MBASIC: Matrix Based Analysis for State-space Inference and Clustering

Chandler Zuo* Sündüz Keleş[†]

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1 Introduction

This document provides an introduction to the power analysis of ChIP-seq data with the MBASIC package. This R package implements MBASIC which stands for Matrix Based Analysis for State-space Inference and Clustering in [Zuo et al., 2014]. MBASIC provides a Bayesian framework for clustering units based on their infered states over a set of experimental conditions.

MBASIC is especially useful for integrative analysis for ChIP-seq experiments. In this case, a set of prespecified loci is clustered based on their activities over celltypes and transcription factors. We build a pipeline in the MBASIC package and will focus on this pipeline in this vignette. We will introduce the general functionalities for the MBASIC model at the end of the vignette.

2 MBASIC Pipeline for Sequencing Data

2.1 Workflow

The MBASIC framework consists of five major steps:

- 1. Matching ChIP replicate files with their inputs: This step matches ChIP replicate files with their matching input files;
- 2. Calculating mapped counts and genomic scores on the target loci: This step calculates the mapped counts from each ChIP and input replicate files on each of the target locus;
- 3. Calculating the mappability and GC-content scores for the target loci: This step computes the average mappability and GC scores for loci from external files;

^{*}Department of Statistics, 1300 University Avenue, Madison, WI, 53706, USA.

[†]Departments of Statistics and of Biostatistics and Medical Informatics, 1300 University Avenue, Madison, WI, 53706, USA.

- 4. Estimating the background means: This step uses regression models for the input data and the M, GC scores to estimate the background means for the ChIP replicates;
- 5. Fitting MBASIC model: This step fits the MBASIC model to identify the binding states for each locus and cluster the loci based on their binding states across different conditions.

MBASIC integrates Step 2-5 in a single function called "MBASIC.pipeline". For Step 1 MBASIC provides a function "ChIPInputMatch" that assists matching the ChIP files with input files based on the naming convention of the ENCODE datasets. We have found that in practice, more often than not, some violations to the ENCODE file name conventions always occur, and manual adjustments to the results of our automated matching are inevitable. Therefore, we do not integrate this function in "MBASIC.pipeline".

2.2 Match ChIP and Input Datasets

To illustrate Step 1 we first generate a set of synthetic data.

MBASIC package provides a function "generateSyntheticData" to assist our demo. This function generates synthetic BED data for ChIP and input samples, as well as mappability and GC scores in a directory specified by the "dir" argument. It also generates a target set of loci for our analysis. By default, the number of loci is 100, each with size 20 bp. All data are generated across 5 chromosomes, each with size 10K bp. ChIP data are from 2 celltypes, and for each celltype there are K=5 TFs. Under each condition randomly 1-3 replicates for the ChIP data are generated. All ChIP data from the same celltype are matched to the same set of 3 input replicates.

```
> library(MBASIC)
> target <- generateSyntheticData(dir = "syntheticData")
GRanges with 100 ranges and 0 metadata columns:
        seqnames
                        ranges strand
            <Rle>
                     <IRanges>
                                 <Rle>
    [1]
             chr1 [ 200, 220]
    [2]
             chr1 [1800, 1820]
             chr1 [2440, 2460]
    [3]
             chr1 [2480, 2500]
    [4]
    [5]
             chr1 [2960, 2980]
    . . .
             chr5 [7400, 7420]
   [96]
   [97]
             chr5 [8720, 8740]
             chr5 [8880, 8900]
   [98]
             chr5 [9560, 9580]
   [99]
  [100]
             chr5 [9970, 9990]
  seqlengths:
   chr1 chr2 chr3 chr4 chr5
          NA
                NA
                     NA
> system("ls syntheticData/*/*")
```

Function "ChIPInputMatch" assists Step 1. It reads all files with suffix ".bed" in directories specified by the argument ".dir", and matches the files assuming ENCODE naming convention. It looks up files up to the number of levels of subdirectories specified by "depth". The output of this function contains multiple columns. The first column contains the file name for each ChIP replicate. The second column is the initial string for the matching input replicates, because for each ChIP replicate there are possibly multiple input replicates. The rest of the columns contains information for lab, experiment identifier, factor and control identifier. This information is parsed from the file names.

```
4 syntheticData/chip/wgEncodeLabExpCell1Fac1CtrlAlnRep2.bed
5 syntheticData/chip/wgEncodeLabExpCell1Fac1CtrlAlnRep2.bed
6 syntheticData/chip/wgEncodeLabExpCell1Fac1CtrlAlnRep2.bed
                                          inputfile
1 syntheticData/input/wgEncodeLabExpCell1InputCtrl
                                               <NA>
4 syntheticData/input/wgEncodeLabExpCell1InputCtrl
5
                                               <NA>
  lab experiment cell factor control chipformat
1 Lab
             Exp Cell1
                         Fac1
                                 Ctrl
2 Lab
             Exp Cell1
                         Fac1
                                 Ctrl
                                              BED
3 Lab
             Exp Cell1
                                 Ctrl
                                              BED
                         Fac1
4 Lab
                                              BED
             Exp Cell1
                         Fac1
                                 Ctrl
             Exp Cell1
                         Fac1
                                 Ctrl
                                              BED
5 Lab
6 Lab
             Exp Cell1
                         Fac1
                                 Ctrl
                                              BED
  inputformat
          BED
2
          BED
3
          BED
4
          BED
5
          BED
          BED
```

We also need to prepare the following meta data information. Below the vector "n" specifies the number of replicates within the same experimental conditions.

```
> conds <- paste(tbl$cell, tbl$factor, sep = ".")</pre>
```

Now we are in a position to continue the next steps in the pipeline. There are two ways to execute these steps: (1) use function "MBASIC.pipeline", which wraps up all the consecutive steps; or (2) execute each step separately.

2.3 Pipeline Execution

The following code calls the function "MBASIC.pipeline":

```
> MBASIC.fit <- MBASIC.pipeline(chipfile = tbl$chipfile,
                                 inputfile = tbl$inputfile,
                                 input.suffix = ".bed",
+
                                 target = target,
                                 chipformat = tbl$chipformat,
                                 inputformat = tbl$inputformat,
+
                                 fragLen = 150,
                                 pairedEnd = FALSE,
                                 unique = TRUE,
                                 m.prefix = "syntheticData/mgc/",
                                 m.suffix = "_M.txt",
                                 gc.prefix = "syntheticData/mgc/",
                                 gc.suffix = "_GC.txt",
                                 fac = conds,
                                 struct = NULL, J = 3,
                                 family = "negbin",
                                 burnin = 20, maxitr = 100,
                                 tol = 1e-4, nsig = 2,
                                 datafile = NULL)
```

"MBASIC.pipeline" requires a number of arguments. We list these arguments in Table 1. For details users are recommended to read our manual.

Table 1: Arguments for the "MBASIC.pipeline" function.						
	Data Sources					
chipfile	A string vector for the ChIP files.					
inputfile	A string vector for the matching input files. The length must be					
	the same as "chipfile".					
input.suffix	A string for the suffix of input files. If NULL, "inputfile" will be					
	treated as the full names of the input files. Otherwise, all input files					
	with the initial "inputfile" and this suffix will be merged.					
chipformat	A string specifying the type of all ChIP (input) files, or a vector					
(input for-	of string specifying the types of each ChIP (input) file. Currently					
mat) three file types are allowed: "BAM", "BED" or "TAG						
	("TAGALIGN" and "BED" files are treated as same). Default:					
	"BAM".					
m.prefix	A string for the prefix of the mappability files.					
m.suffix	A string for the suffix of the mappability files. See our man files					
	for more details. Default: NULL.					
gc.prefix	A string for the prefix of the GC files.					
gc.suffix	A string for the suffix of the GC files. See our man files for more					
	details. Default: NULL.					
Genomic Information						
target	A GenomicRanges object for the target intervals where the reads					
	are mapped.					
fragLen	Either a single value or a 2-column matrix of the fragment lengths					
	for the chip and input files. Default: 150.					
pairedEnd	Either a boolean value or a 2-column boolean matrix for whether					
	each file is a paired-end data set. Currently this function only					
	allows "BAM" files for paired-end data. Default: FALSE.					
unique	A boolean value for whether only reads with distinct genomic					
coordinates or strands are mapped. Default: TRUE.						
_	Model Parameters					
fac	A vector of length N for the experimental condition of each ChIP					
	replicate.					
struct	A matrix indicating the levels of the signal matrix.					
J	The number of clusters to be identified.					
family	The distribution of family to be used. Either "lognormal" or "neg-					
	bin". See our man files for more information.					
nsig	The number of mixture components for the distribution of the					
	signal state. Default: 2.					
	Tuning Parameters					
burnin	An integer value for the number of iterations in initialization. De-					
•,	fault: 20.					
maxitr	The maximum number of iterations in the E-M algorithm. De-					
4-1	fault: 100.					
tol	Tolerance for error in checking the E-M algorithm's convergence.					
dotof1-	Default: 1e-04.					
datafile	The location to save the count matrices.					

2.4 Stepwise Execution

Alternatively, each step in the pipeline can be executed separately.

```
> ## Step 2: Generate mapped count matrices
> dat <- generateReadMatrices(chipfile = tbl$chipfile,</pre>
                               inputfile = tbl$inputfile,
                               input.suffix = ".bed",
                               target = target,
                               chipformat = tbl$chipformat,
                               inputformat = tbl$inputformat,
                               fragLen = 150,
                               pairedEnd = FALSE,
                               unique = TRUE)
> ## Step 3: Compute M and GC scores
> target <- averageMGC(target = target,
                       m.prefix = "syntheticData/mgc/",
                       m.suffix = "_M.txt",
+
                       gc.prefix = "syntheticData/mgc/",
+
                       gc.suffix = "_GC.txt")
> ## Step 4: Compute the background means
> MuO <- bkng_mean(inputdat = dat$input,
                   target = target,
                   family = "negbin")
> ## Step 5: Fit an MBASIC model
> MBASIC.fit <- MBASIC.binary(Y = t(dat$chip),
                               Mu0 = t(Mu0),
+
                               fac = conds,
                               J=3,
+
                               zeta=0.2,
                               maxitr = 100,
                               burnin = 20,
                               outfile=NULL,
                               out=NULL,
                               init.mod = NULL,
                               struct = NULL,
                               family="negbin",
                               tol = 1e-4,
                               nsig = 2)
```

2.5 The "MBASICFit" Class

> showClass("MBASICFit")

The outputs of both "MBASIC.binary" and "MBASIC.pipeline" functions are of S-4 class "MBASICFit".

```
Class "MBASICFit" [package "MBASIC"]
Slots:
Name:
                   Theta
                                          W
Class:
                  matrix
                                     matrix
                        V
                                          Z
Name:
Class:
                  matrix
                                     matrix
Name:
                        b
                                        aic
Class:
                 numeric
                                    numeric
Name:
                     bic
                                       aicc
```

numeric	numeric	Class:
alllik	lik	Name:
numeric	numeric	Class:
Mu	zeta	Name:
matrix	numeric	Class:
sigma0	Sigma	Name:
numeric	matrix	Class:
probz	e	Name:
numeric	numeric	Class:
converged	P	Name:
logical	matrix	Class:
ARI	Theta.err	Name:
numeric	numeric	Class:
MisClassRate	W.err	Name:
numeric	numeric	Class:
AssociationMatrix matrix	Iter numeric	Name: Class:
Struct	Loss	Name:
matrix	list	Class:

Slot "Theta" is a matrix for the estimated state, where each row corresponds to an experimental condition, and each column corresponds to a locus. Each entry is the probability for the locus to be un-binding at the corresponding state.

```
> dim(MBASIC.fit@Theta)
[1] 10 300
> rownames(MBASIC.fit@Theta)
  [1] "Cell1.Fac1" "Cell1.Fac2" "Cell1.Fac3"
  [4] "Cell1.Fac4" "Cell1.Fac5" "Cell2.Fac1"
  [7] "Cell2.Fac2" "Cell2.Fac3" "Cell2.Fac4"
[10] "Cell2.Fac5"
> head(MBASIC.fit@Theta[ 1, ])
[1] 1 1 1 1 1 1
```

Slot "Z" is a matrix for the posterior probablity of each locus to belong to each cluster. The rows correspond to the loci and the columns correspond to the clusters.

Slot "b" is a vector for the probability of each locus not to belong to any cluster.

```
> length(MBASIC.fit@b)
[1] 100
```

```
> head(MBASIC.fit@b)
[1] 0.17699181 0.15474780 0.47290380 0.44647337
[5] 1.000000000 0.07785736
```

Slot "W" is a matrix for the probability of loci in each group to be unenriched at each condition.

```
> rownames(MBASIC.fit@W)
 [1] "Cell1.Fac1" "Cell1.Fac2" "Cell1.Fac3"
 [4] "Cell1.Fac4" "Cell1.Fac5" "Cell2.Fac1"
 [7] "Cell2.Fac2" "Cell2.Fac3" "Cell2.Fac4"
[10] "Cell2.Fac5"
> dim(MBASIC.fit@W)
[1] 10 3
> head(MBASIC.fit@W)
                                [,2]
                                              [,3]
                   [,1]
Cell1.Fac1 1.000000e+00 0.999999999 0.9999999999
Cell1.Fac2 2.824951e-01 0.0012727096 0.9999999999
Cell1.Fac3 5.997807e-01 0.0000000001 0.6887011720
Cell1.Fac4 1.000000e+00 0.999999999 0.0000000001
Cell1.Fac5 9.267086e-05 0.5333470444 0.8000822904
Cell2.Fac1 1.000000e+00 0.5345144593 0.9191454521
```

2.6 Advanced Model Initialization

An important argument that is accessible by the "MBASIC.binary" function but not "MBASIC.pipeline" is "init.mod". This argument allows the user to pass a "MBASICFit" object, whose values are used to initialize the parameters in this function. It can be useful in two conditions:

- 1. The model fitted by a previous call of function "MBASIC.binary" has not yet converged, and the user wishes to increase the number of iterations;
- 2. The user wants to fit a MBASIC model with a larger number of clusters ("J") from a previously fitted model;
- 3. The user wants to fit a MBASIC model with a different structural constraints ("struct") from a previously fitted model.

In both cases, "MBASIC.binary" uses the results from "init.mod" before starting its iterations. This is significantly time saving than restarting the model fitting from scratch. However, this argument is dangerous if the user passes along a fitted model using different data sources or model structures. To avoid that, the user need to check that the arguments used to get "init.mod" and in the current "MBASIC.binary" must be consistent following rules below (which currently cannot be checked by our codes):

- 1. "Y", "Mu0", "fac", "family", "nsig" must be the same;
- 2. "J" for the "MBASIC.binary" function must be larger than or equal to the value used in "init.mod".

```
> ## Fit a MBASIC model with 4 clusters
> MBASIC.binary(Y = t(dat$chip), Mu0 = t(Mu0),
+ fac = conds, J=4, zeta=0.2,
+ maxitr = 100, burnin = 20,
+ init.mod = MBASIC.fit,
+ struct = NULL, family="negbin",
+ tol = 1e-4, nsig = 2)
> ## Fit a MBASIC model with more iterations
> MBASIC.binary(Y = t(dat$chip), Mu0 = t(Mu0),
+ fac = conds, J=3, zeta=0.2,
+ maxitr = 200, burnin = 20,
+ init.mod = MBASIC.fit,
+ struct = NULL, family="negbin",
+ tol = 1e-4, nsig = 2)
```

3 General Functionalities

The MBASIC package also provides additional functions to simulate and fit general MBASIC models with $S \geq 2$ states.

3.1 Simulation for General MBASIC Models

Function "MBASIC.sim" simulates data with "I" units with "J" clusters. The "S" argument specifies the number of different states, and "zeta" is the proportion of unclustered units. "fac" specifies the condition for each experiment. The "xi" argument relates to the magnitude of the simulated data. Its detailed description is included in our manual.

The "MBASIC.sim" function returns a list object. The "Y" field contains the simulated data matrix at each unit (column) for each experiment (row). The "Theta" field is the matrix for the states for each unit (column) and each experimental condition (column). The "W" field is a matrix with dimensions $KS \times J$, where the (S(k-1)+s,j)-th entry is the probability that units in the j-th cluster have state s under the k-th condition.

```
> names(dat.sim)
                   "Y"
                                 "W"
 [1] "Theta"
 [4] "Z"
                   "delta"
                                 "zeta"
 [7] "prior.mean"
                   "prior.sd"
                                 "stdev"
[10] "Mu"
                                 "snr"
                   "bkng"
[13] "non.id"
> dim(dat.sim$Y)
[1]
      20 1000
> dim(dat.sim$W)
[1] 30 3
> dim(dat.sim$Theta)
[1]
      10 1000
```

3.2 Fitting General MBASIC Models

The general MBASIC model is fitted by the function "MBASIC". The arguments here are similar to the "MBASIC.binary" function. The arguments that are not common to both functions are::

Argument	Meaning	"MBASIC.binary"	"MBASIC"
"nsig"	Number of signal components for the binding state.	Y	N
"Mu0"	Background means.	Y	N
"S"	Number of states.	N	Y
"method"	Fitting method.	N	Y
"para"	True parameters.	N	Y

The "method" argument specifies the fitting algorithm to be used. Currently, thre algorithms are provided. "em" is the nested E-M algorithm that is also used by "MBASIC.binary"; "2em" and "naive" are benchmark methods. Both "2em" and "naive" fit the model in two phases. In Phase 1 they estimate the state space matrix, and in Phase 2 "2em" uses a mixture model for clustering the units while "naive" uses hierarchical clustering.

The "para" argument allows users to pass the true model parameters into the model. In the following codes we pass the simulated list object "dat.sim". In that case, the output of the model contains a few slots that calculate the estimation error:

- ARI: Adjusted Rand Index;
- *W.err*: The mean squared error in matrix W;
- Theta.err: The mean squared error in state estimation;

• *MisClassRate*: The mis-classification rate.

For more details of the algorithms, as well as these metrics to assess model fitting, users may refer to [Zuo et al., 2014].

3.3 Degenerate MBASIC Models

In a degenerate MBASIC model, the states for each unit under each condition are directly observed. "MBASIC.sim.state" and "MBASIC.state" functions allows users to simulate and fit such models. The usage of these functions are similar to functions "MBASIC.sim" and "MBASIC".

"MBASIC.sim.state" simulates data from a degenerate MBASIC model. Most arguments are similar to "MBASIC.sim". "MBASIC.sim". "MBASIC.sim.state" does not need arguments "fac" and "family", but it needs the "K" argument, specifying the number of experimental conditions.

"MBASIC.state" fits a degenerate MBASIC model. Compared to function "MBASIC", it does not need arguments "Y" and "family". Instead, it needs the argument "Theta" to pass the observed states.

```
> state.sim.fit <- MBASIC.state(Theta = state.sim$Theta, J = 4,
+ method = "2em", zeta = 0.1,
+ maxitr = 100, tol = 1e-04)</pre>
```

4 Session Information

```
R version 3.1.1 (2014-07-10)
Platform: x86_64-redhat-linux-gnu (64-bit)
locale:
 [1] LC_CTYPE=zh_TW.UTF-8
 [2] LC_NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
 [4] LC_COLLATE=en_US.UTF-8
 [5] LC_MONETARY=en_US.UTF-8
 [6] LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
 [8] LC_NAME=C
 [9] LC_ADDRESS=C
[10] LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8
[12] LC_IDENTIFICATION=C
attached base packages:
[1] parallel stats
                        graphics grDevices
[5] utils
              datasets methods
                                  base
```

```
other attached packages:
```

```
[1] doMC_1.3.3
                          iterators_1.0.7
 [3] foreach_1.4.2
                          MBASIC_0.99.0
 [5] MASS_7.3-33
                          GenomicRanges_1.14.4
 [7] XVector_0.2.0
                          IRanges_1.20.7
 [9] BiocGenerics_0.8.0
                          mclust_4.3
[11] Rcpp_0.11.2
                          {\tt msm\_1.4}
[13] gtools_3.4.1
loaded via a namespace (and not attached):
 [1] codetools_0.2-8 compiler_3.1.1
```

- $[3] expm_0.99-1.1$ grid_3.1.1 [5] lattice_0.20-29 Matrix_1.1-4 [7] mvtnorm_1.0-0 splines_3.1.1 [9] stats4_3.1.1 survival_2.37-7
- [11] tools_3.1.1

References

[Zuo et al., 2014] Zuo, C., Hewitt, K., Bresnick, E., and Keles, S. (2014). A bayesian framework for state space matrix clustering. Submitted.