Enhancing Hi-C contact map resolution with neural network

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1 Introduction

Recently, the high-throughput chromoseome comformation capture (Hi-C) technique has become a powerful tool for studying the three-dimensional structure of chromosomes. Hi-C data is usually expressed as a $n\times n$ matrix. The resolution of Hi-C data is defined as the bin size of each cell of the matrix. Hi-C data at kilobase level are requisite for future genome 3D structure studies. Rao et al. (2014) generated Hi-C data with 1 kilobase resolution. However, millions of sequenced reads are required to archive this resolution with a huge amount of money and time consumption.

Zhang et al. presented a approach to enhance the resolution of Hi-C data called HiCPlus. Which generated low-resolution data by down-sampling the number of sequenced reads and then a neural network was used to create the mapping between high-resolution contact map and low-resolution contact map.

2 Method

2.1 Experiment Flow

Step 1 Data preparation and processing

Since this experiment is to validate the algorithm for mapping low-resolution data to high-resolution data, high-resolution data is required. In order to compare to some state-of-the-art approaches (HiCPlus and HiCNN), we use data sets (such as GM12878 from GSE63525) which are also used in other approaches . We start from generating 10kb resolution contact map using Hi-C Pro. Then we perform down-sampling on high-resolution data. We use bam files to generate low-resolution contact maps by changing bin size bigger. We generate three contact maps with bin sizes are 20kb, 30kb and 40kb, respectively. We use chromsome 1-8 as traning sets, and chromosome 17 as test set.

Step 2 Learning by Neural network

We separate the low-resolution contact map into many 40×40 submatrices. Those submatrices are used as inputs.

2.2 Layer Structure

We consider the