MLExpResso: differential expression and methylation analysis

Case study using RTCGA data

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1 Package (Abstract?)

It is considered that the result of increased methylation is decreased gene expression. While, recent studies suggest that the relationship between methylation and expression is more complex than was previously thought. The package MLExpResso provides methods to test for differential expression and methylation by use of the negative binonial distribution and t-test. Additionaly 'MExpResso allows to visualize results in a simple way.

2 Standard Workflow

In this vignette we will work with the data sets containing information about gene expression and methylation for patients with breast cancer. We will analyze differences in methylation and expression for patients with different subtypes of BRCA cancer.

2.0.0.1 BRCA methylation chr17 data set

In this section, we will work with the methylation level data from TCGA database. Package contains BRCA_methylation_all dataset. BRCA_methylation_all contains information about methylation of CpG islands for patients with breast cancer. Rows of this data set correspond to patients, more precisely, to samples taken from patients. First column SUBTYPEcorresponds to a subtype of BRCA cancer, next columns correspond to CpG islands. Values inside the table indicate the methylation level of CpG island for specified sample.

```
library(MLExpResso)
```

```
## Setting options('download.file.method.GEOquery'='auto')
## Setting options('GEOquery.inmemory.gpl'=FALSE)
## No methods found in "genoset" for requests: toGenomeOrder
##
```

head(BRCA_methylation_all)[1:5,1:4]

```
##
                                SUBTYPE cg00000292 cg00002426 cg00003994
## TCGA-A1-A0SD-01A-11D-A112-05
                                   LumA 0.6055526 0.06197412 0.33345006
## TCGA-A2-A04N-01A-11D-A112-05
                                   LumA 0.7433957 0.07044132 0.32317983
## TCGA-A2-A04P-01A-31D-A032-05
                                  Basal
                                         0.2897206 0.25927969 0.02402149
## TCGA-A2-A04Q-01A-21D-A032-05
                                  Basal 0.7898920 0.63619354 0.10885097
## TCGA-A2-A04T-01A-21D-A032-05
                                  Basal
                                        0.6512270 0.27268734 0.03413620
```

In this analysis we would like to find genes with different methylation and expression. At first we need to use function map_to_gene, which generates new data frame with CpG islands mapped to genes.

```
BRCA_methylation_gen <- aggregate_probes(BRCA_methylation_all[,-1])
head(BRCA_methylation_gen)[1:5,1:4]
```

```
##
                                 {\tt c.0.627525304291071..0.61889803781234..0.561973209745455..0.64687254168}
## TCGA-A1-A0SD-01A-11D-A112-05
                                                                                                        0.6
## TCGA-A2-A04N-01A-11D-A112-05
## TCGA-A2-A04P-01A-31D-A032-05
                                                                                                        0.5
                                                                                                        0.6
## TCGA-A2-A04Q-01A-21D-A032-05
## TCGA-A2-A04T-01A-21D-A032-05
                                                                                                        0.6
##
                                                 A1BG
                                                            A2BP1
                                       X7A5
## TCGA-A1-A0SD-01A-11D-A112-05 0.13199966 0.9686056 0.03378443
## TCGA-A2-A04N-01A-11D-A112-05 0.11862215 0.9785676 0.06679088
## TCGA-A2-A04P-01A-31D-A032-05 0.08032758 0.9793897 0.29396794
```

In this case we have two conditions, connected with subtypes of breast cancer.

TCGA-A2-A04Q-01A-21D-A032-05 0.08958826 0.9718291 0.21287231 ## TCGA-A2-A04T-01A-21D-A032-05 0.13135664 0.9801575 0.21864058

Before we go to the testing, we need to define condition values for each sample. We would like to test for differences between LumA subtype and other subtypes of breast cancer, so we create a vector, which each element corresponds to a sample. Our division into this two groups relies on numbers of occurences of each subtype. The LumA subtype is the most common, in case of breast cancer.

```
condition_met <- ifelse(BRCA_methylation_all$SUBTYPE=="LumA", "LumA", "other")</pre>
head(condition met, 8)
                       "other" "other" "other" "LumA"
```

"other"

2.0.0.2BRCA_mRNAseq_all data set

"LumA"

[1] "LumA"

Data set BRCA_mRNAseq_all contains information about gene expression. This data set contains per-gene read counts computed for genes for 736 patients with breast cancer. Rows of this data set correspond to samples taken from patients. First column SUBTYPEcorresponds to a subtype of BRCA cancer, next columns correspond to genes.

```
BRCA_mRNAseq_all[1:5,1:5]
```

```
##
                                  SUBTYPE A1BG A1CF A2BP1 A2LD1
## TCGA-A1-A0SB-01A-11R-A144-07
                                           164
                                                   0
                                                        22
                                                              127
                                   Normal
## TCGA-A1-A0SD-01A-11R-A115-07
                                     LumA
                                           546
                                                   0
                                                         1
                                                              331
## TCGA-A1-A0SE-01A-11R-A084-07
                                     LumA 1341
                                                   0
                                                         2
                                                              498
## TCGA-A1-A0SF-01A-11R-A144-07
                                     LumA
                                           836
                                                   1
                                                         0
                                                              526
## TCGA-A1-A0SG-01A-11R-A144-07
                                     LumA
                                           512
                                                   3
                                                        25
                                                              451
```

In our example we will test for differential expression between groups with LumA breast cancer subtype and other subtypes of that cancer. Again we will use vector conditions, which consist of two values corresponds

```
to subtype of breast cancer: LumA and other.
condition_exp <- ifelse(BRCA_mRNAseq_all$SUBTYPE=="LumA","LumA","other")
head(condition_exp, 8)</pre>
```

```
## [1] "other" "LumA" "LumA" "LumA" "LumA" "cther" "LumA"
```

3 Testing

```
genes_comparison <- calculate_comparison_table(BRCA_mRNAseq_all[ ,-1], BRCA_methylation_gen, condition_
head(genes_comparison)</pre>
```

```
id nbinom2.log2.fold nbinom2.pval ttest.log2.fold
##
                                                              ttest.pval
## 1
                  -0.4855862 7.330514e-13
       A1BG
                                                 0.01318900 1.119637e-01
## 2 A2BP1
                   -0.8461944 4.315880e-07
                                                -0.05677500 1.019935e-02
## 3
        A2M
                   0.2352576 1.675724e-03
                                                 0.03310777 1.686741e-01
## 4 A2ML1
                   4.1919667 8.164823e-201
                                                 0.17016379 1.019044e-12
## 5 A4GALT
                   -0.1927530 9.741784e-03
                                                -0.00841392 5.203636e-01
## 6 A4GNT
                   0.2208814 8.021080e-02
                                                0.06387711 1.957709e-05
```

3.1 Choosing most different genes

```
Sorting ...
```

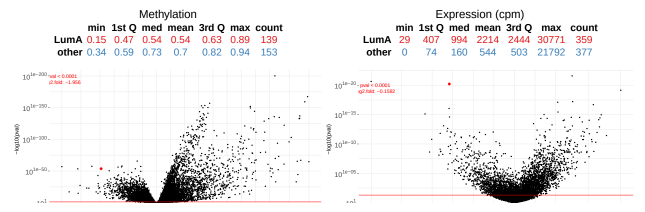
```
genes_comparison_sorted <- genes_comparison[order(genes_comparison$ttest.pval), ]
head(genes_comparison_sorted)</pre>
```

```
id nbinom2.log2.fold nbinom2.pval ttest.log2.fold
## 10601 TNFRSF10A
                           0.2038466 2.272347e-04
                                                         0.1401746
## 10218
            TBX19
                           1.0212930 3.226882e-49
                                                        -0.3479977
## 4958
            IGFALS
                          -1.9559531 1.892160e-54
                                                        -0.1581887
## 6214
            MGST1
                           0.4688093 8.122084e-08
                                                         0.2583506
                          -0.1772576 5.466990e-05
## 5895
            LUC7L
                                                         0.1414415
## 8870
              RRM2
                           1.4786220 5.630314e-103
                                                         0.1051548
##
           ttest.pval
## 10601 2.568571e-22
## 10218 2.252784e-21
## 4958 6.144850e-21
## 6214 7.121403e-20
## 5895
        6.298008e-18
## 8870 7.790260e-18
IGFALS!
```

4 Visualization

```
Visualizing chosen gene - IGFALS.
```

```
test_exp <- genes_comparison[ ,c(1,2,3)]
test_met <- genes_comparison[ ,c(1,4,5)]</pre>
```



Note that plot_gene methylation require data frame with cpg islands, not genes.

