant signals in the MHC region (chr6: 28,510,120-33,480,577) were excluded owing to the complicated LD patterns of this locus.

In loci where we identified a single colocalized gene, we classified whether the candidate genes prioritized in our study have been previously reported as likely trait-relevant genes for the phenotype of interest or not. To do so, we downloaded the NHGRI-EBI GWAS Catalog in May 2018. We retrieved the entries that matched the GWAS phenotype (column “MAPPED_TRAIT”) as well as chromosome nomenclature of the GWAS locus under question (column “REGION”). For each retrieved entry, we checked the hg19 genomic position of the SNP reported (column “SNPs”) and pulled entries that are located within the GWAS locus of interest (within 1 Mb to GWAS lead variant). Among the selected entries, we checked the proportions of studies that reported the colocalized gene as the sole versus among many candidate genes of interest (column “REPORTED.GENE(S)”). Colocalized genes that have not been reported as candidate GWAS genes in NHGRI database were further retrieved in PubMed database using search terms that included the colocalized gene ID in combination with GWAS phenotype. Each retrieved manuscript was then evaluated to determine whether colocalized gene has been previously reported as candidate GWAS gene for the phenotype of interest or not. We note that our literature review may be incomplete as the review was performed by a single author and was limited to manuscripts that were written in English and those that were open.
Results

Annotation of Regulatory Elements and Their Interacting Gene Promoters

H3K4me3 and H3K27ac are epigenetic histone modifications that are enriched in functional non-coding regions of the human genome, including active gene promoters and enhancers.65–67 We performed chromatin immunoprecipitation (ChIP)68 for H3K4me3 and H3K27ac modifications in human liver tissue, sequenced ChIP-ed DNA, and obtained H3K4me3 and H3K27ac ChIP-seq data from 9 and 18 individuals, respectively (Figure 1A and Table S1, see Material and Methods for all cohort and analysis details). Using these data, we annotated 68,600 and 131,293 genomic regions enriched for H3K4me3 and H3K27ac modifications (i.e., ChIP-seq peaks; Figure 2A and Table S2). Similar to previous reports,65–67 we showed that H3K4me3 is highly enriched (p < 2.2 × 10^{-16}) near transcription start sites (TSSs) and H3K27ac is approximately equally present within intronic, distal intergenic, and near-TSS regions (Figure 2B). We also collected genome-wide genotype and RNA-seq data in the liver from two cohorts and, alongside with publicly available GTEx v6p liver data,25 we studied the extent of inter-individual variation in liver gene expression levels in a total sample size of 241 (Figure 1A). Across the three cohorts in our study, we identified 23,271 expressed genes.
Welch two sample t test p values were expressed genes were significantly higher than those between H3K27ac peaks and genes that were not detected as expressed. One-tailed (G) Distribution of distance between interacting bait promoters and histone peaks. genes, respectively.

than 1 Mb apart.

For details), we identified 29,328 significant promoters (Figure 2G). More than 99% of the histone peaks with evidence of looping were within 1 Mb of their interacting promoter (Figure 2E). Furthermore, the interactions between expressed gene promoters and the liver histone peaks were significantly stronger than those observed between histone peaks and genes that were not expressed (p < 2.2 × 10^-16; Figures 2C and 2D). More than 99% of the histone peaks with evidence of looping were within 1 Mb of their interacting promoters (Figure 2G).

Identification of Shared and Liver-Specific cis-eQTLs
We mapped cis-expression quantitative trait loci (cis-eQTL; here defined as associations between the expression level of a gene and a variant within 1 Mb of the gene TSS) within each cohort by linear regression and performed a meta-analysis across three cohorts. We identified 2,625 genes with cis-eQTLs at 5% FDR; we hereafter refer to such genes as eQTL-genes (Figure 1B and Table S4). For each cis-eQTL identified, we estimated the posterior probability that the eQTL effect is present in non-liver GTEx v6p tissues. (Table S5). Overall, cis-eQTLs were significantly more likely to overlap H3K4me3 and H3K27ac histone peaks relative to randomly selected SNPs matched for key properties...
numbers of peaks were drawn from the set of expressed and baited genes that do not interact with hQTL-peaks. eQTL p values of the variants within the background set of histone peaks on their interacting gene(s) are shown in black color. One-sided Wilcoxon rank sum test p value comparing two p value distributions was 3.75 × 10⁻⁷.

(F) Percentage of eQTL-genes among genes that interact with hQTL-peaks is shown in the magenta vertical line. Distribution of expected frequency (MAF), gene density, and distance to nearest gene (one-tailed Fisher’s exact test p value comparing two p value distributions was 3.75 × 10⁻⁷).

Identification of cis-hQTLs in the Human Liver
To identify genetic determinants of H3K4me3 and H3K27ac modifications in the liver, we applied a method that uses both total and allele-specific signals in sequencing data to enable quantitative trait loci (QTL) mapping with relatively small sample sizes.²⁰ We identified cis-QTLs for 51 H3K4me3 and 921 H3K27ac peaks at 5% FDR (Figures 1C and 3C and Table S7). We refer to such variants as histone QTLs (hQTLs) and the peaks that they intersected the hQTL-peaks with transcription factor binding sites (TFBS) data. For each variant within a hQTL-peak, its eQTL-pvalue on the interacting gene(s) is plotted in magenta color. Matching

including linkage disequilibrium (LD), minor allele frequency (MAF), gene density, and distance to nearest gene (Figure 3B and Table S6). Moreover, shared eQTLs overlapped H3K4me3 promoter marks more often than liver-specific eQTLs (Figure 3B and Table S6). Conversely, while shared eQTLs showed similar levels of overlap with H3K27ac enhancer marks across different tissues, liver-specific eQTLs were significantly more likely to overlap H3K27ac marks that we identified in human liver tissue as well as those that were identified in liver-derived HepG2 cell lines, consistent with the previous reports of significantly higher cell-type specificity of enhancers relative to promoters⁹⁹ (Figure 3B and Table S6). Despite implementing multiple approaches, we did not identify any significant trans-eQTLs in the human liver (see Material and Methods for details).
remodeling (JUN and JUND) when compared with randomly selected matching numbers of liver histone peaks from our data (p < 0.05; Figure 3D). Furthermore, using our chromatin capture data, we found that variants within hQTL-peaks were more likely to be significantly associated with the expression of genes with which they are in contact with relative to the variants within histone peaks that do not have hQTLs (p = 3.75 × 10⁻⁹, Figure 3E). Overall, genes that interact with an hQTL-peak were almost twice as likely to have cis-eQTLs relative to randomly selected matching numbers of expressed and baited genes that do not interact with hQTL-peaks (p = 0.001; Figure 3F). These results suggest that genotype-dependent putative functional elements identified here play causal roles in the regulation of gene expression levels and this, at least in part, is mediated via DNA looping interactions.

Putatively Co-regulated Histone Modification States and Gene Expression Levels

Integrating eQTL associations with regulatory element annotations has proven useful for the precise identification of causal regulatory variants and the specific regulatory elements they perturb. Our results highlight the value of analyzing tissue-type-matched gene expression and regulatory element datasets (Figure 3B). These analyses, however, are limited as there are often multiple regulatory elements within each eQTL locus and hence it has remained difficult to systematically link regulatory elements to their respective target genes. To address this, we first identified putatively co-regulated hQTL-peaks and eQTL-genes. Because of the limited sample size of our ChIP-seq data, we identified co-regulated peak-gene pairs based on LD between lead QTL-SNPs (i.e., r² > 0.8 between the lead hQTL and lead eQTL; Figure 4A). We found 116 gene-peak pairs that are likely regulated by the same causal variant (Table S8). These 116 gene-peak pairs corresponded to 104 unique eQTL-genes and 95 unique hQTL-peaks. hQTL-peaks were often not assigned to their nearest gene; in 71% of the co-regulated gene-peak pairs, there was at least one other gene that is closer to the hQTL-peak than the eQTL-gene with which it is co-regulated (Figures 4B and 4C). Figure 4D displays an example of a putatively genetically co-regulated gene-peak pair (r² between lead QTLs = 0.89), supporting the presence of a shared causal effect underlying the activity of an enhancer located in the second intron of SPIRE1 (H3K27ac-84963; chr18: 12,551,731–12,553,678) as well as SPIRE1 gene expression level. The full set of putatively genetically co-regulated gene-peak pairs are included in Table S8.

Identification of Trait-Relevant Genes and Regulatory Elements in GWAS Loci

Next, we asked whether leveraging our hQTL, eQTL, and chromatin capture findings could help fine-map GWAS loci. Throughout this manuscript, we defined “fine-mapping” as evidence of refinement in putatively trait-relevant gene, regulatory element, and variant identification in any individual GWAS locus. We obtained GWAS summary statistics of 20 phenotypes that are commonly studied (based on the number of PubMed IDs in the NHGRI-EBI GWAS Catalog) and that have variable levels of suggested causality manifesting in the liver. These phenotypes included coronary artery disease, HDL cholesterol, LDL cholesterol, total cholesterol, triglycerides, diastolic blood pressure, systolic blood pressure, mean arterial pressure, rheumatoid arthritis, type 2 diabetes, multiple sclerosis, asthma, psoriasis, Parkinson disease, Alzheimer disease, schizophrenia, Crohn disease, ulcerative colitis, inflammatory bowel disease, and age-related macular degeneration. Links to the GWAS summary statistics used are included in Table S9. We used a p value threshold of < 1 × 10⁻⁶, selected a lead variant to represent each 2 Mb region (1 Mb upstream and 1 Mb downstream of the lead variant), and identified 1,614 loci previously associated with these phenotypes. Genetically regulated gene expression levels and histone modification states in GWAS loci can reveal the
Figure 5. Candidate Trait-Relevant Genes and Gene Regulatory Elements in GWAS Loci

(A) Each GWAS locus was defined as the 2 Mb region around the lead GWAS variant. A Bayesian colocalization approach was performed between the GWAS phenotype and each gene with a significant meta cis-eQTL whose TSS resides within the GWAS locus.

(B) An LD threshold of $r^2 > 0.8$ between lead GWAS variants and lead hQTLs was used to identify putatively trait-relevant cis-regulatory elements in GWAS loci.

(legend continued on next page)
mechanisms underlying observed associations between genetic variants and disease phenotypes.\textsuperscript{7–14} We therefore applied a Bayesian colocalization approach\textsuperscript{64} to identify eQTL-genes that likely underlie disease phenotypes (Figure 5A) and used an LD threshold ($r^2 > 0.8$) between lead GWAS variants and lead eQTLs to identify putatively trait-relevant cis-regulatory elements in GWAS loci (Figure 5B). To assess the significance of colocalization analysis,\textsuperscript{64} we used a previously published approach\textsuperscript{16} and first assessed whether there was sufficient power to test for colocalization (PP3+PP4 > 0.8), and for the colocalization pairs that pass the power threshold, we defined the threshold for significance as PP4/(PP3+PP4) > 0.9. In loci where we found at least one eQTL-gene with significant evidence of underlying GWAS associations, we repeated colocalization analysis using all expressed genes within 2 Mb region around each lead GWAS variant. Our rationale was to avoid possibility of excluding genes with moderate eQTL signals, which did not reach genomewide significance threshold of eQTL mapping but displayed significant evidence of eQTL-GWAS colocalization.

We found a total of 125 GWAS-gene and 33 GWAS-peak pairs with evidence of shared genetic causality (Table S10). For 77 GWAS-gene pairs, our dataset contains evidence supporting identification of the trait-relevant gene, as there was only one gene that significantly colocalized with the GWAS phenotype (Figures 5C–5E). We identified several candidate genes and regulatory elements that underlie associations with more than one GWAS phenotype. For instance, 77 GWAS-gene pairs with only one colocalized gene implicated 54 unique genes (Figures 5C–5E). 21 of these genes were previously reported as the likely trait-relevant gene at the locus\textsuperscript{15,56,61,64,72–79} (Figure 5C). For 10 loci, our findings help refine candidate gene identification from among several genes that were suggested to be trait relevant in the literature\textsuperscript{56,72,73,76,79–84} (Figure 5D) and in 21 loci, we discovered candidate trait-relevant genes that have not been previously suggested to underlie the corresponding GWAS phenotype (Figure 5E). We were not able to identify any trait-relevant genes for rheumatoid arthritis, multiple sclerosis, age-related macular degeneration, and Alzheimer disease using the data collected in the human liver.

At the 17q21.32 locus, we identified a genetically regulated enhancer (H3K27ac-10102; chr1: 109,816,977–109,818,871)\textsuperscript{11} as the candidate regulatory element responsible for the GWAS associations with coronary artery disease, HDL cholesterol, LDL cholesterol, and total cholesterol levels (Figures 5F and S15). Our chromatin capture interaction data also revealed an interaction between ChIP-seq peak H3K27ac-10102 and the promoter of the SORT1 (MIM: 602458) gene, supporting the previously reported regulatory role of this enhancer on SORT1 gene expression (Figure S15).\textsuperscript{11}

At the 11q12.2 locus, which has been shown to be a critical component of adaptation to different diets,\textsuperscript{86,87} we identified a genetically regulated enhancer (H3K27ac-33409; chr11: 61,587,373–61,589,527) that likely underlies associations with blood lipid phenotypes, asthma, and Crohn disease (Figure 6B). Interestingly, a previous study has reported a genetically regulated DNA methylation probe that overlaps the enhancer we identified and suggested that DNA methylation differences in this putative enhancer affects FADS1 (MIM: 606148) protein activity in the liver.\textsuperscript{88} While our colocalization analyses identified both FADS1 and FADS3 (MIM: 606150) as candidate trait-relevant genes (Figure 6B), our findings as well as those from Howard et al.\textsuperscript{88} support the identification of the trait-relevant regulatory element (H3K27ac-33409; chr11: 61,587,373–61,589,527) at this locus.

Lastly, while we were not able to identify a target gene at the chromosome 16q12.2 locus, we found a genetically regulated putative enhancer in the first intron of FTO.

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(C) GWAS loci with only one significantly colocalized gene and the gene identified has been reported as the likely trait-relevant gene by overwhelming majority of the literature.
(D) GWAS loci with only one significantly colocalized gene and the gene identified has been reported among several genes that were suggested to be trait relevant in the literature.
(E) GWAS loci with only one significantly colocalized gene and the gene identified has not been previously implicated in the corresponding GWAS phenotype.
(F) GWAS loci with more than one significantly colocalized gene. Genes within the same GWAS locus are shown in alternating shades of yellow and blue. When identified, putatively trait-relevant histone peaks are included next to the colocalized gene names of each locus.
(G) GWAS loci where candidate trait-relevant regulatory elements were identified in the absence of colocalized liver eQTL genes.

Phenotype abbreviations are as follows: CAD, coronary artery disease; HDL, HDL cholesterol; LDL, LDL cholesterol; TC, total cholesterol; TG, triglycerides; DBP, diastolic blood pressure; SBP, systolic blood pressure; MAP, mean arterial pressure; T2D, type 2 diabetes; AST, asthma; PSO, psoriasis; PD, Parkinson disease; SCH, schizophrenia; CD, Crohn disease; UC, ulcerative colitis; IBD, inflammatory bowel disease.
(MIM: 610966) (H3K27ac-70404; chr16: 53,812,377–53,812,817) with evidence of underlying GWAS associations in this locus with type 2 diabetes, HDL, and triglyceride levels (Figure 6D). To our knowledge, there has not been a previous report of a genotype-dependent regulatory element in the human liver that overlaps the...
GWAS interval at this locus, which has received considerable study. 89

Characteristics of Fine-Mapped GWAS Loci

Overall, using the genome-wide data collected in the human liver, we were able to fine-map at least one GWAS locus for 16 out of 20 phenotypes we studied (Figure 5). 30% of the trait-relevant genes had liver-specific cis-eQTLs, 16% had shared cis-eQTLs, and the remaining 54% had cis-eQTLs that were neither classified as liver-specific nor as shared. When we looked at the percentage of GWAS loci with at least one candidate gene (Figure 7A) or regulatory element (Figure 7B) identified in our study, we found that our ability to identify trait-relevant genes and regulatory elements in GWAS loci is correlated with the physiological relevance of the studied phenotype to the human liver. Phenotypes with a known molecular basis in the liver such as blood lipid phenotypes had a larger proportion of GWAS loci with candidate genes or regulatory elements identified in our study (Figures 7A and 7B).

A median of two genes were located closer to the lead GWAS variant than the trait-relevant gene identified in this study. When we compared the candidate trait-relevant genes discovered in our study (Figure 5E) to genes that were previously suggested to be trait-relevant in the literature (Figure 5C), we noted a significant difference in terms of the distance and the number of genes between the lead GWAS variants and the trait-relevant genes identified (Figures 7C and 7D). This discrepancy emphasizes once again that genes reported as trait-relevant in the literature are biased toward those that are closer to the GWAS lead variants and that unbiased genome-wide approaches are required to identify true trait-relevant genes in GWAS loci.

We also note that mutations in four of the complex phenotype-causing genes, LIPA (MIM: 613497), LIPC (MIM: 151670), GPIHBP1 (MIM: 612757), and IRF8 (MIM: 601565), have been implicated in related Mendelian diseases,90–93 and 50% of the trait-relevant genes that have murine models were reported to display similar phenotype in the model organism as well (Mouse Genome Database; Table S11). Lastly, while LIPA, PLTP (MIM: 172425), and SLC39A8 (MIM: 608732) have been previously suggested to affect their associated phenotypes through protein altering mutations,94–96 our findings are in line with those of Wild et al.97 and Hess et al.,98 suggesting that genotype-dependent changes in their gene expression levels also contribute to the complex trait pathogenesis.

Discussion

In 2001, the first published draft of the human genome confirmed that the vast majority of its sequence, approximately 97% of the 3.2 billion bases, has no protein-coding function.99 Following this discovery, the next phase of research focused on understanding and functionally annotating non-coding regions within the human genome. These studies generated reference epigenomic maps for multiple cell lines and tissue types and demonstrated that epigenetic marks on histone proteins are important predictors of gene-regulatory activity.100 Perhaps more interestingly, such gene regulatory regions were subsequently shown to harbor the majority of the complex disease-associated variants,4 making studies of gene regulation an important area of investigation at the interface of basic and disease biology.

In this study, we generated the most comprehensive, in terms of sample size and characterizing inter-individual differences, genome-wide dataset of two epigenetic marks, H3K4me3 and H3K27ac, in the human liver. These studies generated reference epigenomic maps for multiple cell lines and tissue types and demonstrated that epigenetic marks on histone proteins are important predictors of gene-regulatory activity.100 Perhaps more interestingly, such gene regulatory regions were subsequently shown to harbor the majority of the complex disease-associated variants,4 making studies of gene regulation an important area of investigation at the interface of basic and disease biology.
widely functional consequences of natural genetic variation on regulatory element activity and gene expression levels. Furthermore, we showed that a single genetic variant could co-regulate both histone modification states and gene expression levels and this co-regulation is at least partly mediated via DNA looping interactions. We expect that this expansive resource containing functional annotation of non-coding elements and DNA-looping interactions between gene promoters and putative functional gene regulatory elements will greatly facilitate future analyses and stimulate new areas of investigation.

Our results also hold significant relevance for medical genomics. Using genetic colocalization approaches, we fine-mapped a total of 74 GWAS loci associated with at least one complex phenotype. For 21 loci, the gene we prioritized had been previously reported as the likely trait-relevant gene in the majority of the literature. For 10 loci, our findings helped refine identification of the candidate gene from among several genes that were suggested to be trait relevant in the literature. In 21 loci, we discovered candidate trait-relevant genes that have not been previously suggested to underlie the corresponding GWAS phenotype and for a total of 16 loci, we identified candidate trait-relevant gene regulatory elements.

While our efforts constitute the largest GWAS fine-mapping effort performed in the human liver, we were able to identify candidate trait-relevant genes in less than 20% of the GWAS loci even for the most directly liver-related complex phenotypes (i.e., blood lipid levels). This result indicates a need for similar comprehensive studies of the transcriptome and epigenome in a wider range of tissue types and stimulation conditions as well as studies focusing on other complex disease-causing mechanisms. Additionally, we believe performing Capture-C experiments in the human liver tissue as opposed to immortalized HepG2 cell lines could increase our fine-mapping power. We note that while HepG2 cell lines are a widely accepted model system to study liver biology, they display an abnormal hyperdiploid karyotype, which could have effects on chromatin-chromatin interactions. It is also likely that there is inter-individual variation in chromatin-chromatin interactions and when possible Capture-C data should be obtained across several individuals as opposed to a single cell line. Our eQTL and hQTL mapping experiments were performed in whole liver tissue samples and performing similar studies in isolated liver cell types or single cells could also enhance GWAS fine-mapping ability. Furthermore, while statistical colocalization approaches help prioritization of genes in GWAS loci, it is possible that the application of such approaches to eQTL and GWAS summary statistics from larger cohorts may reveal additional colocalization signals in these same GWAS loci. Lastly, while we recognize and value the contribution of genome-wide integrative approaches we and others have undertaken, we note that further in vivo and organism-level validations are necessary to confirm the suggested causality of these findings.

Overall, our findings expand the repertoire of candidate genes and regulatory mechanisms governing complex disease development and contribute to basic understanding of genetic and epigenetic regulation of gene expression in the human liver tissue. Furthermore, by more precisely highlighting genes and regulatory elements with relevance to disease or critical intermediate phenotypes, we believe this study will improve research into the development of therapeutic or preventative measures to mitigate the effects of complex disease. Finally, our approaches to integrate genetic variation and multiple molecular phenotypes across individuals are likely to be applicable to other tissues and traits.

**Accession Numbers**

RNA-seq and ChIP-seq datasets from Penn Cohorts have been deposited in GEO under accession number of GSE128072. Genotype and RNA-seq datasets from GTEx Cohort have been deposited in the dbGaP under ID code of phs000424.v6.p1. HepG2 Capture-C data have been deposited in ArrayExpress with accession number E-MTAB-7144.

**Supplemental Data**

Supplemental Data can be found online at https://doi.org/10.1016/j.ajhg.2019.05.010.

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