# intSEQ: Differential Analysis of RNA-seq Data with Integrated Likelihood Method

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#### 1 Introduction

This vignette is intended to give a brief introduction how to conduct differential analysis with RNA-seq data by means of the intSEQ R package. For further details of the methods, please consult [first paper], [second paper].

Consider the case we have RNA-seq count data arranged in a matrix that each columns describe a sample, and each row describe a genomic event (gene/exon/isoform). We denote the count in ith row and of jth group as  $Y_{ij}$ . Assume we have a predictor variable X, can be treatment/control group, dose of radiation, etc, which may or may or affect the level of expression across samples. We want to determine if it exist any pattern between the predictor X and different samples in each row.

The negative binomial distribution can be seen as Poisson-gamma mixture. If  $Y \sim \text{Poisson}(\lambda)$ , and  $\lambda$  is a random variable following Gamma distribution with shape parameter  $1/\theta$  and rate  $\frac{1}{\mu\theta}$ . Then Y has negative binomial distribution with mean parameter  $\mu = E(Y)$  and dispersion parameter  $\theta$ . The variance function of negative binomial distribution is

$$V(\mu) = \mu + \theta \mu^2$$

Many exsiting methods models RNA-seq data with negative binomial model. However, many of them suffers from false postive problem, that is, there are more than expected rejections occur. [first paper] suggests it may due to unaccounted variability of estimated dispersion parameters. The integrated likelihood method integrate the likelihood over the support of dispersion parameter rather than using a point estimator of the dispersion to control the false positive. It seems outperform other methods that also controls the false postive in term of higher power when the sample size is moderated.

# 2 Prepare the data

#### 2.1 Read the data

The users should arrange data into a numeric matrix with rows corresponds to genomic event (such as genes), and columns corresponds to samples. We also accept object of class "DGEList" in edgeR.

We use the Montgomery and Pickrell data as an example. The data set is RNA-Seq of RNA from lymphoblastoid cell lines from 129 individuals from the HapMap project. There were 60 from the above-referenced CEU subset and 69 from the Yoruba people in Ibadan, Nigeria (YRI). 52,580 unique genes of which 8,124 had a count of at least 129 across the 129 samples. We choose 10 CEU and 10 YRI individuals to conduct differential analysis.

```
library(intSEQ)
data("count.data")
data("condition")
```

```
count = count.data[ , c(1:10, 61:70)]
cond = condition[c(1:10, 61:70)]
count[1:10,]
##
                   NAO6985 NAO6986 NAO6994 NAO7000 NAO7037 NAO7051 NAO7346 NAO7347 NAO7357
                                                                 24
                                        10
                                                                         14
                                                                                 19
## ENSG00000000419
                        11
                                16
                                                 19
                                                          8
                                                                                          20
## ENSG0000000457
                        28
                                22
                                        17
                                                 21
                                                         18
                                                                 15
                                                                         21
                                                                                 13
                                                                                          18
## ENSG0000000460
                         0
                                 7
                                         2
                                                 4
                                                          6
                                                                  3
                                                                          8
                                                                                  5
                                                                                          2
                                                         17
                                                                  4
                                                                         19
                                                                                 35
                                                                                          16
## ENSG0000000938
                         8
                                28
                                        20
                                                 10
## ENSG0000001036
                        19
                                25
                                        13
                                                 22
                                                         10
                                                                 16
                                                                         12
                                                                                 12
                                                                                          23
                                                95
                                                         75
## ENSG0000001167
                               107
                                        89
                                                                         80
                                                                                 73
                                                                                         82
                       112
                                                                 87
## ENSG0000001497
                        10
                                 9
                                        13
                                                 4
                                                         18
                                                                 10
                                                                         10
                                                                                  7
                                                                                          4
                                                                                         108
## ENSG0000001561
                        83
                                62
                                        52
                                                 69
                                                         18
                                                                 44
                                                                         30
                                                                                 50
## ENSG0000002016
                         1
                                 1
                                         1
                                                 1
                                                          0
                                                                  0
                                                                          0
                                                                                  0
                                                                                           0
## ENSG0000002330
                        69
                                82
                                        51
                                                 69
                                                        100
                                                                 88
                                                                         34
                                                                                 47
                                                                                          65
                   NA10847 NA18486 NA18498 NA18499 NA18501
                                                            NA18502 NA18504
                                                                            NA18505 NA18507
##
                                22
                         6
                                       105
                                                 40
                                                         55
                                                                 67
                                                                         37
                                                                                 88
                                                                                         127
## ENSG0000000419
## ENSG0000000457
                        15
                                22
                                       100
                                                107
                                                         53
                                                                 72
                                                                         38
                                                                                 98
                                                                                         69
                         2
## ENSG0000000460
                                 5
                                        23
                                                 10
                                                         18
                                                                 15
                                                                          8
                                                                                 11
                                                                                          16
## ENSG0000000938
                         9
                                36
                                        70
                                                 41
                                                         33
                                                                 59
                                                                         29
                                                                                 22
                                                                                         71
                         4
                                29
                                        79
                                                33
                                                         31
                                                                 29
                                                                         21
## ENSG0000001036
                                                                                 42
                                                                                         62
## ENSG0000001167
                       108
                               301
                                       351
                                                344
                                                        176
                                                                340
                                                                        170
                                                                                238
                                                                                         247
                                32
## ENSG0000001497
                        10
                                        22
                                                 4
                                                          8
                                                                 15
                                                                          5
                                                                                 15
                                                                                         16
## ENSG0000001561
                        31
                                52
                                       419
                                                173
                                                        137
                                                                127
                                                                         72
                                                                                267
                                                                                         247
## ENSG0000002016
                         1
                                 3
                                         8
                                                 6
                                                          6
                                                                  5
                                                                          4
                                                                                  3
                                                                                          4
## ENSG00000002330
                        41
                                48
                                        81
                                                65
                                                         19
                                                                 59
                                                                         57
                                                                                 44
                                                                                         79
                   NA18508 NA18510
## ENSG0000000419
                        70
                                43
## ENSG0000000457
                        66
                                43
                                 7
## ENSG0000000460
                        18
## ENSG0000000938
                        12
                                43
                                25
## ENSG0000001036
                        41
                               227
## ENSG0000001167
                       226
## ENSG0000001497
                        19
                                15
## ENSG0000001561
                       154
                                81
## ENSG0000002016
                         6
                                 3
## ENSG00000002330
                        71
                                79
cond
    ## Levels: CEU YRI
```

## 2.2 Filtering

Since almost all existing methods have very low power for low expressed gene. We recommend filter out those genes with small average row count.

```
count = count[rowMeans(count) >= 1]
```

## 3 intSEQ Approach

#### 3.1 Normalization

The intSEQ function has a built-in normalization method "TMM". The users can use this method by setting "normalize=TRUE". Please don't normalize the count data yourself. Any non-integer input of count data will cause error.

## 3.2 Integrated Likelihood Ratio Test

The intSEQ function will estimate the Cox and Reid's estimator of dispersion, then fit the negative binomial model. After that, the likelihood will be integrated over the dispersion:

$$IL(\mu) = L(Y|\mu) = \int L(Y|\mu, \theta) \pi(\theta|\mu) d\theta$$

Then the integrated likelihood will be treated as likelihood function. The integrated likelihood ratio statistics is compared with a  $\chi^2$  distribution.

```
res=intSEQ(count, cond)
```

Alternatively, we can use DGEList

```
library(edgeR)
dge=DGEList(counts=count, group=cond)
res2=intSEQ(dge)
```

### 3.3 Display the Results

There is a function named *show* to display the result. The users can choose to sort the genes from p values small to large, sort the log fold change from large to small by changing argument *sortby*.

```
show(res, sortby = "pvalue", shownum = 10)
##
                        logFC
                                logCPM
                                                FDR
                                                       intLR
                                                                intPValue ordinaryLR
## ENSG00000185246 2.1449454 24.27938 1.738963e-07 44.83778 2.140525e-11
                                                                            170.9681
## ENSG00000079134 1.4882846 24.43664 4.804814e-07 41.49311 1.182869e-10
                                                                            142.6679
## ENSG00000165792 0.9596607 25.82333 5.542915e-07 40.28158 2.198724e-10
                                                                            177.3485
## ENSG0000108465
                   1.3800198 24.73898 5.542915e-07 39.48873 3.299574e-10
                                                                            138.4683
## ENSG00000084733 0.9930815 24.46722 5.542915e-07 39.42363 3.411445e-10
                                                                            129.6687
## ENSG00000135617 -2.7759272 21.06611 1.067186e-06 36.87906 1.256885e-09
                                                                            132.4831
## ENSG00000178971 1.2574490 26.23903 1.067186e-06 36.68384 1.389260e-09
                                                                            118.0295
## ENSG00000181472 1.0825896 25.51406 1.067186e-06 36.68198 1.390590e-09
                                                                            126.5957
## ENSG00000167136 -1.8795301 24.45529 1.067186e-06 36.67793 1.393482e-09
                                                                            114.4304
## ENSG00000188986 -0.8065574 27.67979 1.067186e-06 36.36040 1.640017e-09
                                                                            115.4995
##
                   ordinaryPValue
## ENSG0000185246
                     4.546990e-39
## ENSG00000079134
                     6.947704e-33
## ENSG00000165792
                    1.838165e-40
## ENSG0000108465
                     5.756722e-32
## ENSG00000084733
                    4.842153e-30
## ENSG0000135617
                    1.172981e-30
## ENSG0000178971
                     1.708275e-27
## ENSG00000181472
                     2.277463e-29
## ENSG0000167136
                    1.048858e-26
## ENSG0000188986
                     6.117582e-27
```

```
show(res, sortby = "LFC", shownum = 10)
                      logFC
                              logCPM
                                              FDR
                                                               intPValue ordinaryLR
                                                      int.I.R.
## ENSG00000144214 5.247848 19.93416 7.941057e-06 28.802358 8.015346e-08
                                                                           41.94483
## ENSG00000174171 5.228006 19.75586 1.952671e-03 13.640402 2.213700e-04
                                                                           19.37467
## ENSG00000166523 5.081534 19.65238 1.220612e-02
                                                  9.146011 2.492609e-03
                                                                           11.87244
## ENSG00000171217 4.937806 19.71183 6.591471e-05 22.787563 1.809328e-06
                                                                           30.60231
## ENSG00000175265 4.836845 22.20792 1.452849e-05 27.249362 1.788341e-07
                                                                           56.69613
## ENSG00000173110 4.827374 21.83189 3.623576e-03 12.112584 5.008268e-04
                                                                           14.96935
## ENSG00000204666 4.659077 19.46568 4.999031e-03 11.318636 7.673304e-04
                                                                           14.77077
## ENSG00000171747 4.484295 19.39571 3.797217e-03 11.980474 5.376090e-04
                                                                           14.73600
## ENSG00000117616 4.398118 20.09435 2.658984e-04 18.936039 1.351749e-05
                                                                           33.98861
## ENSG00000183578 4.342930 19.98446 9.115193e-03 9.882680 1.668418e-03
                                                                           14.16119
##
                   ordinaryPValue
## ENSG0000144214
                     9.388517e-11
## ENSG0000174171
                     1.074225e-05
                     5.697087e-04
## ENSG0000166523
## ENSG0000171217
                     3.167162e-08
## ENSG0000175265
                     5.086367e-14
## ENSG0000173110
                     1.092718e-04
## ENSG00000204666
                     1.214025e-04
## ENSG0000171747
                     1.236622e-04
## ENSG0000117616
                     5.543564e-09
## ENSG0000183578
                   1.677962e-04
```

## 4 Choose the Best Methods

We found the performance of methods fluctuant as the sample size differs. We suggest the strategy to simulate synthetic negative binomial distributed data many times and apply each methods to the simulated data. Then compare their null performance and power. The simulated data uses the means and dispersions estimated from the real data. If user set "null = TRUE", it will simulate two groups of RNA-seq data with same expression level each row and will run FPR analysis. Otherwise, it will use the means of two groups and then conduct power analysis.

#### 4.1 Null performance

Suppose we want to repeat the study in previous section for 10 times with same sample size. More number of simulations are always recommended.

```
set.seed(100)
simu.res.null <- simuComp(res, nsamp=10, ntime = 10, null=TRUE)</pre>
```

The FPR performance can be shown by *summary*, the FPR should be controlled under levels.

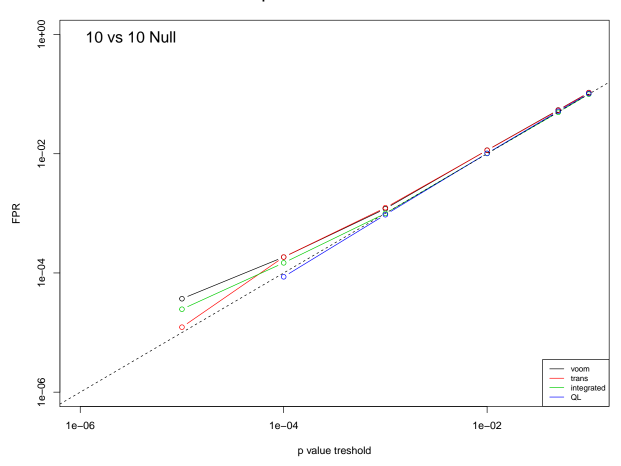
```
summary(simu.res.null)
## The four methods has FPR of:
##
               treshold = 1e-06 treshold = 1e-05 treshold = 1e-04 treshold = 0.001
## limma.voom
                               0
                                             4e-05
                                                             0.00018
                                                                               0.00119
## limma.trans
                               0
                                             1e-05
                                                             0.00018
                                                                               0.00124
                               0
## IntSEQ
                                             2e-05
                                                             0.00015
                                                                               0.00100
```

```
## edgeR.Quasi
                                            0e+00
                                                            0.00009
                                                                              0.00095
##
               treshold = 0.01 treshold = 0.05 treshold = 0.1
                       0.01148
                                        0.05423
## limma.voom
                                                        0.10624
                       0.01141
                                        0.05357
                                                        0.10586
## limma.trans
## IntSEQ
                       0.01009
                                        0.04956
                                                        0.09993
## edgeR.Quasi
                       0.01016
                                        0.05158
                                                        0.10254
```

To plot the levels against FPR:

```
plotComp(simu.res.null, text = " 10 vs 10 Null")
```

#### **Empirical FPR for four methods**



The four method all maintained the expected size.

#### 4.2 Power Analysis

This is similar with the null case, except setting the "null = FALSE":

```
set.seed(100)
simu.res.full <- simuComp(res, nsamp=10, ntime = 10, null=FALSE)</pre>
```

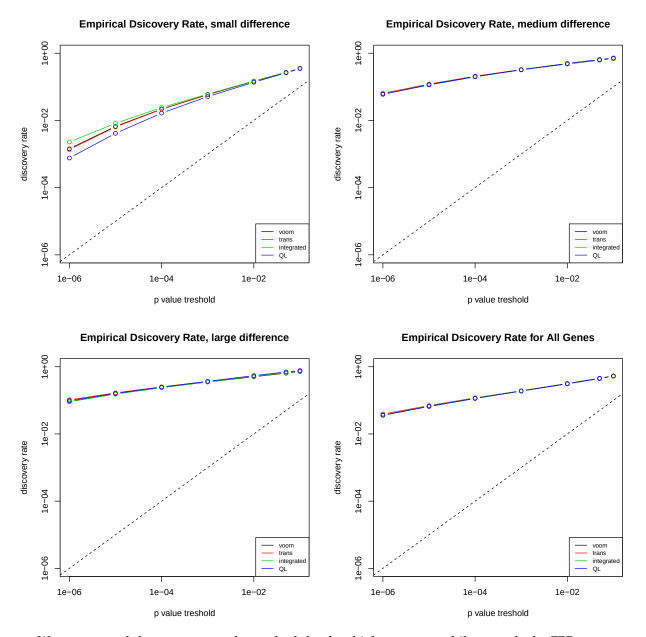
The summary function first will display the discovery rate of four method under FDR level equals to the argument "fdrlevel" passed to *simuComp*. Then we categorize genes into different groups with respect

to their log fold change. The users can choose "small" to display power of genes that are diminutively different, "medium" and "large" for medium and large log fold change genes, and "all" for all genes.

```
summary(simu.res.full, difflevel = "small")
## The number of genes in each category are:
## small medium large
  4602 2408
              1751
##
## Under FDR threshold 0.1 the four method has average discovery rate of:
  0.4036681 0.4109060 0.4110044
                                0.4145618
##
##
##
## For those genes with small difference the four method has average discovery rate of:
##
           treshold = 1e-06 treshold = 1e-05 treshold = 1e-04 treshold = 0.001
##
                 ## limma.voom
## limma.trans
                  0.00137
                               0.00641
                                             0.02153
                                                          0.05865
## IntSEQ
                 0.00228
                               0.00819
                                            0.02419
                                                         0.06076
                 0.00076
                               0.00413
                                             0.01649
## edgeR.Quasi
                                                          0.05126
  treshold = 0.01 treshold = 0.05 treshold = 0.1
## limma.voom 0.14598 0.26860
                                     0.35376
## limma.trans
                0.14752
                             0.27045
                                         0.35515
                 0.14887
                             0.26925
## IntSEQ
                                         0.35282
## edgeR.Quasi
              0.13674
                              0.26028
                                         0.34726
summary(simu.res.full, difflevel = "all")
## Under FDR threshold 0.1 the four method has average discovery rate of:
0.4036681 0.4109060 0.4110044 0.4145618
##
## -----
## For those genes with all difference the four method has average discovery rate of:
##
##
           treshold = 1e-06 treshold = 1e-05 treshold = 1e-04 treshold = 0.001
## limma.voom
                 0.03561
                           0.06578
                                         0.11373
                                                          0.18870
## limma.trans
                  0.03897
                                0.06996
                                             0.11855
                                                          0.19370
## IntSEQ
                 0.03589
                               0.06736
                                             0.11574
                                                          0.19141
## edgeR.Quasi 0.03651
                               0.06597
                                             0.11398
##
   treshold = 0.01 treshold = 0.05 treshold = 0.1
## limma.voom
           0.30853
                         0.44142
                                       0.51929
## limma.trans
                 0.31458
                             0.44802
                                          0.52443
## IntSEQ
                0.31584
                             0.44933
                                         0.52710
             0.31718
                          0.45509
                                       0.53549
## edgeR.Quasi
```

There will be four plot showing discovery rate for small, medium, large group genes.

```
par(mfrow=c(2,2))
plotComp(simu.res.full, text = "")
```



We recommend the users to use the method that has highest power while controls the FPR.

# 5 Session Information

```
## R version 3.3.3 (2017-03-06)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 10 x64 (build 14393)
##
## locale:
## [1] LC_COLLATE=Chinese (Simplified)_China.936 LC_CTYPE=Chinese (Simplified)_China.936
## [3] LC_MONETARY=Chinese (Simplified)_China.936 LC_NUMERIC=C
```

```
## [5] LC_TIME=Chinese (Simplified)_China.936
##
## attached base packages:
## [1] stats graphics grDevices utils datasets methods
                                                                   base
##
## other attached packages:
## [1] intSEQ_0.99.0 edgeR_3.16.5 limma_3.30.13 knitr_1.16
## loaded via a namespace (and not attached):
## [1] magrittr_1.5 tools_3.3.2
                                             Rcpp_0.12.11
                                                                fastGHQuad_0.2
## [5] splines_3.3.2 stringi_1.1.5
## [9] locfit_1.5-9.1 stringr_1.2.0
                                                                grid_3.3.2
                                             highr_0.6
                                             matrixStats_0.52.2 lattice_0.20-35
## [13] evaluate_0.10
```