"Ultra Chrome"

Unique Deep CNN model for predicting gene expression from histone modifications

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Abstract-Histone modifications the most important factors that control gene regulation. Computational methods that predict gene expression from histone modification signals are highly desirable for understanding their combinatorial effects in gene regulation. This knowledge can help in developing "epigenetic drugs" for diseases like cancer. Previous studies for quantifying the relationship between histone modifications and gene expression levels either failed to capture combinatorial effects or relied on multiple methods that separate predictions and combinatorial analysis. This paper develops a unified discriminative framework using a deep convolutional neural network to classify gene expression using histone modification data as input. Our system, called DeepChrome, allows automatic extraction of complex interactions among important features. To simultaneously visualize the combinatorial interactions among histone modifications, we propose a novel optimization-based technique that generates feature pattern maps from the learnt deep model. This provides an intuitive description of underlying epigenetic mechanisms that regulate genes.

We show that DeepChrome outperforms state-of-the-art models like Support Vector Machines and Random Forests for gene expression classification task on 56 different cell-types from REMC database. The output of our visualization technique not only validates the previous observations but also allows novel insights about combinatorial interactions among histone modification marks, some of which have recently been observed by experimental studies.

Index Terms—Histone Modifications, Deep Learning, Convolutional Neural Networks, Recurrent Neural Networks.

I. Introduction

Gene regulation and the Histone Code Hypothesis

Gene regulation is a complex process that controls gene expression to be high or low. One of the most important factor that influence gene regulation is histone modifications. Histones are basic proteins that prevent DNA damage by wrapping it up in tight chromatin packs. Their special function directly determines the spatial arrangement of the DNA causing the activation or deactivation of different binding locations and leading to changes in gene regulation. The importance of these epigenetic changes are supported by evidence that particular modifications profiles are linked to cancer (*Bannister* (2011) [1]).

However, unlike DNA mutation, these changes are reversible. This fact makes the study of histone modifications interesting for the development of "epigenetic drugs".

The role of histone modifications in gene regulation is

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expressed by the **Histone Code Hypothesis** (*Jenuwein et al.* (2001) [2]), which states that modifications of histone proteins (changes to DNA scaffolding) are directly involved in DNA transcription and gene regulation.

Previous studied already investigated the correlation between histone modifications and gene expression. The problem now has shifted to predicting if a gene is expressed or not solely based on the histone modifications signals.

Contributions

In this paper I present *UltraChrome* a Deep CNN model that extends previous deep learning architectures and whose main contributions can be summarized as follows:

- Is the first implementation of Spatial (2D) Convolutional Neural Network used to tackle gene expression prediction tasks
- Generalize over the cell type leading to one unique and comprehensive model for all the available cell types
- Sightly improves previous state-of-the-art Deep Learning models if compared over all the available cell types

II. RELATED WORK

In the last decade, different studies addressed the combinatorial interactions of histone modifications in gene regulation. An initial study confirmed the correlation between histon modifications and gene regulation by *Lim et al.*(2009) [3]. Later *Dong et al.*(2012) [4] implemented Random Forests for classifying gene expression as high or low.

Most recent studies applied deep learning models to automatically learn combinatorial interactions among histone modifications. Singh et al. (2016) [5] introduced DeepChrome, a unified CNN framework and the first deep learning implementation for gene expression prediction task. More important than the CNN model itself, the main contribution of DeepChrome was to create a visualization of combinatorial interactions among histone modification signals. One important drawback of DeepChrome is the use of temporal convolution (1D) which takes into consideration only sequences and do not capture local patterns between histone modifications.

Later, *Singh et al.* (2017) [6] developed *AttentiveChrome*: the first attention-based deep learning method applied to this particular task. The novelty of this work is about providing an interpretation of the deep learning black-box model by using the attention scores and visualizing what the model "sees"

when making predictions.

However both of these implementation failed to tackle the predictive challenge from a global point of view and they specialize their deep learning models over one particular cell type at the time.

III. PROCESSING PIPELINE

The input data consists of 56 cells types each of them containing different genes represented by 5 different histone modifications over 100 bins.

In some implementations of my models I merged all the 56 cell types datasets into one comprehensive dataset to train a *global* model. This strategy will be detailed in Sec.VI.

Starting from the raw input generated and formatted by Singh et al. (2016) [5] I transformed the 5×100 matrix in one 3D tensor of shape $5\times 100\times 1$ in order to correctly feed it to the convolution layer at the beginning of the deep learning model. After the convolution I placed some regularization layers and then a multi-layer architecture. The last step is simply a softmax operation to reduce the computation into a binary classification task.

The processing pipeline it graphically summarized by the Fig.1.

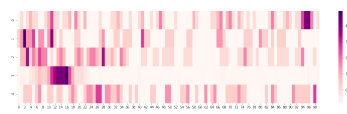


Fig. 1: Processing pipeline

IV. SIGNALS AND FEATURES

My setup is identical to *Singh et al.* (2016) [5]. Starting from the REMC¹ database, they divided the 10 000 basepair (bp) DNA region (65000 bp) around the transcription start site (TSS) of each gene into bins of length 100 bp. Each bin includes 100 bp long adjacent positions flanking the TSS of a gene. In total, they consider five core histone modification marks from REMC database (*Kundaje et al.* (2015) [7]). These five histone modifications are selected as they are uniformly profiled across all cell-types considered in that study. This makes the input for each gene a 5×100 matrix, where columns represent different bins and rows represent histone modifications. For each bin, they report the value of all 5 histone signals as the input features for that bin. A visual example of an input matrix is reported in Fig.2.

Here I formulate the gene expression prediction as a binary classification task. Specifically, the outputs of my models are labels 1 and -1, representing gene expression level as high or low, respectively.

Following what done by Singh et al. (2016) [5] I split the

¹REMC: http://www.roadmapepigenomics.org/

dataset into training (6601), validation (6601) and test (6600). In addition to this procedure I standardized the dataset using mean and standard deviation of the training set. I applied this scaling to improve the performance of the neural networks learning algorithm.

V. LEARNING FRAMEWORK

UltraChrome architecture is composed of different layers. The diagram in Fig.(??) provides a visual reference of the model that can be summarized as follows:

- Input layer: the input is a tensor of shape $5 \times 100 \times 1$.
- Convolution: I applied a spatial (2D) convolution with
 F filters with a kernel size of 5. This operation performs
 a sliding window over all bins positions and across all
 5 histone modifications. The convolution enables the
 model to extract visual, local and global features of the
 distribution of the modifications.
- Pooling: to preserve local features I inserted a max pool layer of shape (1 × 5): the particular choice of the shape aimed to capture the variability just over the bins and not also across the different modifications.
- **Dropout**: I inserted a *dropout* layer with a rate of 0.5 after the pooling to help the generalization ability of the model and reduce overfitting.
- **Flattening**: this flattening step is a necessary step to format the input for the following layers.
- Fully Connected layers: the core of the learning is achieved by exploiting a multi layer, fully connected, architecture composed of 3 different layers with N_1 , N_2 , N_3 neurons each of them with *relu* activation function to produce non-linearities.
- **Batch normalization**: in order to reduce the gradient vanishing problem I inserted 2 *Batch Normalization* layers between the 3 *fully connected* layers.
- Output layer: the final layer is just a Dense layer with one neuron that maps to the desired output in the form of a probability score.

VI. RESULTS

A. Validation of Hyper-parameters

In order to select the best model I chose the hyper parameters validating them on the validation set. I tested different options and values:

- Number of filters: $\{50|100\}$
- Number of neurons per layer:
 - Low: [32, 128, 32] (3 layers)
 - High: [64, 256, 64] (3 layers)
 - Deep: [64, 256, 256, 64] (4 layers)
- Dropout: $\{Yes|No\}$
- Batch Normalization: $\{Yes|No\}$

In the Tab. 1 are reported a compact subset of the most significative results of the validation step. As we can see, the best combination is obtain with F=100 filters, high number of neurons (64, 256, 64 respectively), dropout with probability

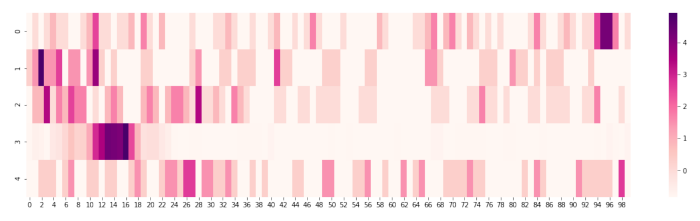


Fig. 2: Example of the matrix representation of the input

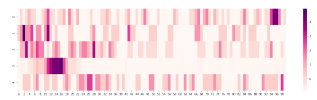


Fig. 3: UltraChrome Architecture

Hyper-parameters				AUC validation set		
Filters	Neurons	Dropout	BN	Min	Max	Mean
50	Low	No	No	0.653	0.935	0.764
100	Low	No	No	0.653	0.939	0.765
100	High	No	No	0.653	0.936	0.763
100	High	Yes	No	0.654	0.933	0.763
100	High	Yes	Yes	0.652	0.937	0.766
100	Deep	Yes	Yes	0.648	0.937	0.765

TABLE 1: Hyper parameter tuning for *UltraChrome* on validation set

0.5 and batch normalization. However the differences are not large from one combination to another: small changes in the second part of the architecture (the fully connected multilayers) are not so relevant on the overall results, suggesting that the main contribution maybe come from the convolution itself rather than the particular deep models. Moreover, the last combination reported, with a deeper architecture, do not lead to better results, but only increases the complexity of the network.

B. Other models

While developing my main model *UltraChrome* I tested other models:

- Random Forest Classifier (RFC) with 150 estimators and max depth of 20. This model was used in one of the first attempts to investigate correlation between histone modifications and gene expression, as reported in this paper (???).
- Base Neural Network (BNN): simple Deep Neural Network with two fully connected layers with 64 and

- 32 neurons respectively. This naive model was used as a baseline for more complex architectures.
- ChromeR (CR): Recurrent Neural Network with a Bidirectional LSTM implementation. In particular, after validation, I used 128 units for the LSTM layer. After the Bidirectional layer I added a Dropout layer with a probability 0.5 to overcome overfitting, then I flattened the data and fed them to two fully connected layers with 625 and 125 neurons respectively and relu activation function.

C. Global and individual models

As mentioned in the introduction I, my contribution is also related to the type of training approach: I trained my models over the full dataset, unaware of the different cell types (the same principle applies to validation and test set). However, in order to compare the original baselines of *DeepChrome* and *AttentiveChrome* I also trained my neural networks in an individual way: for each cell type I trained a model, validated on its set and tested only on his test set. These two configurations could be summarized as *global* and *individual*. Moreover, implementing *DeepChrome* model myself allowed me to reproduce both configurations.

Individual configurations led to better results in terms of AUC score but required to differentiate for the cell type also at test time: this particular setup poses a strong hypothesis on the problem formulation and execution in the sense that we must obtain the cell type separately before testing the unseen data with our model.

D. Test Set AUC

In the Tab 2 are reported the AUC scores for all the implemented methods and the two baselines from previous related work.

As we can see from the results, *AttentiveChrome* is the best *individual* model in terms of mean AUC. However, reporting just the mean AUC averaged over all 56 cell types is not so interesting as we look at only one single score. This is the reason why I also trained my global model individually for all the cell types. This arrangement puts my algorithm on the

Model	Training type	AUC
Random Forest Classifier	Global	0.7942
Base Neural Network	Global	0.7930
ChromeR	Global	0.7961
DeepChrome	Global	0.8032
UltraChrome	Global	0.8045
Base Neural Network	Individual	0.7986
ChromeR	Individual	0.8009
DeepChrome	Individual	0.8068
AttentiveChrome	Individual	0.8115
UltraChrome	Individual	0.8114

TABLE 2: AUC scores comparison on test set

same level as *DeepChrome* and *AttentiveChrome*, enabling me to discriminate the single improvement with single cell precision.

The plot Fig. 4 shows the performance in terms of AUC of the best models for each of the 56 different cell types. The data is organized as follows: first I sorted the types from highest to lowest referring to *AttentiveChrome* scores; second I reported the other models scores following the same sorting strategy. This arrangement has two benefits:

- we can compare single AUC scores between different models
- allows us to identify an underlying trend common to all the models: gene expression on some cell types are easier to predict than others

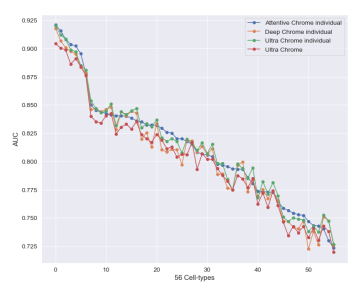


Fig. 4: AUC score on test set

With reference to Fig. 4 we can see that individual *UltraChrome* is almost always better than its global version, and it even compares well with both *DeepChrome* and *AttentiveChrome*.

E. Improvements Matrix

To graphically visualize the number of improvements from one model to another I built an improvements matrix, defined as follows:

$$IM = [x_{ij}] \tag{1}$$

where:

• $i, j \in \mathcal{M}$

 \bullet \mathcal{M} : models set

• x_{ij} : # improvements of model i over model j.

The matrix defined by Eq. (1) is reported in the Fig. (5) as an heatmap for all the models available. The matrix is properly sorted with the best performing model at the top and the darker a row the better a model compares to others. The models set \mathcal{M} is defined as follows: **X-UC** (individual *UltraChrome*), **X-AC** (individual *AttentiveChrome*), **X-DC** (individual *DeepChrome*), **UC** (*UltraChrome*), **X-CR** (individual *ChromeR*), **X-BNN** (individual *Base NN*), **CR** (*ChromeR*), **RFC** (*Random Forest Classifier*), **BNN** (*Base NN*).

Observing the improvements over all the cell types, individual

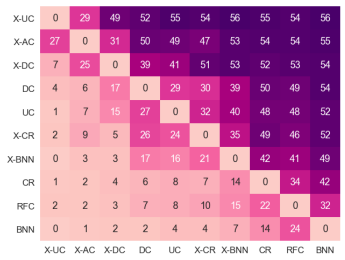


Fig. 5: Improvements Matrix

UltraChrome (X-UC) comes as best model even if compared to AttentiveChrome (X-AC). However if we look at global UltraChrome (UC) the results are not as good, but the different problem setup generates an unfair comparison: the individual models are specifically trained, validated and tested on one cell type at the time; instead UltraChrome reaches good AUC scores in a global scenario.

There are several implications to these results:

- global model incorporates all the cell types characteristics and could describe the phenomenon as a whole
- lower probability of overfitting the model as we increase both the complexity of the task and the amount of training data
- better generalization ability over new cell types

VII. CONCLUDING REMARKS

In this paper I have presented *UltraChrome* a Deep Convolutional Neural Network that can effectively predict the gene expression by simultaneously analyzing spatial distribution of histone signals between all 5 kinds of modifications. Moreover, the model in cell type independent and do not need to know the cell type at test time, improving the generalization ability of the model and extending the previous models with this global approach.

The spatial convolution of *UltraChrome* intercepts patterns and interactions between different modifications. This approach is vital to better capture the gene expression mechanism: as reported by *Bannister et al.* [1], histone modifications can positively or negatively affect other modifications. This phenomenon is called *crosstalk*.

In the future, as new cell types and their relative histone modifications are added to the database we may be interested in instantly predict the gene expression without training a specific model for the new cell type. In this scenario a global model like *UltraChrome* can be useful to assign an initial indication of the gene expression to be refined later on with an individual and cell-specific model.

Future developments could improve the interpretability of *UltraChrome* model: the deep convolution internally maps the input features and extracts meaningful relations, but still miss a way to describe the biology underpinning these processes. This is relevant from a epigenetic point of view. Another improvement could be the integration of new histone modifications in addition to the 5 of the current setup. The new modifications could contribute to shed a light in more complex relations and *crosstalk* effects.

A. Project Remarks

Personally I found the challenge of this project pretty tough mainly because the domain was very specific and required me some time and effort to properly understand the context of the problem.

On the other hand, the project was also instructive. First, I had the chance to develop from scratch deep learning models and test several architectures and hyper-parameters. Second, I was forced to build a complete learning framework, starting from data collection all the way to the communication of the results. In particular I reckon to have been very useful to think about new ways of communicate my results in an effective and elegant manner.

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