DECODE: a Deep-learning framework for Condensing enhancers and refining boundaries with large-scale functional assays

Zhanlin Chen1,†, Jing Zhang2,*,†, Jason Liu3, Yi Dai2, Donghoon Lee4, Martin Renqiang Min5, Min Xu6 and Mark Gerstein1,3,7,*

1Department of Statistics & Data Science, Yale University, New Haven, CT 06520, USA, 2Department of Computer Science, University of California, Irvine, CA 92617, USA, 3Department of Molecular Biophysics and Biochemistry, Yale University, New Haven, CT 06520, USA, 4Genetics and Genomic Sciences, The Icahn School of Medicine at Mount Sinai, New York, NY 10029-6574, USA, 5NEC Laboratories America, Princeton, NJ 08540, USA, 6Computational Biology Department, School of Computer Science, Carnegie Mellon University, Pittsburgh, PA 15213, USA and 7Department of Computer Science, Yale University, New Haven, CT 06520, USA

*corresponding author: pi@gersteinlab.org.
†The authors wish it to be known that, in their opinion, the first two authors should be regarded as Joint First Authors.

Abstract

Motivation: Mapping distal regulatory elements, such as enhancers, is a cornerstone for elucidating how genetic variations may influence diseases. Previous enhancer-prediction methods have used either unsupervised approaches or supervised methods with limited training data. Moreover, past approaches have implemented enhancer discovery as a binary classification problem without accurate boundary detection, producing low-resolution annotations with superfluous regions and reducing the statistical power for downstream analyses (e.g. causal variant mapping and functional validations). Here, we addressed these challenges via a two-step model called DECODE for Condensing enhancers and refining boundaries with large-scale functional assays (DECODE). First, we employed direct enhancer-activity readouts from novel functional characterization assays, such as STARR-seq, to train a deep neural network for accurate cell-type-specific enhancer prediction. Second, to improve the annotation resolution, we implemented a weakly supervised object detection framework for enhancer localization with precise boundary detection (to a 10 bp resolution) using Gradient-weighted Class Activation Mapping.

Results: Our DECODE binary classifier outperformed a state-of-the-art enhancer prediction method by 24% in transgenic mouse validation. Furthermore, the object detection framework can condense enhancer annotations to only 13% of their original size, and these compact annotations have significantly higher conservation scores and genome-wide association study variant enrichments than the original predictions. Overall, DECODE is an effective tool for enhancer classification and precise localization.

Availability and implementation: DECODE source code and pre-processing scripts are available at decode. gersteinlab.org.

Contact: pi@gersteinlab.org

Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Transcriptional regulation in eukaryotes is the most common and fundamental form of gene regulation for maintaining cell identity during differentiation, determining how cells or organisms respond to intra- and extra-cellular signals, and coordinating various cellular activities (Cramer, 2019; Sperling, 2007). It undergoes precise spatial and temporal regulation via complex interactions of numerous cis-regulatory elements, such as enhancers and promoters, transcription factors (TFs) and chromatin remodelers (Abeel et al., 2009; Dao and Spicuglia, 2018; Jothi et al., 2009; Klemm et al., 2019; Lewis et al., 2019). Hence, enhancer discovery is a cornerstone for understanding transcription control and gene regulation.

The computational methods traditionally utilized for enhancer discovery mainly fall into two categories. First, some methods use the combinatory patterns of various epigenetic features within a
genomic region (e.g., 200 bp bins) to infer the existence of enhancers with unsupervised approaches (Ernst and Kellis, 2012; Hoffman et al., 2012; Moore et al., 2020). These methods use unlabeled datasets to characterize chromatin states and require human interpretation of the discovered states. They are highly transferable for predicting enhancers in a new cell type because the epigenetic patterns are preserved across cell types (Boyle et al., 2014). However, the simple combinatorial, and usually binary, epigenetic patterns may not fully capture the complexity of transcriptional regulation. Second, some methods employ supervised approaches to identify target regulatory elements using hypothetical enhancer loci or a limited number of validated enhancers, which are underpowered for training a reliable model for accurate prediction (Alipanahi et al., 2015; Chen et al., 2018; Li et al., 2018; Lu et al., 2015; Min et al., 2017; Tang et al., 2020). Moreover, we recently developed a linear predictive model based on shape-matching filters from multiple epigenetic features trained from genome-scale STARR-seq experiments on Drosophila (Sethi et al., 2020). However, almost all existing methods can only make binary predictions within a given region (>200 bp) and do not have a high enough resolution for more precise enhancer localization and boundary detection. Previous studies have shown that enhancers can range from 50 to 1500 bp long (Dao et al., 2017; Li and Wunderlich, 2017). Hence, current enhancer annotations usually contain both active enhancer regions and superfluous non-functional regions, introducing noise and reducing statistical power for downstream analyses such as casual variant mapping and functional validation.

Distinct from previous binary classification efforts, we reformulate the enhancer discovery problem into a weakly supervised object detection problem originated from computer vision by answering two questions: (i) Is there an enhancer within a given genomic region? (ii) If yes, exactly where is it? To accomplish these tasks, we take advantage of recent advances in functional characterization assays and utilize direct human enhancer activity readouts from STARR-seq experiments as fuzzy ground-truth labels (Mueller et al., 2015). First, we propose a deep convolutional neural network (CNN) based on a simple but validated hypothesis—the magnitude, shape patterns and cross-coordinate of epigenetic features are important aspects in characterizing the identity of an enhancer (Schmidhuber, 2015). Concretely, we hypothesize that the interaction of open chromatin and histone marks provides a platform for TF binding, which allows epigenetic features to be predictive of enhancers (Sprintz and Furlong, 2012). Second, we add a weakly supervised object detection module to precisely localize the target enhancer in the input genomic region. Specifically, we use the visual activation maps of width and shape-based features from either chromatin accessibility or genomic-wide association study (GWAS) variant enrichment via stratified linkage disequilibrium score regression (LDSC).

2 Materials and methods

We structured the task of cell-type-specific enhancer discovery as a weakly supervised object detection problem with two modules. First, we constructed a CNN binary classifier to predict the existence of enhancers. The model takes a matrix of high-resolution epigenetic features over a large genomic window as input. Second, we developed an object detection module to locate the enhancer boundaries in the positive genomic windows based on the most informative subset of epigenetic features indicated by Grad-CAM. With a trained model, we can carry out cell-type-specific enhancer discovery in a novel cell type with common epigenetic profiles and obtain high-resolution core enhancer coordinates. We benchmarked and validated our framework using various internal and external evaluation metrics. By evaluating through different biological perspectives, we demonstrate DECODE’s ability to generate high-quality cell-type-specific enhancer annotations with a strong regulatory impact.

2.1 Training data processing

We collected STARR-seq data for five human cell lines (HepG2, K562, A549, MCF-7, HCT116), along with chromatin accessibility (ATAC-seq and/or DNase-seq) and ChIP-seq for H3K27ac, H3K4me3, H3K4me1 and H3K9ac, from the ENCODE data portal (Appendix Table A1) (ENCODE Project Consortium and others, 2004). To call STARR-seq peaks, we applied STARRpeaker, which adjusts for GC content and RNA thermodynamic stability during peak calling (Lee et al., 2020). STARR-seq peaks overlapping with a chromatin peak and a peak of an active histone enhancer mark were defined as active enhancers and were considered positive training samples (Zhang et al., 2008). Negative regions were down-sampled from the background at a 1:10 positive to negative ratio. The positive and negative samples were extended to 4 kb, and the signals were aggregated over 10 bp bins.

The resolution of signal aggregation determines the precision of boundary detection. Here, every value in the input matrix represents the average epigenetic signal of a 10 bp bin. In the end, each input value is assigned a Grad-CAM importance score. Hence, filtering by the importance score obtains core enhancers at a 10 bp resolution. It is possible to extract higher-resolution signals for higher precision, but 10 bp was the experimentally determined optimum resolution and the highest resolution for most ChIP-seq experiments.

For the ATAC-seq version, there were 211 097 STARR-seq peaks and 459 321 ATAC-seq peaks for both (HepG2, K562) cell types. Only 23 420 of the STARR-seq and ATAC-seq overlap regions intersected with another ChIP-seq, which were selected as the positives. For the DNase-seq version, there were 912 967 DNase-seq peaks for all five cell types. Only 73 271 of the STARR-seq and ATAC-seq overlap regions intersected with another ChIP-seq, which were selected as the positives. For the selected positive regions, we observed a distinct signal shape for each assay. The peak in chromatin accessibility and peak-trough-peak in other histone marks validate our selection for the training regions and provide a basis for CNN pattern recognition (Fig. 5).

2.2 Binary classifier construction

As shown in Figure 1, the model is a ResNet-inspired CNN that contains convolutional layers, pooling layers and dense fully connected layers (He et al., 2016). The input is a data matrix (of size $5 \times 400$) containing values from the signal tracks of the five epigenetic assays extracted from a 4 kb region by aggregating the signals over 10 bp bins. Each value in the input matrix represents the signal of an epigenetic assay at a genomic loci. We use only epigenetic features because they are more generalizable compared to sequence-based features, especially when transferring predictions to unseen cell types (Zhou et al., 2011). Our model contains seven convolutional layers, each of which uses its $k$ convolutional filters to produce $k$ activation maps of width $i$ and height $h$:

$$A^i = W^i \cdot X^i + b^i$$  \hspace{1cm} (1)$$

The first several convolutional layers extensively capture altitude and shape-based features from either chromatin accessibility or ChIP-seq with different convolution filters. Moreover, the filters blend signals across different tracks to allow for combinatorial feature extraction. Then, max-pooling layers are used to reduce the number of parameters and abstract features trained in the previous convolutional layers. If $b, w$ denotes the dimensions of a pooling
operation, then max-pooling over the activation map takes the max value of each $h \times w$ window to produce an output of size $X^{l+1} = \text{max-pooling}(A^{k,l})$ as input for the next layer.

$$X^{l+1} = \max - \text{pooling}(A^{k,l})$$

(2)

We hypothesize that the interaction of open chromatin states and histone marks coordinate TF binding in enhancers (Spitz and Furlong, 2012). Therefore, we placed chromatin accessibility in the middle of the input data matrix to allow for maximal interaction with other histone marks. The first few filters have a kernel size of $5 \times 10$ in order to span all five assays. With padding removed, the features are then convoluted and down-sampled to one dimension to represent linear genomic windows. Further convolutions on the linear feature use a one-dimensional (1D) filter of size $1 \times 4$. The pooled layers are then fed into fully connected layers to make a sigmoid prediction on the probability of enhancers being in the region.

$$P(\text{enhancer}) = \frac{1}{1 + e^{-X}}$$

(3)

Between every convolution layer are squeeze-and-excitation blocks that calculate the residual features and act as a gate for how much original feature is passed through (Hu et al., 2018).

2.3 Object boundary detection via weakly supervised learning framework (Grad-CAM)

We operationalized the task of enhancer localization using a weakly supervised object detection method in computer vision. Grad-CAM extracts the implicit localization of the target from classification models and obtains a high-resolution subset of the image with the most informative content regarding the target (Selvaraju et al., 2017). For a genomic region with a positive classification, we used Grad-CAM to extract the implicit enhancer localization as a subset of the original input genomic region, thereby increasing the resolution of our core enhancer annotations (Fig. 2). Utilizing Grad-CAM to revisit the positive predictions can refine our annotations by finding the most salient enhancer regions. Further, our method is much more interpretable compared to previous supervised blackbox prediction models because we can trace and visualize the process of decision-making in our network.

During the training process, each convolutional filter learns to extract features that are important in accurately predicting enhancers in the genomic window and outputs an activation feature map. Each activation feature map highlights genomic regions that contain important features for enhancer prediction. Superimposing all activation feature maps sums together all the highlighted regions and forms a silhouette of locations, with respect to the original input, that activated the greatest number of neurons in our neural network, thereby extracting the implicit localization information underlying a classification task.

In detail, we produce a scalar importance score for each activation map using the global-average-pooled gradient of the positive class with respect to the feature map activation (Equation 4).
\[ a'_k = \frac{1}{Z} \sum_i \sum_j \frac{\partial y^{i,j-1}}{\partial A^k_{i,j}} \]  

We use the scalar importance scores as weights for the linear combination of all k activation feature maps. We then multiply each importance score with its respective activation feature map and sum element-wise over all activation feature maps in the layer. Next, we apply RELU to filter for only positive values, and interpolate to obtain a Grad-CAM score map of size \( i \times j \), the same size as the input of the convolutional layers (Equation 5).

\[ L_{Grad-CAM} = \text{ReLU} \left( \sum_k a'_k A^k \right) \]  

Grad-CAM maps can be generated for different convolutional layers in our architecture. Concretely, the Grad-CAM map from the last 1D convolutional layer is a 1D map, or a position-wise score, describing the highest-level feature importance of each 10 bp bin in the genomic window. To refine our predictions, we use the position-wise score to select a subset of the input region (or a subset of the 10 bp bins) that corresponds to a high Grad-CAM score. Here, our cutoff is set as the average Grad-CAM score over all positive genomic positions; hence, the filtering is performed after all predictions have been processed by Grad-CAM. Furthermore, Grad-CAM maps can also be generated using gradients from the first few convolutional layers to observe how much each epigenetic assay contributes to a positive prediction, or a feature-wise score. We constructed our custom CNN such that the high- and low-level Grad-CAM maps correspond to position-wise or feature-wise biological interpretations, respectively. This enables us to study the epigenetic features of enhancers by visualizing and interpreting the biological process of decision-making of our model.

2.4 Model training configurations

We used the Adam optimizer at a learning rate of \( 5e-5 \) and added dropout layers to prevent overfitting. The model is set to train for 100 epochs, but we included early stopping with monitoring of validation loss to prevent overfitting. The data was split 80–20 for training and validation. We also added class weights to address the 1:10 positive and negative sample imbalance in our dataset. Training and model prediction were accelerated using NVIDIA Tesla K80 GPUs.

2.5 Enhancer challenge data for benchmarking against the state-of-the-art model

We used transgenic mouse enhancer data from the VISTA enhancer database to benchmark against the state-of-the-art method (Visel et al., 2007). Hypothesized enhancers were cloned into a plasmid with a promoter and lacZ reporter gene, which were injected into mouse embryos. After reimplantation with surrogate mothers, the transgenic embryos were collected at e11.5 to score for enhancer activity. Target regions were considered positive for enhancers if at least three transgenic embryos had reporter-gene expression across three sample tissues and were considered negative if we observed no reproducible pattern across at least five samples. Tissue-specific enhancers were pooled from all six tissues (forebrain, heart, hindbrain, limb, midbrain, neural tube). In order to mediate cross-species effects, DECODE was trained with all available human cell line data, and then fine-tuned with out-of-sample mouse enhancers.

2.6 Cell-line case study evaluation

We utilized a variety of methods to evaluate the accuracy of our supervised model and the regulatory impact of our core enhancer annotations. As a case study, we predicted enhancers in neural progenitor cells (NPCs), which lack STARR-seq data. These cells play an important role in psychiatric disorders.

The NPC signal tracks for DNase-seq and four active histone marks (H3K27ac, H3K4me3, H3K9ac, H3K4me1) were downloaded from the ENCODE portal. We created sliding windows of size 4 kb with 500 bp steps across the whole genome. For DECODE classifier input, the average signal was extracted for 10 bp bins in each genomic window. After classification, the positive 4 kb genomic windows made up the original annotation. We then applied Grad-CAM to refine each positive 4 kb window to define our core enhancer annotation.

GWAS LDSC Enrichment: We characterized the disease-variant impact of our NPC annotations by calculating stratified LDSC enrichment from GWAS (Bulik-Sullivan et al., 2015). LDSC regresses the chi-square statistics (\( \chi^2 \)) with the linkage disequilibrium (LD, \( r^2 \)) to estimate the heritability in a disease-specific manner. This method calculates the partitioned heritability of certain regions or annotations using GWAS summary statistics. We utilized LD scores from the 1000 Genomes Project and GWAS summary statistics from Bulik-Sullivan et al. and the Psychiatric Genomic Consortium (Siva, 2008; Turley et al., 2018).

Conservation Score Analysis: We measured inter- and intra-species conservation by 100-way PhastCons and rare derived allele frequency (DAF) SNP enrichment, respectively. The PhastCons score is a phylogenetic hidden Markov model trained on genetic sequences across 100 different species and quantifies the conservation of a given genetic annotation (Yang, 1995). We calculated the ratio of DAF (\( <0.5\% \)) SNP enrichment using SNPs from the Genome Aggregation Database (gnomAD) and Pan-Cancer Analysis of Whole Genomes (PCAWG) resources (Campbell et al., 2020; Karczewski et al., 2020).

3 Results

In contrast to traditional enhancer classification methods, we reformulated enhancer discovery into a two-step weakly supervised object detection problem. Specifically, we first utilized the direct enhancer activity readout from novel functional characterization assays to train a deep neural network. Sections 3.1 and 3.2 demonstrate the ability of our deep learning binary classifier to make accurate predictions. Then, for the object detection module, we used Grad-CAM to define high-resolution enhancer boundaries; Sections 3.3–3.6 describe our results for enhancer localization. In short, we applied our two-step DECODE model to various real-world datasets for comprehensive performance benchmarking and demonstrate its benefits in constructing compact genome annotations to facilitate variant interpretations.

3.1 The DECODE binary classifier is a transferrable model for accurate cell-type-specific enhancer prediction

A trained binary classifier can predict enhancers on sliding windows across the genome. We merged and shuffled positive- and negative-labeled data from all cell types to train a binary classifier for predicting the existence of enhancers in any given 4 kb genomic region using cell-type-matched epigenetic features (details in Section 2.4). To verify our results, we performed fivefold cross-validation by partitioning the merged data into fivefolds and iteratively using each fold as the out-of-sample validation set. High out-of-sample area under the receiver operating characteristic curve (auROC; ATAC-seq: 0.972, DNase-seq: 0.989) metrics demonstrate the ability of the binary classifier module in DECODE to accurately predict enhancers using combinatorial epigenetic features. In addition, we did not observe any divergence between training and validation loss during backpropagation. Moreover, the validation metrics remained high across all folds (>0.95), suggesting that there was no overfitting under our training configurations.

In real-world scenarios, a model would predict enhancers in cell types that it has not yet seen during training. Therefore, we further tested the robustness of the DECODE binary classifier in transferring the learned features to new cell types. We also performed leave-one-chromosome-out cross-validation to evaluate the process by which DECODE generalizes to new genomic loci (Schreiber et al., 2020). By leaving out one cell type or chromosome for validation, we trained the model using the rest of the cell types (for both
ATAC-seq and DNase-seq) or chromosomes (for DNase-seq only) and evaluated it using the out-of-sample cell type or chromosome. Similar to the validation performance from merged data, we observed consistently high cross-cell-type and cross-chromosome validation metrics, demonstrating our model’s ability to transfer its predictions across cell types and genomic loci (Fig. 3). For instance, our DECODE model trained on DNase-seq and four other ChIP-seq datasets achieved consistently high validation auROC (0.996–0.999) and auPRC (0.954–0.998) scores. Cross cell-line performance remained high even when using ATAC-seq for chromatin accessibility. Therefore, our DECODE model can accurately predict enhancers and robustly transfer the predictions onto novel, unexplored cell lines and loci.

3.2 Decode outperforms an existing state-of-the-art method on experimentally validated mouse enhancers

In addition to making accurate predictions for internal evaluations, we further compared the efficacy of our model with existing methods on an external experimentally validated dataset. To do so, we applied the DECODE binary classifier on 3244 experimentally validated regions from six mouse tissues, which were also used in the official ENCODE enhancer challenge. Specifically, we downloaded the signal tracks for the five epigenetic features from the ENCODE portal, extracted the signals for each given region as input, and predicted for the existence of enhancers using our trained model. We compared our predictions with Matched-Filter, a leading method in the state-of-the-art model using transgenic mouse enhancers.

Benchmarking Against Matched-Filter-Based Model: We benchmarked our trained model against the state-of-the-art model using transgenic mouse enhancers. DECODE produced validation metrics (a, auROC; b, auPRC) that outperformed the state-of-the-art model in all tissue types.

As shown in Figure 4, our DECODE model obtained an average auPRC of 0.46, which was 24% higher than the auPRC of Matched-Filter on the same dataset (Fig. 4a). It is also worth pointing out that DECODE outperformed Matched-Filter in all six tissues with a decent margin. Specifically, our model demonstrated higher auROC scores ranging from 0.82 to 0.85 in all six tissues (versus 0.76–0.85 for Matched-Filter) and noticeably improved auPRC scores (0.39–0.57 in DECODE versus 0.27–0.43 in Matched-Filter) with an average margin ranging from 0.02 to 0.18.

Two main reasons could explain DECODE’s improvement in performance over existing methods. First, we used genome-wide large-scale training data from direct human enhancer readouts of five cell types compared to the Drosophila data used for the training of Matched-Filter. This allows our model to recognize more complicated features that are important for enhancer predictions. Second, we believe the interactions among epigenetic features guarantee active regulatory activity in functional enhancer regions, which has been demonstrated in previous literature (Spitz and Furlong, 2012). As a result, we designed the convolutional filters in our deep learning framework to span multiple epigenetic marks to model non-linear epigenetic interactions. In contrast, Matched-Filter considers epigenetic marks independently with its linear support vector machine-based methods. The capacity of DECODE to learn complex combinations of features provides the basis for achieving better performance than the current state-of-the-art method (Supplementary Figs S4 and S5).

3.3 DECODE’s object detection module generates interpretable visual explanations for enhancer boundary refinements

After demonstrating the efficacy of DECODE’s binary classifier for accurate enhancer predictions, we seek to uncover more information regarding the basis of neural network decisions through our weakly supervised framework. The DECODE object detection module extracts interpretable feature-wise and position-wise importance scores as a visual explanation. Here, we examine the benefits of feature-wise importance scores generated by Grad-CAM from the lower-level convolutional layers.

Grad-CAM uses gradients to identify input locations that activate the greatest number of neurons in a given layer. We extracted the feature-wise Grad-CAM scores from the positive training samples to assess the basis of model predictions. A high feature-wise Grad-CAM score corresponds to higher importance placed on that epigenetic feature. Grad-CAM scores peak at the center of the 4 kb windows for each epigenetic assay (Fig. 5), which corresponds to the greatest amount of regulatory activity as indicated by the original signal. This result shows that DECODE predicts the presence of enhancers using highly active regulatory regions.

Next, we compared the Grad-CAM scores across different epigenetic features to show that our model prioritizes key features for enhancer prediction. As shown in Figure 5, DNase-seq and H3K27ac ChIP-seq demonstrate the highest Grad-CAM scores (mean at center ≫0.35) as compared to other features (mean at center ≲0.25), indicating their important role in defining enhancers in the genome. Our findings recapitulate known biology, as these features were also used in the official ENCODE3 encyclopedia annotation (Moore et al., 2020). In contrast, H3K4me1 exhibited the lowest feature-wise Grad-CAM score (mean from 0.05 to 0.2),
which implies that it played a relatively less important role in our model decision.

3.4 DECODE provides high-resolution enhancer boundary localization

Position-wise Grad-CAM importance scores from DECODE’s object detection module can be used to interpolate high-resolution cell-type-specific enhancer coordinates and condense the annotation of an enhancer to its core functional regions. In contrast, this feature is missing in most existing methods.

To demonstrate this function, we utilized DECODE to predict compact enhancers in NPCs, which play important roles in neurodevelopment and have been implicated in a wide variety of psychiatric disorders (Castreñ, 2014; Das et al., 2020). Specifically, we utilized the five epigenetic marks on NPCs to predict enhancers using our DECODE framework trained on all available data (see details in Section 2.4). We divided the genome into 4 kb windows with 500 bp steps between each window. The corresponding epigenetic signals were extracted at a 10 bp resolution from each window as inputs to our model. Windows with a binary classifier output greater than 0.5 were identified as positive predictions. In total, we identified 1,515,431 overlapping windows across the genome, and 1.1% (17,622) among them showed predicted enhancer activities. We merged the positive predictions to create the original prediction set, which contained 16,522 elements with a mean length of 4,188 bp.

To condense the annotation using DECODE’s object detection module, we extracted the position-wise Grad-CAM scores from the positive regions for each 10 bp bin (Fig. 7a). Bins with Grad-CAM scores larger than the mean across all windows were merged to create the refined set. This process resulted in a total of 23,505 refined positive elements with a mean length of 371 bp.

In Figure 6, we show an example positive genomic window (chr1:8680000–8684000) and its feature-wise and position-wise Grad-CAM values. The example shows the process by which high-level position-wise scores are derived from low-level feature-wise scores, which increases interpretability by tracking neuron activation through our neural network classifier.

In addition, the true enhancer in the example is shifted to the right of the genomic window. Due to the lack of a quantitative boundary detection algorithm, most existing methods take the entire input window as an enhancer region, which potentially confounds various downstream analyses such as validation region selection and disease causal variant mapping. In contrast, the object detection module in our DECODE model does not rely on interpolating from the center, but rather localizes enhancer coordinates based on the importance of the loci using the Grad-CAM outputs. Therefore, we are still able to rescue the shifted enhancers and discover true functional regions with high regulatory impacts.

This boundary detection module can remove a significant portion of the noise regions in our enhancer prediction and noticeably condense our genome annotation. For instance, analysis on the refined set shows that it is only 12.6% in coverage as compared to the original set but includes a disproportionately large amount (71%) of transcription start sites, which indicate a more enriched transcriptional regulatory footprint (Fig. 7b). In the following sections, we demonstrate the compactness of the refined set through conservation and GWAS variant enrichment analysis.

3.5 DECODE’s compact enhancer predictions are highly conserved across species and populations

To test whether the object detection module accurately selects true regulatory regions, we compared cross-species conservation scores of the original and refined enhancer regions. If the enhancer annotations serve important regulatory functions, then those annotations should be conserved, as any negative mutations would increase the likelihood of disadvantage phenotypes that eliminate that allele from the gene pool. Hence, higher conservation usually indicates higher enhancer quality (Yang, 1995).

Specifically, we downloaded the 100-way PhastCons signals and calculated the average PhastCons conservation scores within the original and refined regions. The median PhastCons score in the refined regions was 0.117, which is significantly higher than those in
the original regions (median 0.092, one-sided Wilcoxon test \(P < 0.001\), Fig. 7c). The original regions also exhibited higher conservation compared to other existing annotations [i.e. ENCODE NPC candidate cis-regulatory elements (cCREs), FANTOM human enhancers and FANTOM neuronal stem cell enhancers; Supplementary Fig. S1].

We further evaluated the quality of our compact enhancer annotation from DECODE via the enrichment of rare variants with a simple but validated assumption—key functional regions in the genome are under strong negative selection and hence are depleted in common variants (Fu et al., 2014; Zhang et al., 2020b). Therefore, we compared the rare variant enrichment in the refined and original positive enhancer regions (and against ENCODE NPC cCREs, FANTOM human enhancers, FANTOM neuronal stem cell enhancers in Supplementary Fig. S2). Specifically, we downloaded the entire human genetic variation set from gnomAD and PCAWG (Campbell et al., 2020; Karczewski et al., 2020), and defined rare variants as those with a DAF less than 0.5% over the entire population. We calculated the percentage of rare variants within each merged annotation set. The refined enhancer regions demonstrated significantly higher percentages of rare variants in both the gnomAD and PCAWG dataset. For instance, we observed a rare variant percentage of 0.922 and 0.857 in gnomAD and PCAWG, respectively; this number decreased to 0.910 and 0.843 for the original positive input regions \((P < 10^{-3.22})\) for binomial test in both datasets).

Altogether, the condensed compact enhancers refined in our model showed higher cross-species and cross-population conservations, indicating DECODE’s ability to remove noisy regions in our enhancer predictions and provide high-quality genome annotations.

### 3.6 Compact enhancer annotations predicted by DECODE can better explore GWAS variants in neurodevelopmental and psychiatric disorders

Disease-causal variant mapping is one of the most important applications of distal regulatory element mapping. Lines of evidence have demonstrated that accurate and compact annotations can significantly increase the statistical power for both somatic and germline variant mapping in disease studies (Fu et al., 2014; Zhang et al., 2020a,b). Therefore, we further tested whether our condensed enhancer definitions can benefit variant prioritization and interpretations.

Here, we predicted two sets of enhancers in NPCs—a coarse set of predictions using the binary classifier (similar to existing methods) and a refined set of core predictions using the DECODE object detection module. We extracted the summary statistics from GWAS for around three million SNPs for nine traits. For each phenotype, we used stratified LDSC to test whether the heritability of a GWAS phenotype is enriched in one set of annotated genome regions in NPCs, where a high LDSC enrichment for a GWAS trait would indicate that the set of annotations has a high partitioned heritability for the corresponding trait. Using these summary statistics, we calculated the enrichment of the original and refined set for each trait, represented by the \(P\)-value (Fig. 7d).

We found that both annotations demonstrated significant LDSC enrichment in three of the nine phenotypes (with \(log P\)-value ranging from 5.28 to 12.87), but the refined set consistently showed higher LDSC enrichment for all five neurodevelopmental and psychiatric phenotypes. For example, the \(P\)-value enrichment of the original set for bipolar disorder was 0.052, while the condensed enhancer set increased the statistical power by about \(10^{5} \times (P\)-value < 0.005). Even with only 12.6% of the coverage, the refined set improved the overall quality of the annotations and obtained a range of 2- to 10-fold increases in GWAS enrichment compared to the original set of annotations. We observed no enrichment for cardiovascular, metabolic or immunological phenotypes, demonstrating DECODE’s ability to discover cell-specific enhancers. Both predictions also exhibited higher LDSC enrichment compared to other existing annotations [i.e. ENCODE NPC cCREs, FANTOM human enhancers and FANTOM neuronal stem cell enhancers; Supplementary Fig. S3].

We believe the compactness of our refined annotation accounts for the increase in statistical power compared to the original set. Because of Grad-CAM, we are able to remove regions of variable lengths that do not contribute to the identity of an enhancer and...
shrink the prediction to 12.6% of its original size. Therefore, the refined annotations are more condensed and are more suitable for a wider range of analyses because many calculations, such as functional validation, require compact definitions to obtain statistical power.

4 Discussion

Here, we propose a DECODE. Our model has two distinct parts: a binary classifier and an object detection module, both of which provide substantial benefits over previous methods.

For the binary classifier, we trained a deep learning model on direct readouts of human enhancer data to classify enhancer windows based on common epigenetic profiles. The classifier outperformed the state-of-the-art method in predicting cell-type-specific enhancers by using a larger set of training data and a deep learning model. CNNs have a larger capacity to learn non-linear, complex feature interactions compared to previous linear methods. We also emphasize that our deep learning-based DECODE model will benefit from the rapid development of novel functional characterization assays (e.g. MPRA and CRISPR-based screens) and the exponential growth of training data to further improve the accuracy and performance in enhancer predictions.

In addition to an improvement in prediction accuracy, our DECODE model also has a unique boundary detection module via Grad-CAM, which is not found in previous methods. The resultant feature-wise importance scores increase the interpretability by visualizing feature prioritization, while the position-wise importance scores can be used to condense the coarse enhancer annotations to the core functional regions. In particular, we show that our compact enhancer definition has strong regulatory impact and are essential for disease causal variant mapping in disease studies.

In summary, we introduce a powerful tool that could be widely deployed for enhancer discovery. With corresponding epigenetic features, DECODE can not only accurately predict the existence of enhancers in disease causal variant mapping in disease studies. We also model. CNNs have a larger capacity to learn non-linear, complex feature interactions compared to previous linear methods. We also emphasize that our deep learning-based DECODE model will benefit from the rapid development of novel functional characterization assays (e.g. MPRA and CRISPR-based screens) and the exponential growth of training data to further improve the accuracy and performance in enhancer predictions.

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Appendix A

Table A1. Data availability matrix from ENCODE. Data was available for all four histone marks across all cell types, but ATAC-seq was only available for K562 and HepG2 cells

| Cell Type | STARR-seq | ATAC-seq | DNase-seq | H3K27ac ChIP-seq | H3K4me3 ChIP-seq | H3K4me1 ChIP-seq | H3K9ac ChIP-seq |
|-----------|-----------|----------|-----------|------------------|-----------------|------------------|----------------|
| K562      | ✓         | ✓        | ✓         | ✓                | ✓               | ✓                | ✓              |
| HepG2     | ✓         | ✓        | ✓         | ✓                | ✓               | ✓                | ✓              |
| A549      | ✓         | ✓        | ✓         | ✓                | ✓               | ✓                | ✓              |
| HCT116    | ✓         | ✓        | ✓         | ✓                | ✓               | ✓                | ✓              |
| MCF-7     | ✓         | ✓        | ✓         | ✓                | ✓               | ✓                | ✓              |