The ChIPanalyser User's Guide

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Introduction - What is this package?

ChIPanalyser provides a quick an easy method to predict and explain TF binding. The package uses a statistical thermodynamic framework to model the binding of proteins to DNA.

The model assumes that there are four main driver to TF binding:

- Chromatin State affinity
- Binding Energy
- Number of bound Molecules
- Scaling Factor modulating binding affinity

While binding energy is given by PWM matrices, the other parameters will be inferred by maximizing or minimizing a goodness of fit score between ChIP data and ChIPanalyser predictions. To do so, ChIPanalyser provides a genetic algorithm to infer optimal values for each parameter.

Methods - The Chromatin State Model

The chromatin state model is described by the following equation derived from statistical thermodynamics:

$$P(N, a, \lambda, \omega)_{j} = \frac{N \cdot a_{j} \cdot e^{(\frac{1}{\lambda} \cdot \omega_{j})}}{N \cdot a_{j} \cdot e^{(\frac{1}{\lambda} \cdot \omega_{j})} + L \cdot n \cdot [a_{i} \cdot e^{(\frac{1}{\lambda} \cdot \omega_{j})}]_{i}}$$

- \bullet N, the average number of bound molecules
- a_j , chromatin state affinity at site j
- ω , the binding energy required for a TF to bind to site j in the form of a Position Weight Matrix Score
- λ , a scaling factor for the Position Weight Matrix score
- L, the length of the genome of interest
- n, the ploidy level of the organism

Chromatin state affinity is defined as the following:

$$a_j = \sum_k \alpha_k \cdot c_j^k$$

with α the chromatin state affinity scores for a given TF and c the chromatin state at site j.

Shortly, the model above generates ChIP like profiles and describes the affinity of a TF to chromatin states. The affinity scores a, the number of bound molecules N, and lambda λ are inferred by optimising the fit between predicted profiles and ChIP data using a genetic algorithm.

Using ChIPanalyser

Loading data

To demonstrate the use of ChIPanalyser and the genetic algorithm, we provide some internal data sets. We complement this data with external data source such a DNA sequence sets taken from the BSgenomepackage suit.

ChIPanalyser

First, we load ChIPanlyser and the internal data

```
library(ChIPanalyser)
# Input data
data(ChIPanalyserData)
# PFM Matrix
PFM <- file.path(system.file("extdata",package="ChIPanalyser"),"BEAF-32.pfm")</pre>
```

External Data

To reduce the size of the internal data size, we complement our data with the *Drosophila DNA* sequence taken from the BSgenome package suit.

```
library(BSgenome.Dmelanogaster.UCSC.dm6)

DNASequenceSet <- getSeq(BSgenome.Dmelanogaster.UCSC.dm6)</pre>
```

Input data: What is it?

Now we that have loaded some data, what exactly are we looking at? The environment should contain the following new objects:

- chip GRanges object containing ChIP scores. Essentially, ChIP data that will be used to train our model.
- ullet cs GRanges object containing Chromatin State (CS) information. Essentially, where each CS can be found in the genome.
- top GRanges object containing regions of interest. Esssentially, the genomic regions we will use for training and testing.
- *PFM* Path to file containing the Position Frequency Matrix. Essentially, this represents the binding affinity of our TF (here we are using BEAF-32)
- DNASequenceSet DNAStringSet containing the DNASequenceSet of the organism we are looking at.
- geneRef GRanges object containing gene reference taken from the (Genome UCSC website)[https://genome.ucsc.edu/]

Setting Parameters

The next step consists in setting initial parameters to run the genetic algorithm.

```
# Number of individuals per generation
pop <- 10

# Number of generations
gen <- 2</pre>
```

```
# Mutation Probability
mut <- 0.3

# Children - Number of ofspring passed to the next generation
child <- 2

# Method - Goodness of fit metric used to optimise the Genetic algorithm
method <- "MSE"</pre>
```

Please note that the parameters presented here are for the sake of the vignette. We recommend using the following parameters for a first trial with your own data. Please be advised that these parameters will depend on the nature of the data you are using.

```
# Number of individuals per generation
pop <- 100

# Number of generations
gen <- 50

# Mutation Probability
mut <- 0.3

# Children - Number of ofspring passed to the next generation
child <- 10

# Method - Goodness of fit metric used to optimise the Genetic algorithm
method <- "MSE"</pre>
```

We can also define which parameters we wish to optimise. In this example, we will optimise the number of bound molecules (N), lambda (λ) , the PWM Threshold and 11 different chromatin states.

```
# Parameters to optimised
params <- c("N","lambda","PWMThreshold", paste0("CS",seq(1:11)))</pre>
```

Alternatively, we can also set custom ranges for the parameters. We will use custom ranges here to reduce computational time.

```
params_custom <- vector("list", 14)
names(params_custom) <- c("N","lambda","PWMThreshold", paste0("CS",seq(1:11)))

# vector in the format of min value, max value and number of values
params_custom$N <- c(1,1000000,5)

params_custom$lambda <- c(1,5,5)

# Bound between 0 and 1
params_custom$PWMThreshold <- c(0.1,0.9,5)

# Bound between 0 and 1

CS <- c(0,1,5)

CS_loc <- grep("CS",names(params_custom))
for(i in CS_loc){
    params_custom[[i]] <- CS
}</pre>
```

Initializing ChIPanalyser

Building Initial objects

The first step of any ChIPanalyser analysis is to create parameter object. These object contain input parameter and hold basic data. Here, we will load the PFM as PWM and compute the Base Pair Frequency from the DNASequenceSet.

```
GPP <- genomicProfiles(PFM=PFM,PFMFormat="JASPAR", BPFrequency=DNASequenceSet)
```

Generating a starting population

For clairty, we will show what a starting population looks like. This step is not strictly required as the evolve function can take the number of individuals in the population (defined above by pop) and the parameters to be optimised (defined above by params or params_custom).

```
start_pop <- generateStartingPopulation(pop, params_custom)</pre>
```

Pre-processing ChIP data

We will pre-process the ChIP data by reducing noise and converting GRanges to a ChIPscore object. This object contains normalised and smoothed ChIP scores that will be used to train and test the model.

```
chipProfile <- processingChIP(chip,loci = top)

# Splitting data into training and testing
# We recommend setting dist to 20/80. However, here we only have 4 loci.
splitdata <- splitData(chipProfile,dist = c(50,50),as.proportion = TRUE)

trainingSet <- splitdata$trainingSet
testingSet <- splitdata$testingSet</pre>
```

Evolution

We can run the Genetic algorithm provided by ChIPanalyser using a single function. The function will generate intermediate files that allow you to check the status of the algorithm while it is running. It should be noted that the intermediate files are updated at each generation. This method also provides a lambda database. This allows for a faster run time on larger data sets as all values for lambda are pre-computed.

If you do not want intermediate files, set the checkpoint argument to FALSE.

If you do not want to pre-compute lambda values, set the lambda argument to FALSE.

For the purpose of this vignette, we will not save intermediate files.

```
evo <- evolve(population = pop,
    DNASequenceSet = DNASequenceSet,
    ChIPScore = trainingSet,
    genomicProfiles = GPP,
    parameters = params_custom,
    mutationProbability = mut,
    generations = gen,
    offsprings = child,
    chromatinState = cs,
    method = method,
    filename = "This_TF_is_Best_TF",
    checkpoint = FALSE,
    cores= 1)</pre>
```

```
## Generating Lambda DataBase
## Generation: 1
## Generation Fitness: 0.015338136833197
## Generation: 2
## Generation Fitness: 0.015338136833197
```

The output of this function returns a list of 3 elements:

- database a data frame with all parameters that have been computed through out each generation
- poppulation list containing the last generated population
- fitest list containing goodness of metric for best perfomring individuals

Fitest of them all

Once we have the best performing parameters, we can plug them into a single run of ChIPanalyser and use these parameters on a tesing set.

Get fitest individual

The first step is to extract the best performing individual and its associated traits from the population.

Running ChIPanalyser with fitest individual

We can use these parameters as input to the singleRun function to obtain the ChIP like profiles. This run represents the predicted run and TF binding affinity of our TF of choice. We will set fitness to all. This means that the function will return all goodness of fit metrics available.

```
## Warning in ks.test.default(predicted, locusProfile): p-value will be
## approximate in the presence of ties
## Warning in ks.test.default(predicted, locusProfile): p-value will be
## approximate in the presence of ties
```

This final sections returns a list containing 3 elements:

- occupnacy genomicProfiles object containing occupancy scores for each region
- ChIP genomicProfiles object containing ChIP like scores for each region
- gof goodness of fit scores for each regions and mean scores over all regions.

Plotting

Finally, we plot the resulting profiles.

```
par(mfrow = c(2,1))
plotOccupancyProfile(predictedProfile = superFit$ChIP,
      ChIPScore = testingSet,
      chromatinState = cs_single,
      occupancy = superFit$occupancy,
      goodnessOfFit = superFit$gof,
      geneRef = geneRef,
      addLegend = TRUE)
pearson = -0.0768
AUC = 0.3408
MSE = 0.003984
                                                                                                                                   CS6
                                                                                                                                   CS5
CS7
CS2
                                                                                                                 ChIP Profile
Predicted Profile
                                                                                                                                   CS1
CS4
CS11
      transcript
                                                                                                                 Binding Site
      exon
      3UTR
CDS
5UTR
                                                                                                                                   CS10
                                start_codon
                                                         Fer2
      stop_codon
             16140001 16142223 16144445
                                          16146667 16148889 16151112 16153334 16155556 16157778 16160000
                                          Occupancy at Position chr3R 16140001:16160000
 pearson = 0.3825
AUC = 0.8606
MSE = 0.01555
                                                                                                                                   CS2
CS1
CS4
CS11
                                                                                                                 ChIP Profile
                                                                                                                 Predicted Profile
      transcript
                                                                                                                 Binding Site
      exon
                                                                                                                                   CS7
3UTR
     CDS
5UTR
start_codon
      stop_codon
             19500001 19502223 19504445
                                          19506667
                                                   19508889
                                                             19511112 19513334 19515556 19517778 19520000
```

Occupancy at Position chr2L 19500001:19520000