Multiple Beta t-Tests of Differential Transcription of mRNAs Between Two Conditions with Small Samples

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February 3, 2018

Abstract

A major task in the analysis of count data of RNA reads from RNA-Seq is the detection of differentially expressed genes or isoforms. The count data are presented as a matrix consisting of RNA isoform annotation and the number of reads. Analogous analyses also arise for other assay types, such as comparative ChIP-Seq. The package MBttest provides a powerful method to test for differential expression by use of the beta distribution and gene- or isoform-specific variable ρ to control fudge effect due to small sample size. 1 . This vignette explains the use of the package. For more detail of the statistical method, please see our paper [8].

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1 Introduction

This vignette is intended to give a rapid introduction to the commands used in implementing new beta t-test methods of evaluating differential expression in high-throughput sequencing data by means of the $MBttest\mathbf{R}$ package. For fuller details on the methods being used, consult Tan et al (2015) [8] .

We assume that we have count data from a set of sequencing or other high-throughput experiments, arranged in an array such that except gene annotation information and id, each column describes a library and each row describes RNA tag or isoform for which data have been acquired. For example, the rows may correspond to the different sequences observed in a sequencing experiment. The data then consists of the number of each sequence observed in each sample. We wish to determine which, if any, rows of the data correspond to some patterns of differential expression across the samples.

MBttest, also called *mBetat* test, uses new beta t-test method to identify differential expression for each row. This approach introduces a gene- or isoform-specific variable, called ρ , into t-statistics to control fudge effect resulted from

¹Other Bioconductor packages with similar aims are edgeR, baySeq, DESeq and DESeq2.

small samples. It has higher work efficiency than existing methods for identifying differential expressions of genes or isoforms either by inflating t-values with

$$\rho > \omega$$

threshold or by shrinking those with

$$\rho < \omega$$

[8] when number of replicate libraries in each condition is small, for example, equal to or less than 8.

Different from the exiting methods such as baySeq [2], edgeR Exact test [6] and [7], edgeR GLM [5] and [7], DESeq [1] and DESeq2 [4], etc, MBttest requires performance of simulation to determine threshold ω before running program mbetattest. MBttest provides negative binomial simulation program to generate null count data without inputting arguments. User should repeat five or more simulations, perform program smbettest to produce null results and calculate ω using the method given in our paper [8].

2 Data Preparation and Input

We begin by loading the MBttest package.

> library(MBttest)

MBttest requires data file contain two parts: Annotation information and count data. Information consists of tagid, geneid, gene name, chromosome id, DNA strand, etc. Information is in left side. It has at least one column for geneid or tagid (isoformid). The data contain two conditions each having several replicate libraries and must be in right side. Here is an example:

- > data(jkttcell)
- > jkttcell[1:10,]

	tagid	geneid	name	chr	strand	pos	anno	Jurk.NS.A	Jurk.NS.B	Jurk.NS.C		
1	54	58998	COMT	chr22	+	19956542	sg	66.80	43.48	4.65		
2	111	59029	CRKL	chr22	+	21308033	tu-ce	68.75	63.94	66.46		
3	171	59104	SLC2A11	chr22	+	24227723	sg	2.86	2.67	8.15		
4	231	59157	ADRBK2	chr22	+	26118985	tu	12.88	8.45	8.59		
5	242	59164	SRRD	chr22	+	26887904	sg	62.54	59.88	83.27		
6	265	59184	HSCB	chr22	+	29153206	tu	4.02	2.07	6.85		
7	306	59209	UCRC	chr22	+	30165939	sg	516.71	594.71	83.84		
8	327	59212	MTMR3	chr22	+	30426495	tu	4.08	3.99	2.93		
9	445	59310	NCF4	chr22	+	37274056	sg	0.00	1.30	0.03		
10	472	59321	CYTH4	chr22	+	37711384	sg	4.40	1.60	0.12		
	Jurk.48h.A Jurk.48h.B Jurk.48h.C											

1	32.99	25.49	14.68
2	80.42	63.89	72.48
3	12.95	12.70	8.81
4	13.13	35.34	9.78
5	54.99	51.15	66.61
6	9.66	4.02	5.87
7	254.81	142.26	156.72
8	10.38	14.23	6.85
9	71.26	25.89	11.73
10	37.61	18.54	16.64

User also may use head to display the data with top 6 lines:

> head(jkttcell)

	tagid	geneid	name	chr	strand	pos	anno	Jurk.NS.A	Jurk.NS.B	Jurk.NS.C
1	54	58998	COMT	chr22	+	19956542	sg	66.80	43.48	4.65
2	111	59029	CRKL	chr22	+	21308033	tu-ce	68.75	63.94	66.46
3	171	59104	SLC2A11	chr22	+	24227723	sg	2.86	2.67	8.15
4	231	59157	ADRBK2	chr22	+	26118985	tu	12.88	8.45	8.59
5	242	59164	SRRD	chr22	+	26887904	sg	62.54	59.88	83.27

6	265 5918	4 HSCB cl	nr22 +	29153206	tu	4.02	2.07	6.85
	Jurk.48h.A	Jurk.48h.B	Jurk.48h.C					
1	32.99	25.49	14.68					
2	80.42	63.89	72.48					
3	12.95	12.70	8.81					
4	13.13	35.34	9.78					
5	54.99	51.15	66.61					
6	9.66	4.02	5.87					

If the data are *csv* file, user can use read.csv function to input data into **R Console** or If the data are *txt* file, user can use read.delim function to load data into **R Console**. After loading data, user should check the data inputted. jkttcell shows an example. In this example, 7 columns in the left side are information of *poly(A)* sites. The count data are listed in the right side.

3 Simulation for Calculating omega Value

Before performing mbetattest on the real data, user needs simulation to determine omega value. There are three steps for doing so:

3.1 Step1: Simulate null count data

Use the following function to generate null simulation data

where

yy is real data.

r1 and r2 are replicate numbers in conditions 1 and 2.

p is proportion of genes differentially expressed in m genes, default value is 0.

q is proportion of genes artificial noise. Its default value is 0.

A is effect value. Its default value is 0. nci: column number of information of data.

Here is an example:

```
> sjknull1<-simulat(yy=jkttcell[1:500,],nci=7,r1=3,r2=3,q=0.2)
```

Our data is *jkttcell*. It has 7 columns for information of poly(A) sites. Two conditions are resting and stimulation. Each has 3 replicate libraries, r1=3 and r2=3. Since this is null simulation, we set p=0 and q=0.2 for artificial noise. With the same read data and parameters, you can generate a set of 4-6 null datasets: sjknull2, ..., sjknull6 for calculating omega value.

3.2 Step2: Perform multiple beta t-tests

Use function smbetattest to perform multiple beta t-test with ho=1 on the simulated null data:

where

X=simulated data.

na and nb are numbers of replicate libraries in conditions 1 and 2.

alpha is probabilistic threshold. User can set alpha=0.05 or 0.01.

The example is

> mysim1<-smbetattest(X=sjknull1,na=3,nb=3,alpha=0.05)

Save them to csv files

```
write.csv(mysim1, file='simulatedNullData1Result.csv")
```

After performing smbetattest on each simulated null dataset, user would have results recorded in file like *simulatedNullData1Result.csv* and open it with excel.

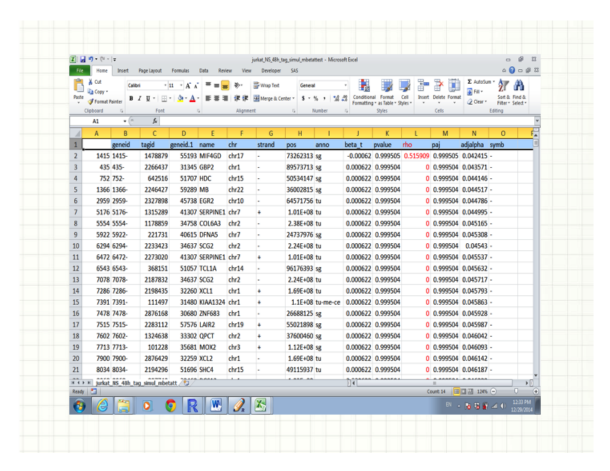


Figure 1: This is an example showing the results obtained by smbetattest from simulated null data. Column A is row number, geneid is set in simulation, and geneid.1 is original geneid.

In symbol column, mbeta t-test gives test result: symb = "-" means that the gene or tag is not chosen while symb = "+" indicates that the gene or isoform is found to be differentially expressed.

In this example, 12 genes would be found to be falsely positive.

3.3 Step3: Calculate omega value

Here is a demo for calculating omega (since we can't use greek letter omega in R function, we use W to represent omega). In Figure 3, red highlighted column is rho column. We copied the ρ values of these 12 genes into another empty column and sorted them from the smallest to the largest. Then we gave sequence numbers from 1 to 12 corresponding to ρ -values and calculate q-value for each ordered ρ value:

We chose the 10th ρ value (1.09166) as the first W value because the 11th rho value has q-value > 0.85. Repeat this process in 4-6 simulated null datasets and we took the averaged W value as W value in the real data.

4 Normalize the count data

As a second processing step, we need to estimate the effective library size. This step is also called *normalization*, even though it may not make the count data be of normal distribution. If the counts of expressed genes in one condition are, on average, twice as high as in another (because the library was sequenced twice as deeply), the size factor for the first condition should be twice higher than the second one, then differential analysis would give error results. For this reason, we must make all libraries have the same size before performing any statistical method. For doing so, user can the function estimateSizeFactors of **R** package *DESeq* [1] or *DESeq2* [4] to estimate the size factors from the count data or use the following simple method to normalize the the count data: In excel sheet, use function

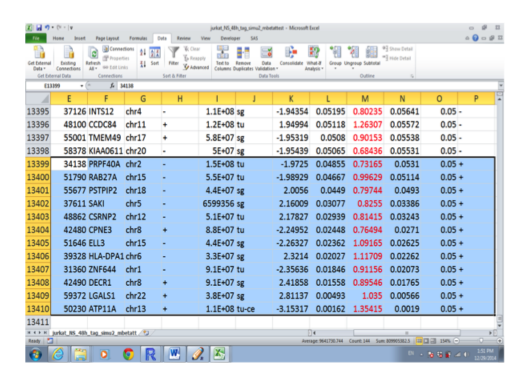


Figure 2: The row number in column A in Figure 1 was deleted. To show explicitly, we here hided geneid (for simulation), tagid column and selected rows are genes that were identified to be differentially expressed.

sum to calculate sizes of all libraries, and then use excel function average to calculate averaged library size. The last step is to use the following equation to convert the original count data to new count data with the same library size:

$$Y_{ik} = \frac{y_{ij}\bar{N}}{N_i}$$

where $i=2,\,\cdots,\,n$ (number of genes or isofoms) in rows in a sheet, $j=nci+1,\,\cdots,\,nci+c$ where c=na+nb and nci is column number of annotation information; $k=nci+c+2+j;\,N_j$ is size of library j and \bar{N} is mean of sizes over all libraries; y_{ij} is original count of RNA reads in row i and column j.

5 Perform Multiple Beta *t*-Tests on The Real Data

Suppose the data have been normalized so that all libraries have the same size. After obtaining W value, user can use the function and the real data to perform mbeta t-test: mbetattest(X, na, nb, W, alpha, file) where X is real data. In our current example, X=jkttcell. na and nb are respectively numbers of replicate libraries in conditions 1 and 2. For jktcell data, na = nb = 3. W is omega value. According to our calculation above step, W = 1. alpha is the probabilistic threshold. You can set alpha=0.05 or 0.01 or the other values; file is csv file for saving the results. The example is

> res<-mbetattest(X=jkttcell[1:1000,],na=3,nb=3,W=1,alpha=0.05,file="jurkat_NS_48h_tag_mbetattest.csv")

mbetattest has two output results: one is saved in csv file and the other is dat for maplet and for heatman. The

mbetattest has two output results: one is saved in *csv* file and the other is dat for *maplot* and for *heatmap*. The package MBttest has this result obtained the whole data. We here load it for making MAplot:

- > data(dat)
- > head(dat)

Image: Section of the control of												
Paste	Copy *		11 · A A	=	₩rap Tex			Conditional Fo		Insert Delete Form	Σ AutoSum *	ZI III
Clipb	Format Painter oard 5	Font	6	All	gnment		Number G	Formatting * as T Style		Cells	② Clear ▼ Edi	Filter * Select * ting
Q13398 • (* fe rho												
	Н	1	J	K	L	M	N	0	Р	Q	R	S
13395	-	1.1E+08	sg	-1.94354	0.05195	0.80235	0.05641	0.05	-			
13396	+	1.2E+08	tu	1.94994	0.05118	1.26307	0.05572	0.05	-			
13397	+	5.8E+07	sg	-1.95319	0.0508	0.90153	0.05538	0.05	-			
13398	-	5E+07	sg	-1.95439	0.05065	0.68436	0.05531	0.05	-	rho	sequence	q-value
13399	-	1.5E+08	tu	-1.9725	0.04855	0.73165	0.0531	0.05	+	0.73165	1	0.08333
13400	-	5.5E+07	tu	-1.98929	0.04667	0.99629	0.05114	0.05	+	0.76494	2	0.16667
13401	-	4.4E+07	sg	2.0056	0.0449	0.79744	0.0493	0.05	+	0.79744	3	0.25
13402	-	6599356	sg	2.16009	0.03077	0.8255	0.03386	0.05	+	0.81415	4	0.33333
13403	-	5.1E+07	tu	2.17827	0.02939	0.81415	0.03243	0.05	+	0.8255	5	0.41667
13404	+	8.8E+07	tu	-2.24952	0.02448	0.76494	0.0271	0.05	+	0.89546	6	0.5
13405	-	4.4E+07	sg	-2.26327	0.02362	1.09165	0.02625	0.05	+	0.91156	7	0.58333
13406	-	3.3E+07	sg	2.3214	0.02027	1.11709	0.02262	0.05	+	0.99629	8	0.66667
13407	-	9.1E+07	tu	-2.35636	0.01846	0.91156	0.02073	0.05	+	1.035	9	0.75
13408	+	9.1E+07	sg	2.41858	0.01558	0.89546	0.01765	0.05	+	1.09165	10	0.83333
13409	+	3.8E+07	sg	2.81137	0.00493	1.035	0.00566	0.05	+	1.11709	11	0.91667
13410	+	1.1E+08	tu-ce	-3.15317	0.00162	1.35415	0.0019	0.05	+	1.35415	12	1
13411												
Ready 2		h_tag_simu2_mb	etatt 😲					erage: 2.66208044		Sum: 95.83489585	II)

Figure 3: Demo for calculating W

	tagid	geneid	name	chr	strand	pos	${\tt anno}$	Jurk.NS.A	Jurk.NS.B	Jurk.NS.C	Jurk.48h.A
1	83344	58782	MX1	chr21	+	42831139	sg	0	0.00	0.00	29.61
2	197313	56792	CD22	chr19	+	35838262	sg	0	0.45	3.81	96.33
3	202264	53072	CD19	chr16	+	28950664	sg	0	0.00	0.00	37.65
4	232007	37653	BASP1	chr5	+	17276943	sg	0	0.55	1.31	63.15
5	301820	46661	HBB	chr11	_	5246697	sg	0	0.00	0.00	4.52
6	368151	51057	TCL1A	chr14	_	96176393	sg	0	1.03	5.15	153.51
	Jurk.48	Bh.B Jui	ck.48h	.C beta	a_t rho	symb					
1	(0.48		0	0 0	_					
2	8	3.07		0	0 0	_					
3	().52		0	0 0	_					
4	().58		0	0 0	_					
5	39	9.52		0	0 0	-					
6	Ş	9.93		0	0 0	-					

> maplot(dat=dat,r1=3,r2=3,TT=350,matitle="MA plot")

myheatmap has multiple options: both-side, row and column cluster trees with distance methods: "euclidean", "pear-son", "spearman", and "kendall" correlation coefficients and color label with "redgreen", "greenred", "redblue", "bluered" or "heat.colors" and angles for genes or isoforms in row and cases (conditions) in column. User can use default without any choice like (Figure 6) which has tree="both" for both-side tree, or choose tree="column" like (Figure 8) if columns are species or cancer cases or not choose tree with tree="none" (see (Figure 7)). User may change heatmap color with colors, for example, in (Figure 8), we chose colors="redblue". If user find that default column name or row name does no have good angles, then user can adjust them with rwangle (row angle) or clangle(column angle). rwangle and clangle values are from 0 to 90.

> myheatmap(dat=dat,r1=3,r2=3,maptitle="Jurkat T-cell heatmap2")

> maplot(dat=dat,r1=3,r2=3,TT=50,matitle="MA plot")

MA plot

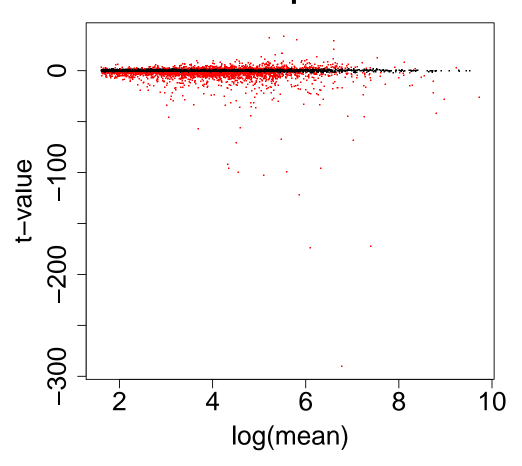


Figure 4: 'MA'-plot of t-value against log mean over all replicate libraries across two conditions. The isoforms who were given differential transcripts in simulation had absolute larger t-values that were highlighted in red than the threshold given in multiple tests. Those who were given no differential expression had very small absolute t-values close to zero labeled in black across long means. Here threshold for truncating t-values is set to be 350, since none of absolute t-values are over 350, the t-Maplot is an outline t-Maplot in which red and black dots are not explicitly seen.

- > myheatmap(dat=dat,r1=3,r2=3,tree="none",maptitle="Jurkat T-cell heatmap3")
- > myheatmap(dat=dat,r1=3,r2=3,colrs="redblue", tree="column",
- + method="pearson", maptitle="Jurkat T-cell heatmap")

Figure 5: 'MA'-plot of t-value against log mean over all replicate libraries across two conditions. To explicit display the t-values across log mean, absolute t-values >= 50 were truncated. One can explicitly see that truly differential transcripts in simulation had absolute larger t-values that were highlighted in red than the threshold given in multiple tests and those who had no differential expressions had very smaller absolute t-values that were labeled in black than the threshold.

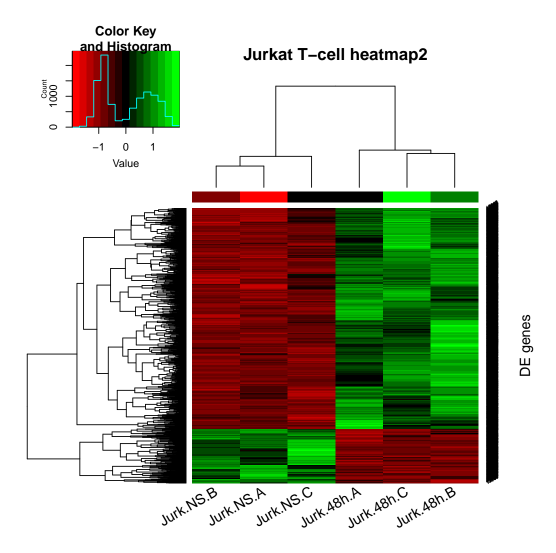


Figure 6: The *heatmap* has both-side trees and displays explicitly differential expression between stimulating and rest. Most of genes were up-expressed by stimulation but a small part of genes were down-expressed. The tree in column divides columns into two groups: NS and 48h. The tree in row is tree of differentially expressed genes and also divide genes in row into two big groups.

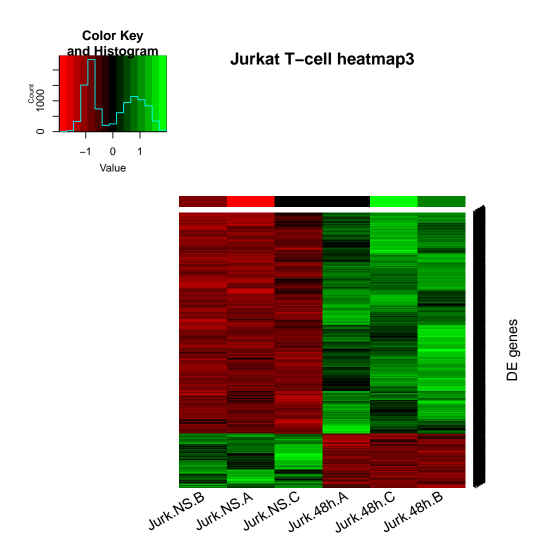


Figure 7: The heatmap did not give trees on both sides, but the heatmap is the same with (Figure 6)

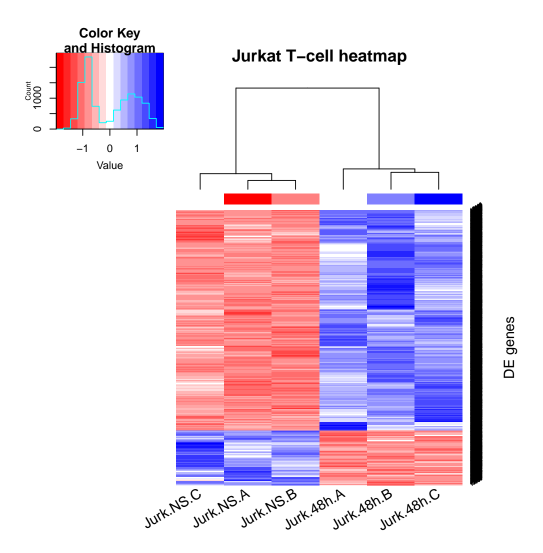


Figure 8: This *heatmap* was labeled with red and blue and gave column trees

6 Session Info

```
> sessionInfo()
R version 3.3.1 (2016-06-21)
Platform: x86_64-apple-darwin11.0.0 (64-bit)
Running under: OS X 10.11.6 (El Capitan)
locale:
[1] C/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
attached base packages:
[1] stats
              graphics grDevices utils
                                            datasets methods
                                                                 base
other attached packages:
[1] MBttest_1.7.1 gtools_3.5.0 gplots_3.0.1
loaded via a namespace (and not attached):
[1] BiocStyle_2.2.1
                       tools_3.3.1
                                          KernSmooth_2.23-15 gdata_2.18.0
[5] caTools_1.17.1
                       bitops_1.0-6
```

References

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