Transcription factor and chromatin features predict genes associated with eQTLs

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ABSTRACT

Cell type-specific gene expression in humans involves complex interactions between regulatory factors and DNA at enhancers and promoters. Mapping studies for expression quantitative trait loci (eQTLs), transcription factors (TFs) and chromatin markers have become widely used tools for identifying gene regulatory elements, but prediction of target genes remains a major challenge. Here, we integrate genome-wide data on TF-binding sites, chromatin markers and functional annotations to predict genes associated with human eQTLs. Using the random forest classifier, we found that genomic proximity plus five TF and chromatin features are able to predict >90% of target genes within 1 megabase of eQTLs. Despite being regularly used to map target genes, proximity is not a good indicator of eQTL targets for genes >150 kilobases away, but insulators, TF co-occurrence, open chromatin and functional similarities between TFs and genes are better indicators. Using all six features in the classifier achieved an area under the specificity and sensitivity curve of 0.91, much better compared with at most 0.75 for using any single feature. We hope this study will not only provide validation of eQTL-mapping studies, but also provide insight into the molecular mechanisms explaining how genetic variation can influence gene expression.

INTRODUCTION

Regulation of gene expression in eukaryotes involves multiple transcription factors (TFs) and cofactors acting on DNA at specific genomic loci defined as regulatory elements. Given the complexity of genomes in higher organisms such as humans, the program for this process is even more difficult to decipher (1,2). The first steps toward understanding the regulatory program involve determining the target specificity of TFs that are modulated by interactions with other factors and by the local chromatin structure. Many of these interactions occur in promoter regions that are in proximity to the transcription start sites (TSS) of target genes. Yet, in recent years, experimental evidence has shown that interactions between regulatory elements and target genes can occur over long genomic distances (3–5).

The importance of distal gene regulatory elements for coordinating cell type–specific expression of their target genes has motivated whole-genome surveys of different human cell types. The ENCODE consortium is an international collaborative effort initially set up to build a comprehensive list of all functional elements in the human genome (6). Since then, the consortium has identified cis-regulatory elements (CREs) in a wide variety of cell types (7). The identification of these regulatory elements through mapping protein–DNA interactions has been greatly accelerated by the advent of high-throughput chromatin immunoprecipitation assays like ChIP-seq. The majority of TFs profiled by ChIP-seq have >50% of their binding sites beyond 2.5 kilobases (kb) of a TSS. By combining binding site data on multiple TFs across cell types, the relationship between TFs and genomic features can be revealed (8). However, not all TF-binding sites detected in a particular cell may localize at active regulatory elements. To detect activated elements, DNase-seq and FAIRE-seq (Formaldehyde Assisted Isolation of Regulatory Elements) are used to map genomic regions of open chromatin. The co-localization of TF-binding sites with open chromatin sites gives more confidence to the location of regulatory elements, where most of the cell type–specific sites are typically located away from the TSS (9,10).

Despite the vast array of assays to detect regulatory elements, it remains a challenge to identify their target genes. To date, the most common approach used in computational and experimental studies of regulatory elements has been based on genomic distance. A target

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gene is mapped to a regulatory element if it is the nearest or if it is within a genomic distance threshold. Although it is often the case that genes in proximity to a regulatory element are targets (11–14), there are many important genes regulated by distal elements. One example is the sonic hedgehog gene (Shh), which is regulated by an enhancer located 1 megabase (Mb) upstream (15). Another example is enhancers of c-Myc found in ‘gene deserts’ located a few hundred kilobases away from the nearest gene (16). Other genomic features like the preserved co-localization of genes on chromosomes of different species, or conserved synteny, are also considered, as it reflects co-evolution between regulatory elements and target genes (17,18). However, low conserved synteny is reported between upregulated genes and enhancers detected by ChIP-seq (19). For these reasons, there remains a need for a systematic approach of predicting and validating target genes of regulatory elements.

This area of research is limited by the paucity of in vivo data describing interactions between regulatory elements, such as enhancers, and target genes. Reporter assays, which place the element in question at the promoter of a reporter gene, allow investigation of enhancer activity (20,21), but do not capture the chromatin structure, which allows distal enhancers to interact with target genes. Because of this, methods to capture chromosome conformation were developed to provide evidence of long-range physical interactions (22,23). However, genome-wide data sets of chromosome conformation have a resolution on the order of megabases, too large to screen for interactions between multiple enhancers and target genes. A more indirect approach is to use co-variation between gene expression and enhancer-associated chromatin markers to map enhancers to target genes (24). It has been suggested to use markers of histone modification, such as H3K4me1, to identify active enhancers. But because H3K4me1 occurs ubiquitously along the genome, there are still too many potential enhancers that can be mapped to a particular gene. This means that such approaches lack the specificity needed to detect direct interactions between enhancers and targets. Nevertheless, analysis of histone modification sites has identified unique chromatin signatures for distal regulatory elements (24,25). The frequent positioning of disease-related single nucleotide polymorphisms (SNPs) within regulatory elements defined by chromatin markers suggests that integrative modeling of multiple chromatin features may help decipher the connection between regulatory elements and diseases (24).

Another approach for finding associations between target genes and regulatory elements relies on analyzing SNPs. The decreased cost of genome sequencing has resulted in the identification of genetic variants in different individuals and cell types. Genome-wide association studies that compared genetic variants with the occurrence of disease have implicated numerous enhancer regions (26,27). This information on genotypes can also be analyzed together with gene expression data to find associations between specific genetic variants and gene expression levels as determined through expression quantitative trait loci (eQTL) studies (28–30). Numerous validated regulatory elements have been identified using eQTL data, progressing the field of functional genomics (31). Because many SNPs in linkage disequilibrium could be associated with the expression of target genes, prediction of target genes using eQTL data suffers from the same low specificity as using chromatin markers. Efforts to integrate eQTL data with other genomic features have provided better estimates of the regulatory effect on target genes (32,33). In particular, Gaffney et al. (33) have shown that TF-binding sites and chromatin markers are enriched in regions with causal eQTL SNPs. Nevertheless, there exists no systematic study of how well associations found in eQTL studies agree with results from previously mentioned studies on TF binding and chromatin structure.

Addressing this question, we looked for putative regulatory elements defined by co-localization of TF-binding, chromatin signatures and eQTL signals. Using these elements, we present a method to predict their target genes. The method uses a combination of features as predictors, including genomic proximity, TF binding, gene expression, open chromatin, Gene Ontology (GO) similarities and insulators. We evaluated the performance of the target gene predictions using eQTL data from lymphoblastoid cell lines (LCL), fibroblasts and T-cells (30). Our tests showed substantially higher accuracy for predictions made using a combination of features compared with using any single feature in isolation. Our method is particularly useful for finding additional target genes when eQTL studies are underpowered to do so. In addition, the features we use describe cis-interactions between regulatory elements and target genes. This is especially useful when trying to identify genes that are directly regulated by eQTLs rather than indirectly regulated through trans-interactions.

MATERIALS AND METHODS

TF-binding sites

We obtained ChIP-Seq data on the binding sites of 29 TFs plus “insulator associated DNA-binding protein,” CTCF in the GM12878 cell line from the ENCODE consortium (Additional file 1, Supplementary Table S1). Each data set contains a signal map of ChIP DNA fragments, where the signal height is the number of overlapping fragments at each nucleotide position in the genome (UCSC hg18). ChIP-seq signals mapped to hg19 were converted to hg18 coordinates using the UCSC liftover tool. Enrichment of genomic regions for protein binding was tested against a set of input DNA control (P ≤ 0.01). Peaks indicating regions with sufficient signal above peak-height threshold (false discovery rate < 0.05) were identified using the PeakSeq algorithm (34). After conducting a genome-wide scan for peaks, we examined tracks of putative binding sites for each of the TFs. The position of each binding site was defined by the center of each ChIP-Seq peak. Adjacent binding sites that are within 500 bp of each other were grouped to form non-overlapping regions of TF-binding sites. The boundaries of each region are defined by the left-most and right-most binding sites.
Chromatin marker data
A multivariate hidden Markov model was applied to the combinatorial patterns of nine histone modification markers to distinguish 15 chromatin states (24). The states were learned de novo on the basis of the patterns of chromatin marks and their spatial relationships in the GM12878 genome. Six broad classes of chromatin states were defined as promoter, enhancer, insulator, transcribed, repressed and inactive states. Within each class, active, weak and poised promoters (states 1–3) differ in expression level; strong and weak candidate enhancers (states 4–7) differ in expression of proximal genes; strongly and weakly transcribed regions (states 9–11) also differ in their positional enrichments along transcripts; and repressed, heterochromic and repetitive states (states 13–15) are enriched in H3K9me3. Regions with state annotations vary in length from 500 bp on average for promoter and enhancer states to 10 kb on average for inactive states. Open chromatin sites in GM12878 cell lines were profiled using FAIRE, and the data were downloaded from ENCODE. Enrichment of sequence fragments from FAIRE was identified using a feature density estimator, F-Seq (35). For each enriched region, the maximum F-Seq density signal value has been calculated, and \( P \)-values for peaks were determined by fitting the data to a gamma distribution. A \( P \)-value threshold of 0.01 was considered to be significant.

Gene expression data
The gene expression profiles of 38 distinct populations of human hematopoietic cells were downloaded from the Broad Institute DMap Project (36). Quantile normalization was applied across expression arrays, and the log expression intensities for each gene was mean centered. Probe sets were mapped to a gene’s TSS via transcript identifiers and probe set annotations provided by the Ensembl database (release 54). For cases where there were more than one probe set mapping to a gene’s TSS, we filtered for the probe set with the highest variance in expression across cell samples. In total, 8968 genes were profiled in the data set.

eQTL data
Gene expression profiling and association testing with genetic variants was performed on primary fibroblasts, LCLs (Epstein–Barr virus–immortalized B cells) and T-cells from umbilical cords of 75 individuals of Western European origin (30). Dimas et al. (30) conducted association testing between genotypes and gene expression values using the Spearman rank correlation on all SNPs within a 2-Mb window centered on the TSS of each gene. After filtering for significance at the 0.001 threshold, there are 427, 442 and 430 genes with significant associations between SNPs and gene expression in fibroblasts, LCLs and T-cells, respectively. We extended the list of associated eQTL SNPs based on linkage disequilibrium with SNPs in CEU HapMap panels detected from low-coverage sequencing pilot (Pilot 1) of the 1000 Genomes Project. The SNAP tool was used to find the additional SNPs that are above the correlation coefficient \( r^2 \) threshold of 0.8 (37).

Identifying candidate regulatory elements
Regions with chromatin state annotation were mapped to non-overlapping regions of TF-binding sites if they share any base pair. If two different chromatin state regions overlap with one TF-binding region, the chromatin state with the greater number of overlapping base pairs is mapped. All regions with co-localization of TF-binding sites and having chromatin state annotations were referred to as CREs (Supplementary Figure S1). The boundary of a CRE is defined by the left-most and right-most centers of TF binding peaks in a non-overlapping region. Because the width of a TF’s binding signal peak is estimated to be 200 bp, we assumed an eQTL SNP is likely to affect TF binding if it is within 100 bp from the center of the signal peak. We filtered for CREs with eQTL SNPs within 100 bases from the CREs’ boundaries, and used those to test for target gene prediction. CREs can be linked to multiple target genes if the co-localized eQTL SNPs are associated with different genes.

Genomic distances between regulatory element and target gene
For each gene within 1 Mb from the center of each CRE, we calculated the genomic distance between the gene’s TSS and the nearest associated eQTL SNP in the CRE. The positions of gene TSS are the same as those used in the mapping microarray probe sets to genes.

Modeling gene co-expression
We used a generalized additive model (GAM) to describe the relationship between potential target genes and the expression of TFs that occupy CREs (38). The GAM implementation in the R package ‘mgcv’ provides the option of smoothing spline functions for each predictor term, which gave us the flexibility of incorporating non-linear relationships between TFs and genes. For each gene–CRE pair, we considered a model with one or more additive functions:

\[
E(y_i) = \beta_0 + \sum_{j=1}^{n} s_j(x_{ij}) + \sum_{1 \leq j < k \leq n} s_{jk}(x_{ij}, x_{ik})
\]

where \( E(y_i) \) is the expected log expression of the target gene in cell type \( i \), \( \beta_0 \) is the mean expression set to zero, \( x_{ij} \) is the log expression of TF \( j \) in cell type \( i \), \( n \) is the number of TFs in the CRE and \( s_j \) is a spline function, where the degree of smoothing is chosen by cross-validation in the mgcv package. As opposed to using linear predictors, the estimated non-parametric function can reveal non-linearities in the effect of TF on target gene. In this model, we also allow for second-order interactions where \( s_{jk}(x_{ij}, x_{ik}) \) is a set of unknown partial bidimensional smoothing functions.

We modeled gene co-expression for every gene within 1 Mb of each CRE. For each CRE–gene pair, we inferred the parameters \( \beta_0, s_j \) and \( s_{jk} \) for the aforementioned equation using the expression profiles \( x_{ij} \) of the co-localized TFs \( j \) and \( y_i \) of the gene across samples \( i \in S_T \) from the training set \( S_T \). We then predicted gene expression across the samples \( i \in S_P \) in the test set \( S_P \).
using the TF expression $x_{ij}$ in those samples as predictors. The prediction step gave us a predicted gene expression value $\hat{y}_i$ for each target gene in a sample $i \in S_p$. The prediction accuracy was then measured by calculating the square of the Pearson correlation coefficient (denoted by $r^2$) between the predicted expression $\hat{y}_i$ and the observed expression $y_i$ for all samples in $S_p$. We also calculated the adjusted $R^2$ for the model to estimate the proportion of gene expression variation in the training set explained by TF expression, while taking into account the number of predictors. The closer the $R^2$ is to 1, the better the model fits to the data. Whenever the regression line fits worse than the horizontal mean line, $R^2$ is negative. This can easily happen for non-linear regressions. Five-fold cross-validation was performed to assess how well predictions would generalize to new sample data sets. The $r^2$ and $R^2$ values reported were averaged over the cross-validations. We used the $R^2$ value as a feature in the prediction of eQTL targets.

**TF co-occurrence**

When we examined the non-overlapping regions of TF-binding sites, we counted the different pairs of TFs co-localized to the same region. The co-occurrence between pairs of TFs $i$ and $j$ is measured by the log-odds score of the observed number of regions containing the binding site pairs $b_{ij}$ over the expected number $\hat{b}_{ij}$.

$$c_{ij} := \log(b_{ij}/\hat{b}_{ij})$$

To estimate the expected number of pairs, we repeatedly permuted ($n_{perm} = 1000$) the TF labels for each TF-binding site and then recounted the pairs of TFs. The average number of pairs from all recounts after permutations is the expected number of pairs.

$$\hat{b}_{ij} = \frac{1}{n_{perm}} \sum_{p=1}^{n_{perm}} b_{ijp}$$

We use the co-occurrence score for different pairs of TFs to measure the co-occurrence between TFs binding in a CRE and TFs binding to the promoter region ($\pm 1$ kb from TSS) of a gene. Calculating the overall co-occurrence score between the two sets of TFs, $Y$ and $Z$, is analogous to maximum weighted matching for bipartite graphs. If we define an indicator variable $x_{y,z}$ for each edge between a TF $y \in Y$ and a TF $z \in Z$, the weights are the co-occurrence score for TF pairs $c_{y,z}$.

$$C_{Y,Z} = \max \left[ \sum_{y \in Y, z \in Z} x_{y,z} c_{y,z} \right]$$

This sum is used as a feature for classifying whether the gene is a target or non-target. Genes without any TF-binding sites in the promoter region is assigned the minimum overall score recorded for genes with TF-binding sites.

**Regions of open chromatin**

Identified FAIRE peaks were mapped to a gene’s promoter if the center of the FAIRE peak is within 1 kb of the gene’s TSS. The FAIRE signal value was used as a feature for eQTL target prediction. A higher signal indicated that chromatin is open in the promoter region and the gene is more likely to be a target. Genes without a detected FAIRE peak at the promoter region were assigned a signal value of 0.

**GO similarity**

The information content (IC) of a GO term $t$ is defined by $IC(t) = -\log p_t$, where $p_t$ is the probability of GO term $t$ occurring. The GO database provides an association table mapping genes to GO terms. We compute $p_t$ as the number of genes annotated by GO term $t$ divided by the total number of annotated genes. The pairwise similarity between GO terms $t$ and $t'$ was calculated as the IC of their most informative common ancestor from the set of all common ancestors $Pa(t,t')$.

$$sim(t,t') = \max_{t \in Pa(t,t')} [IC(t)]$$

We then searched the GO database to find GO terms $t_1, \ldots, t_n$ mapped to the gene $g$ within 1 Mb of a CRE, and GO terms $t'_{1}, \ldots, t'_{m}$ mapped to the gene $g'$ encoding a TF that binds to the CRE. As previously proposed (39), we assigned each GO term $t_i$ occurring in gene $g$ to its best matching partner $t'_i$ in gene $g'$ to calculate the GO similarity measure $sim(g,g')$.

$$sim(g,g') = \text{avg} \left[ \sum_{i=1}^{m} \max_{t \in Pa(t,t'_i)} [IC(t)] \right]$$

When the TF and the target gene have an unequal number of GO terms, multiple terms in gene $g$ can be assigned to one term from TF $g'$, and the result may be different for $sim(g',g)$. Therefore, we take $\max [sim(g',g), sim(g,g')]$ as the symmetric version of GO similarity. For CREs with multiple TFs, the average similarity between the gene and TFs was used for target prediction.

**Insulators as enhancer-blocking elements**

We used CTCF-binding sites as markers for the position of insulators that prevent interaction between a CRE and a gene. For each gene, we examined the genomic region in between the TSS and the nearest TF-binding site of a CRE. The CTCF binding peaks were called from the GM12878 ChIP-Seq data provided by ENCODE, and we averaged the signal values across all peaks within the region. If no CTCF binding peaks were found, the signal value was set to zero. The average signal value was used as a feature for target gene prediction.

**Prediction of target genes**

For each gene whose TSS is 1 Mb upstream or downstream from either side of a CRE, we predicted whether that gene is a target of an eQTL SNP within the CRE. Some or all the features used for prediction were combined by a random forest classifier to decide whether a single gene is a target of an eQTL SNP (Figure 1). We used the implementation of random forests in the R package ‘randomForest’ (version 4.6-6).
and trained a random forest of 500 randomly generated decision trees. At each node of a decision tree, the classifier splits the data using a randomly chosen subset of \( m/3 \) features, where \( m \) is the total number of features given to the classifier. The classifier uses the Gini index to determine the best split at each node. The classifier learns to classify each gene as either ‘target’ or ‘non-target’ and outputs the ratio of trees voting ‘target’ for each gene. The same cutoff ratio \( \Theta \) was applied to all genes when making final predictions. Implementation of random forest classifier for eQTL target prediction can be found at http://sysbio.mrc-bsu.cam.ac.uk/eqtlPredictor.

**Evaluating prediction performance**

To evaluate the classification of genes using a specified cutoff \( \Theta \), we calculated three performance statistics:

\[
\text{sensitivity} = \frac{\text{sensitivity}}{\text{sensitivity}}, \quad \text{specificity} = \frac{\text{specificity}}{\text{specificity}}, \quad \text{precision} = \frac{\text{precision}}{\text{precision}}
\]

Sensitivity, also known as recall, indicates the proportion of target genes that are predicted correctly, and precision denotes the probability that a prediction for a target gene is correct. Because both sensitivity and precision only evaluate classification of target genes, we also assess specificity, which measures the proportion of non-targets that are predicted correctly. To evaluate the overall performance of each classifier for various \( \Theta \) values, we used the area under the receiver operating characteristic (ROC) curve, AUC, as the probability that a classifier will assign a higher target probability score to a randomly chosen target gene than to a randomly chosen non-target gene. Alternatively, the \( F \) measure combines precision and recall by representing the harmonic mean of the two measures:

\[
F = \frac{2 \cdot (\text{precision})}{(\text{precision}) + (\text{recall})}
\]

To control for overfitting and test the classifier on new data, we partitioned the CREs into 10 subsets. Of the 10 subsets, 9 were used for training, and the remaining was used for testing. The cross-validation process was repeated 10 times, where each of the 10 subsets was used only once for testing. The values presented for sensitivity, specificity, precision and AUC are the means over the 10-fold cross-validations. Only for the case of predicting eQTL targets in fibroblasts and T-cells did we not cross-validate, as we trained only on LCL data and tested on other cell types.

**RESULTS**

**TF and histone modifications co-localize with eQTLs**

Analysis of binding sites for 29 different TFs (Additional file 1, Supplementary Table S1) from the GM12878 (lymphoblastoid) cell line revealed 221,926 non-overlapping regions with one or more distinct TF-binding sites. Of those, 221,653 regions are annotated to have distinct chromatin states as defined by histone modifications. These regions with both histone modification markers and TFs positioned near the same site, which we call candidate CREs, are quite common throughout the genome (10,25,40). However, it is unclear as to what extent eQTLs are co-localized with these regions. We examined the locations of 551 eQTL SNPs significantly associated with the expression of genes in LCL cells (30). Figure 2A shows the densities of TF-binding sites, histone modifications and open chromatin signals relative...
to genomic distance from the eQTL SNPs. Regions closer to eQTLs seem to have a higher occurrence of TF and chromatin marks. Since the eQTL data set was published, additional genotyping data became available through the 1000 Genomes Project (41). Therefore, we expanded the list of eQTL SNPs to 10,719 by adding proxy SNPs based on linkage disequilibrium. Analysis of CRE locations in relation to eQTL SNPs showed 1303 CREs positioned within 1000 bases of eQTL SNPs, 971 CREs within 500 bases and 369 CREs within 100 bases. Similar across all three cell types is the distribution of eQTL SNPs around CREs (Figure 2B).

Gene co-expression is a weak predictor of distal genes

Reconstruction of gene regulatory networks often made the assumption that the regulatory potential of a TF on a gene decreases the further away it is from the gene (42,43). Network edges were inferred using the expression of TFs to find associations with the expression of target genes, which we refer to as co-expression. Therefore, we examined the expression of 29 TFs binding to CREs to see whether this assumption holds and to see whether there is potential in predicting target genes based on co-expression. The expression profiles of TFs across 38 distinct hematopoietic cell types were compared with the expression profiles of potential target genes near CREs. Regression analysis was used to fit TF expression to target gene expression, and then the expressions of target genes were predicted for different conditions. We applied GAMs to the gene expression data, as it is well suited for describing non-linear relationships between the expression profiles of TFs and target genes (44).

We examined the model fit for each CRE–target gene pair to see how the proportion of expression explained by TFs varies according to the distance between the CRE and the gene’s TSS. In Figure 3, we show that the average proportion of gene expression explained by our regression models, as measured by the $R^2$ statistic, decreases as we considered CREs positioned further from the TSS. However, we could not detect any significant difference in model fit for CREs within 150 kb of the TSS compared with more distal CREs. For CREs containing eQTL SNPs, the $R^2$ statistic also does not differ significantly depending on distance between the CRE and the eQTL’s target gene. Another way to assess co-expression is to assess how well the expression of TFs predicts the expression of target genes. The square of the Pearson correlation between observed and predicted expression also does not seem to be strongly affected by genomic distance. However, a look at expression prediction at a higher resolution (Supplementary Figure S3) reveals that models generated from CREs, which lie within 1 kb of the TSS, explain significantly more expression variation ($P < 2.2 \times 10^{-16}$) than more distal CREs that are beyond 1 kb. The average $R^2$ for models using CREs that lie within the 1 kb promoter region of target genes is 0.221. Beyond 1 kb, the expression prediction models that are generated from the distal CREs have an average $R^2$ of 0.179. Another interesting feature is that there is also a significant drop in prediction accuracy ($P < 2.2 \times 10^{-16}$) beyond 1 kb even if the CRE is the closest CRE to a target gene (Supplementary Figure S3). Overall, the accuracy of the expression prediction models does not improve even if we choose only the closest CREs. This suggests that sometimes more distal CREs may control gene expression over regulatory elements that are positioned closer to the gene. Consideration of the chromatin structure might clarify the relationship of CREs and regulated genes further.

Other weak predictors may help distinguish target genes

Because we believed that gene co-expression is a weak predictor of target genes, we explored other features of genes, which may predict targets better. For each of the 369 CREs that contain eQTL SNPs, we examined all genes within 1 Mb. A gene is considered a target of a CRE if it is significantly associated ($P < 0.001$) with the
eQTL SNP located within the CRE. All other genes were classified as non-targets. We identified 20 CREs that are associated with two different target genes, while the rest are each associated with a single target gene. A total of 3638 genes were classified as non-targets, but they are not evenly distributed across CREs, as some CREs are located in gene deserts (Supplementary Figure S4).

CREs, such as enhancers, can interact with gene promoters by bringing activator proteins to the promoter (3,5); therefore, the likelihood of co-occurrence between TFs bound to the enhancer and TFs bound to the promoter may help determine regulatory potential. We used the ChIP-seq data from the GM12878 cell line to calculate the co-occurrence score for TFs bound at the promoters of each gene, and hypothesized that this feature may help identify genes targeted for regulation. Another feature of gene promoters that may help determine regulatory activity is the presence of open chromatin (9,45). Signal from the FAIRE assay on GM12878 is used to measure open chromatin at each gene’s promoter region. It has been shown that functional similarities between p300 factors and target genes are greater for upregulated genes (19). Therefore, for each of the TFs bound to a CRE, we measured the amount of overlap their GO annotations have with those of target and non-target genes. It has also been shown that CTCF insulator sites block enhancers from interacting with gene promoters (46,47); therefore, we tested to see whether there are more CTCF markers located between CREs and non-target genes than between CREs and targets.

In Figure 4, we compare target and non-target genes in terms of these four features in addition to genomic distance and gene co-expression. Besides genomic distance, we do not see an obvious difference between target and non-target genes based on the features. Nevertheless, we do detect a significantly higher average co-occurrence score and open chromatin signal for target genes ($t$-test $P < 1 \times 10^{-2}$). There is also a significantly

![Figure 3](https://academic.oup.com/nar/article-abstract/41/3/1450/2903273/1456)
lower signal for CTCF markers between target genes and CREs ($t$-test $P < 1 \times 10^{-6}$). It is unlikely that each feature by itself would accurately predict target genes, but the slight difference between the two classes of genes can be exploited if we can combine the features to increase their predictive power.

Integration of TF and chromatin features accurately predict target genes

Recent advances in ‘ensemble learning’, methods that generate many classifiers and aggregate their results, have resulted in better predictive models. One of these new approaches proposed by Leo Breiman is called random forests (48). The random forests algorithm generates a collection of decision trees, where the node in each tree is split using the best split among a subset of predictors randomly chosen at that node. Each fully grown tree will have a classification for every gene as either target or non-target. The probability of a gene being a target is the ratio of trees that classified the gene as a target. The final prediction of a gene’s class depends on the cutoff we choose for this probability, which we refer to as the target probability cutoff $\Theta$. The ROC curves allow us to visualize the different sensitivity and specificity measurements achieved at various cutoffs.

We evaluated prediction of target genes using random forests for 369 CREs that contain eQTL SNPs. The method classified each gene within 1 Mb of a CRE as a target or non-target based on the genomic distance plus five TF and chromatin features that describe the relationship between the gene and the CRE. We compared the predicted target genes with the list of genes associated with eQTL SNPs that are within the CREs. The specificity and sensitivity of the predictions are shown in Figure 5. For $\Theta = 0.5$, the classifier achieves 70% sensitivity, 97% specificity and 69% precision for its predictions. Decreasing $\Theta$ will allow the classifier to achieve >90% sensitivity while still maintaining 90% specificity. Another measure of prediction performance, which can be used to compare classifiers, is the area under the ROC curve (AUC). The classifier that used all six of the features achieved an AUC of 0.96, while the classifier that used only the genomic distance feature achieved an AUC of
This may seem like a small decrease in performance for the classifier using only genomic distance, but at $\Theta = 0.5$, it achieves only 52% sensitivity and 43% precision. The specificity remains high at 93%, as there are many more genes that are non-targets than genes that are targets.

### Prediction of eQTL targets in other cell types

To test whether the target gene classifier might be applicable in other cell types, we analyzed eQTL data from T-cells and fibroblasts (30). In total, 9053 eQTL SNPs were identified in the fibroblast cells, and 307 of the CREs identified using ChIP-Seq data from the GM12878 cell were found to co-localize within 100 bases of the eQTL SNPs. Similarly, 8464 eQTL SNPs were identified in T-cells, and 273 CREs were found to co-localize with eQTL SNPs. We trained the classifiers using the LCL data set of 369 CREs and eQTL SNPs, and then predicted target genes of the 307 eQTL SNPs in fibroblasts and 273 eQTL SNPs in T-cells. Forty-three CREs are common between fibroblasts and LCLs, 33 CREs are common between T-cells and LCLs and 14 CREs are common to all three cell types. The classifiers performed less accurately for T-cells (AUC = 0.83) and fibroblasts (AUC = 0.90) compared with the prediction of LCL eQTL targets (Figure 6). This was expected given that the TF and chromatin features were identified using only data from GM12878, a LCL. Despite this, the prediction of fibroblast and T-cell eQTL targets using only genomic distance still had a lower AUC (0.83 and 0.80, respectively) than prediction using all six features. The decrease in prediction accuracy for target genes in fibroblasts and T-cells may be due to the LCL eQTL data used to train the classifier. As shown in Supplementary Figure S10, when we trained the classifier on cell type–specific eQTL data, prediction accuracy improved for T-cells (AUC = 0.95) and fibroblasts (AUC = 0.91).

We further compared the associations between CREs and target genes in T-cells with chromatin interactions detected by the ChIA-PET assay. CREs that overlap with H3K4me2 regions were identified in CD4+ T-cells. Chromatin interactions detected between these regions and gene promoters were compared with the location of target genes predicted for the CREs (49). Of the 33 CREs that were profiled by ChIA-PET, chromatin interactions are found between 10 CREs and their predicted target genes (Supplementary Table S2). This evidence suggests that our method predicts eQTL–target relationships that are *cis*-regulatory and may be mediated by chromatin structure.

### Accuracy of target prediction is robust for distal genes

Combining genomic distance with five TF and chromatin features helped our random forest classifiers achieve a moderate increase in performance compared with using genomic distance alone. As shown in Figure 4A and Supplementary Figure S2, the majority of target genes are located in proximity to CREs, whereas non-targets are more uniformly distributed across the genome. This characteristic of target genes allows the classifier to achieve high accuracy using genomic distance alone. The real challenge as mentioned earlier is to accurately predict the few target genes that are distal from CREs and eQTLs. The performance of our random forest classifier using all six features remains high (AUC > 0.91), even for classifying genes located at least 150 kb away from a CRE (Figure 7). In contrast, classification of these distal genes using only genomic distance or any other feature alone achieves an AUC of no greater than 0.75. As shown in Figure 7, the performance of genomic distance as a

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**Figure 6.** Comparison of prediction performance for targets of eQTLs in different cell types. For prediction of fibroblast and T-cell targets, we also trained the classifiers on LCL eQTLs and used features generated from GM12878 ChIP-seq and FAIRE data. Because LCL data were used to train the model, it is not surprising to see that the prediction of targets for LCLs achieved a better ROC curve. The performance of predictions using all six features (solid lines) is compared with the performance of predictions using only genomic distance (dotted line).

**Figure 7.** Performance of random forests at classifying genes of a minimum distance away from the CRE. The performance of using all six features (ensemble) is compared with classifiers using only single features.
predictor also drops more than any other predictor for distal genes. At a distance of >150 kb, the presence of insulator markers between the gene and CRE seems to accurately distinguish target genes from non-targets. The feature is able to predict 58% of distal target genes while maintaining >80% specificity (Figure 8). With additional features, we are able to further increase sensitivity without sacrificing much specificity. However, adding the gene co-expression feature actually decreased the performance of a classifier already using four features. This supports the argument that gene co-expression is a weak predictor of target genes.

Chromatin state of regulatory elements affects prediction of eQTL targets

At the beginning of our analysis, we noted that CREs may have different functionalities as independently classified by the chromatin signatures at those regions (24). The CREs can therefore be divided into groups based on their chromatin state annotation. Regions of the GM12878 genome classified as strong enhancer or active promoter elements had chromatin markers associated with high levels of gene transcription. The promoter elements were also confirmed by the enrichment of RNAPII-binding sites, and luciferase reporter assays were used to validate the functionality of predicted enhancers. We examined the performance of target gene prediction for each chromatin state group using $\Theta = 0.5$. As expected, target prediction for CREs annotated as strong enhancers or active promoters had high sensitivity, specificity and precision (Figure 9). Also as expected, the classifier failed to recall or precisely predict many of the target genes of weak promoters, weak enhancers and insulators. The performance of classifiers seems to reflect the chromatin state predictions of likely and unlikely regulatory elements. Yet, CREs annotated as inactive states of heterochromatin have relatively high target prediction accuracy. The heterochromatin regions, which are characterized by low chromatin marker signal, have been previously found to be associated with a subset of transcribed genes (50). It would be interesting to further investigate the regulatory effect that TF binding has on regions thought to be inactive.

Prediction of long-range intra-chromosomal interactions

For longer-range interactions between eQTLs and target genes, the structure of chromosomes may play a role. Regulatory elements may directly interact with distal target genes if the chromosome conformation is such that distant genomic regions are in contact. To identify intra-chromosomal interactions between CREs and predicted target genes in LCLs, we analyzed a HiC data set in the GM06690 cell line (23). In the HiC assay,
intra-chromosomal interactions between two loci are estimated by the number of ligation products containing the two loci. A significantly higher \( (P < 2.2 \times 10^{-16}) \) number of ligation products have loci containing CREs and target genes compared with non-targets (Supplementary Figure S7). Target genes that were predicted by the classifier, but not associated with a eQTL SNP, also had significantly higher \( (P = 7.23 \times 10^{-7}) \) signal for intra-chromosomal interaction. We further examined how the features used in the random forest classifier relate to the frequency of intra-chromosomal interactions. For each CRE, we examined all genes in the same chromosome and compared their features with the interaction frequency between loci containing the CRE and the genes (Supplementary Figure S8). A total of 143,498 CRE–gene interactions were examined. There seems to be a higher interaction frequency when there is higher gene co-expression, TF co-occurrence, open chromatin and insulators between CRE and gene. The relationships are weak, as genomic regions only cover a small portion of each chromosome. Also because each ligation product covers approximately 100 kb, the assay does not have good resolution to detect interactions for specific genes. Despite this, we predicted intra-chromosomal interactions using the six CRE–gene features and the random forest classifier trained on HiC data. The predicted interaction frequencies across 10 cross-validations for all CRE-gene pairs achieved an average \( r^2 \) of 0.78 when compared with observed interaction frequencies (Supplementary Figure S9). This shows that the features can help predict not only statistical associations between genes and eQTLs but also direct interactions between genomic loci.

**DISCUSSION**

The influx in data on gene regulators has enabled researchers to identify regulatory elements by examining histone modification markers and TF-binding hot spots (25,40). Previous studies have also shown that detection of co-localization sites between chromatin features and genetic variants in humans is useful for identifying regulatory elements (24,51). They postulated that the SNPs that influence gene expression may also affect the local chromatin structure. For instance, Degner et al. (51) defined dsQTLs as loci with SNPs significantly correlated with DNase I sensitivity sites. Fifty-five percent of the detected dsQTLs are also eQTLs in LCLs, suggesting that genetic variation may influence both chromatin accessibility and phenotypic variation. The enrichment of TF-binding sites in those dsQTLs suggested possible mechanisms for the elements to act as repressors or enhancers (51), whereas the presence of CTCF insulator elements between the dsQTL and the gene’s TSS was observed to reduce the probability that a dsQTL is an eQTL. This is consistent with observations that CTCF insulators block interactions between regulatory elements and genes (46,47). The analysis of dsQTL and eQTL interactions with genes was restricted to only the closest genes, and knowledge about long-range interactions is still limited.

Because the majority of identified interactions with eQTLs are with nearby genes, most studies ignore long-range interactions. Gene regulatory networks based on co-expression could be used to identify relationships between the TFs bound to eQTL regions and target genes, but inference of the networks is difficult for combinatorial regulation involving many TFs (52). Rdelsperger et al. (19) suggested using evidence of protein–protein interactions between enhancer TFs and TFs binding to the gene’s promoter to identify targets. We considered including protein–protein interactions as a feature, but 11 of the 29 TFs that were profiled did not have any experimental evidence of physical interactions according to the STRING database (53). Instead, we used functional similarity and co-occurrence of TFs as proxies for physical interaction. Detection of shared functional similarity through GO terms is able to recover as much as 88% of known protein–protein interactions across multiple species and identify novel interactions in humans (54,55). Although co-occurrence of TFs has been shown to be a good predictor of protein–protein interactions in yeast (56), until now, there was not enough data on TF-binding sites in humans to exploit this feature.

We are able to use the random forest classifier to integrate the different TF and chromatin features without introducing a complex parametric model. Using multiple features not only allows us to predict 70% of the genes associated with genetic variants, but also we have the possibility of identifying novel gene regulatory interactions. This is especially useful when most eQTL data in human LCLs are underpowered for identifying the genetic pathways involved in complex traits (57). For the purposes of this study, we only examined CREs co-localized with eQTLs, but there are thousands of other CREs without eQTLs that can be predicted using our classifier. It is also important to note that the CRE–target gene associations provided by the eQTL data are based on correlations and may describe some indirect regulatory interactions (58). The random forest classifier, which is trained on eQTL data, may not be predicting only direct targets of CREs, but it also does help identify TF and chromatin signals that link non-coding SNPs to nearby genes. Without better experimental evidence of direct regulatory relationships, we are limited when validating our predictions. Our hope is that new experimental approaches, such as chromatin interaction analysis by paired-end tag sequencing (ChIA-PET), can one day be conducted in a high-throughput manner to validate interactions between promoters and enhancers (49). Our results also show that the quality of these features for inference is limited by cell specificity. This represents a limitation in our method, where features detected in the GM12878 cell type were not as useful for predicting target genes of eQTLs in fibroblasts and T-cells. The dynamic nature of TF occupancy results in dramatically different TF-binding site profiles detected across different cell types (7,59), and in turn causes differential gene expression.

For each gene, the random forest classifier outputs the ratio of trees in the forest voting for the target
classification. This ratio could represent a prioritized list of candidate CRE–target gene interactions to be validated experimentally. Alternatively, we could predict a target by selecting the gene within 1 Mb of a CRE that has the highest ratio. This would make target gene predictions more precise, but we would also have to make the assumption that each element can only regulate one target gene. Making such a strong assumption would hinder the possibility of discovering novel targets. To gain further insight into gene regulatory mechanisms, we could examine the measures for variable importance returned by the classifier (48). However, correlations between TF-binding sites and chromatin markers mean that multicollinearity exists in models combining multiple features (43). This does not reduce the predictive power of the random forest classifier, but makes it difficult to evaluate the contribution of individual features toward the prediction of target genes. To account for correlations between predictor variable when assessing their relative importance, we can permute the values of each feature in the random forests while conditioning on the other features (60). When we did this, we found that the genomic distance feature has the highest conditional importance score, which is consistent with empirical results showing that it is the best predictor of gene-eQTL associations. The ranking of features based on importance scores, shown in Supplementary Figure S5, further suggests that the insulator, TF co-occurrence and GO similarity have some impact on the prediction of target genes, especially for those that are distal (Figure 7). Finding a reduced model based on these features may allow us to identify specific factors influencing long-range regulatory interactions. This will be especially important when more experimental data sets become publicly available, increasing the number of features to consider for integrative analysis.

In summary, we have examined TF and chromatin features that co-localize along the genome. Specifically, we proposed using genomic distance, gene co-expression, open chromatin, TF co-occurrence, GO similarity and insulator marks as possible features for predicting genes associated with eQTLs. Although all the features are weak predictors for distal genes, we have shown that an ensemble of these features can significantly improve eQTL target prediction. It is, however, crucial for these features to be obtained from the same cell population, owing to the cell type specificity of these features. Through this approach, we can propose mechanistic explanations for how genetic variants influence target genes through chromatin state and TF binding dynamics.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online: Supplementary Tables 1 and 2, Supplementary Figures 1–10 and Supplementary Dataset 1.

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