RnaSeqSampleSize: Sample size estimation based on real RNA-seq data

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Abstract

In this vignette, we demonstrated the application of RnaSeqSampleSize as an sample size estimation tool for RNA-seq data. A user friendly web interface is also provided at http://cqs.mc.vanderbilt.edu/shiny/RnaSeqSampleSize/ for researchers not familar with R.

RnaSeqSampleSize package provides the following features:

- Estimation of sample size or power by single read count and dispersion;
- Estimation of sample size or power by prior real data;
- Visualization of sample size and power by power curves;
- Optimization by power or sample size matrix;

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1 Introduction

Sample size estimation is the most important issue in the design of RNA sequencing experiments. However, thousands of genes are quantified and tested for differential expression simultaneously in RNA-seq experiments. The false discovery rate for statistic tests should be controlled. At the same time, the thousands of genes have widely distributed read counts and dispersions, which were often estimated by experience or set at the most conservative values in previous sample size estimation methods. As a result, the estimated sample size will be inaccurate or over-estimated.

To solve these issues, we developed a sample size estimation method based on the distributions of gene read counts and dispersions from real data. Datasets from the user's preliminary experiments or the Cancer Genome Atlas (TCGA) can be used as reference. The read counts and their related dispersions will be selected randomly from the reference based on their distributions, and from that, the power and sample size will be estimated and summarized.

2 User friendly web interface

A user friendly web interface for *RnaSeqSampleSize* package is provided at http: //cqs.mc.vanderbilt.edu/shiny/RnaSeqSampleSize/. Most of the functions in Examples section can be performed in this website.

3 Examples

First we will load the *RnaSeqSampleSize* package.

```
library(RnaSeqSampleSize)
```

Loading required package: RnaSeqSampleSizeData

Loading required package: edgeR

Loading required package: limma

3.1 Estimation of sample size or power by single read count and dispersion

3.1.1 Power estimation

For example, if we are estimating the power of finding significant genes for RNAseq data with specified sample size, and we have the following parameters:

- Number of samples in each group: 63;
- Minimal fold change between two groups: 2;
- Minimal average read counts: 5;
- Maximal dispersion: 0.5;
- False discovery rate: 0.01;

As a result, the estimated power is 0.8 by est_power function. It means that we have 80% probability to find the significant genes with 63 samples in each group.

```
example(est_power)
##
## est_pw> n<-63;rho<-2;lambda0<-5;phi0<-0.5;f<-0.01
##
## est_pw> est_power(n=n, rho=rho, lambda0=lambda0, phi0=phi0,f=f)
## [1] 0.8
```

3.1.2 Sample size estimation

For example, if we are estimating the sample size for RNA-seq data to achieve desired power of finding significant genes, and we have the following parameters:

- Desired power of finding significant genes: 0.8;
- Minimal fold change between two groups: 2;
- Minimal average read counts: 5;
- Maximal dispersion: 0.5;
- False discovery rate: 0.01;

As a result, the estimated sample size is 63 by sample_size function. It means that if we want to have 80% probability to find the significant genes, we need 63 samples in each group.

```
example(sample_size)
##
## smpl_s> power<-0.8;rho<-2;lambda0<-5;phi0<-0.5;f<-0.01
##
## smpl_s> sample_size(power=power, f=f,rho=rho, lambda0=lambda0, phi0=phi0)
## [1] 63
```

3.2 Estimation of sample size or power by reference data

3.2.1 Power estimation with datasets in RnaSeqSampleSizeData package

RnaSeqSampleSizeData package contains the read counts and dispersion distribution from some real datasets and can be used as prior data for sample size or power estimation. They can be called with following names:

```
## [1] "TCGA_BLCA" "TCGA_BRCA" "TCGA_CESC" "TCGA_COAD" "TCGA_HNSC" "TCGA_KIRC"
## [7] "TCGA_LGG" "TCGA_LUAD" "TCGA_LUSC" "TCGA_PRAD" "TCGA_READ" "TCGA_THCA"
## [13] "TCGA_UCEC"
```

For example, if we are estimating the power of finding significant genes for RNAseq data with specified sample size, and we have the following parameters:

- Number of samples in each group: 65;
- Minimal fold change between two groups: 2;
- Prior data: TCGA READ data, stored in *RnaSeqSampleSizeData* package, can be used with name TCGA_READ;
- False discovery rate: 0.01;

Here we demonstrated the power estimation by prior data in three different situations.

 If we are intesested in all genes, we can use repNumber parameter to specify random number of genes to perform power estimation;

```
## [1] 0.9759823
```

Please note here the parameter repNumber was very small (5) to make the example code faster. We suggest repNumber should be at least set as 100 in real analysis.

 If we are only intesested in a list of genes, we can use selectedGenes parameter to specify the list of genes to perform power estimation;

```
#Power estimation based on some interested genes.
#We use storeProcess=TRUE to return the details for all selected genes.
selectedGenes<-names(TCGA_READ$pseudo.counts.mean)[c(1,3,5,7,9,12:30)]</pre>
powerDistribution<-est_power_distribution(n=65,f=0.01,rho=2,</pre>
                        distributionObject="TCGA_READ",
                        selectedGenes=selectedGenes,
                        storeProcess=TRUE)
str(powerDistribution)
## List of 3
## $ power
              : num [1:24] 0.8161 0.0547 0.9781 0.9785 0.979 ...
## $ count
              : Named num [1:24] 43 36 2000 260 1950 ...
## ..- attr(*, "names")= chr [1:24] "A1BG" "A2BP1" "A2M" "A4GALT" ...
## $ dispersion: num [1:24] 0.6 2.3 0.4 0.5 0.1 1.3 0.4 0.1 0.4 0.1 ...
mean(powerDistribution$power)
## [1] 0.8098202
```

 If we are only intesested a specified pathway, we can use pathway and species parameters to specify the genes in a pathway to perform power estimation.

As a result, we use <u>est_power_distribution</u> function and find the estimated power is 0.91 for random genes, 0.81 for specified gene list, and 0.77 for genes in Glycolysis and Gluconeogenesis (pathway 00010) pathway.

3.2.2 Sample size estimation with datasets in RnaSeqSampleSize-Data package

For example, if we are estimating the sample size for RNA-seq data to achieve desired power of finding significant genes, and we have the following parameters:

- Desired power of finding significant genes: 0.8;
- Minimal fold change between two groups: 2;
- Prior data: TCGA READ data, stored in *RnaSeqSampleSizeData* package, can be used with name TCGA_READ;
- False discovery rate (FDR): 0.01;

As a result, we use *sample_size_distribution* function and find the estimated sample size is 41 for random genes.

```
## [1] "x= 1 f(x)= -0.8"
## [1] "x= 33 f(x)= -0.00967581809745977"
## [1] "x= 65 f(x)= 0.173627660386663"
## [1] "x= 49 f(x)= 0.116489870766909"
## [1] "x= 41 f(x)= 0.0625692432067934"
## [1] "x= 37 f(x)= 0.028503205303583"
## [1] "x= 35 f(x)= 0.00992054428479727"
## [1] "x= 34 f(x)= 0.000216707014488948"
## [1] 34
```

Please note here the parameter repNumber was very small (5) to make the example code faster. We suggest repNumber should be at least set as 100 in real analysis.

3.2.3 Sample size or power estimation with user's prior dataset

For example, if the user has a RNA-seq data with 10000 genes and 10 samples as prior dataset:

```
# Generate a 10000*10 RNA-seq data as prior dataset
set.seed(123)
dataMatrix <- matrix(sample(0:3000, 1e+05, replace = TRUE), nrow = 10000, ncol = 10)
colnames(dataMatrix) <- c(paste0("Control", 1:5), paste0("Treatment", 1:5))
row.names(dataMatrix) <- paste0("gene", 1:10000)</pre>
```

```
head(dataMatrix)
```

##	Control1 Co	ontrol2 Co	ntrol3	Control4	Control5	Treatment1	Treatment2
## gene1	2462	56	421	219	2344	1636	1486
## gene2	2510	1353	1606	591	251	538	2230
## gene3	2226	2680	269	2668	170	2976	625
## gene4	525	461	1948	1589	1315	1580	1915
## gene5	194	2048	259	1761	2564	1862	2498
## gene6	2985	2678	2364	2111	704	2629	1852
##	Treatment3	Treatment	4 Treat	tment5			
## gene1	1961	192	4	1412			
## gene2	382	276	4	1491			
## gene3	1685	210	0	1248			
## gene4	1280	22	8	441			
## gene5	2947	109	3	1761			
## gene6	353	78	3	1097			

Then we are estimating the power of finding significant genes for RNA-seq data with specified sample size, and we have the following parameters:

- Number of samples in each group: 65;
- Minimal fold change between two groups: 2;
- Prior data: User's prior dataset with 10000 genes and 10 samples;
- False discovery rate: 0.01;

We will use est_count_dispersion to estitamete the gene read count and dispersion distribution of user's prior dataset. And then est_power_distribution function will be used to estimate power.

Disp = 0.603 , BCV = 0.7765

distributionObject=dataMatrixDistribution,repNumber=5)

[1] 0.7933217

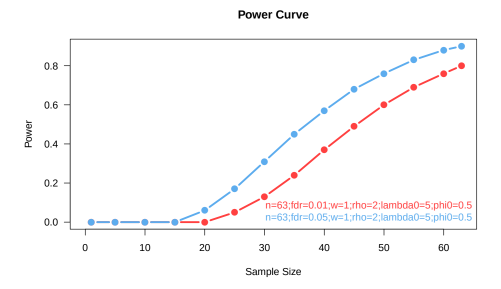
As a result, we can find the estimated power is 0.91. Please note here the parameter repNumber was very small (5) to make the example code faster. We suggest repNumber should be at least set as 100 in real analysis.

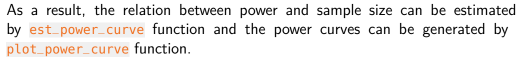
3.3 Power curve visualization

For example, if we are going to compare the power of finding significant genes for different false discovery rate, and we have the following parameters:

- Number of samples in each group: 63;
- Minimal fold change between two groups: 2;
- Minimal average read counts: 5;
- Maximal dispersion: 0.5;
- False discovery rate: 0.01 and 0.05;

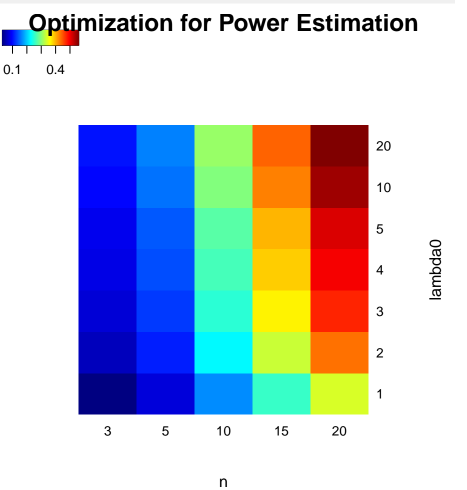
```
result1 <- est_power_curve(n = 63, f = 0.01, rho = 2, lambda0 = 5, phi0 = 0.5)
result2 <- est_power_curve(n = 63, f = 0.05, rho = 2, lambda0 = 5, phi0 = 0.5)
plot_power_curve(list(result1, result2))</pre>
```





3.4 Optimization by power or sample size matrix

For example, if the budget is limited, we need to balance the number of replications and sequence depth. We can use the **optimize_parameter** function to find the relation between sample size, read counts, and estimated power. And then the optimized parameters can be determined.



As a result, the estimated power distribution indicates that the number of replications plays a more significant role in determining the power than the number of read counts.