# Vignette Title

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## 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 ... provides methods to test for differential expression and methylation by use of the negative binonial distribution and t-test. Additionally package ... allows to visualize results in a simple way.

#### 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.

#### Function test\_ diff

The main function of the package is test\_diff. It allows to find differences between genes methylation or expression, taking into account additional information about samples.

#### Methylation

Methylation is a process by which methyl groups are added to the DNA molecule. It can change the activity of a DNA without changing the sequence. DNA methylation typically acts to repress gene transcription. But there exist situations in which adding the methyl groups intensify it. DNA methylation is associated with a lots of key processes including genomic imprinting, repression of transposable elements, aging and carcinogenesis. In our work we want to bind methylation process and carcinogenesis.

#### BRCA\_methylation\_chr17 data set

In this section, we will work with the methylation level data from TCGA database. Package contains BRCA\_methylation\_chr17 dataset. BRCA\_methylation\_chr17 contains information about methylation of CpG islands located on 17th chromosome for patienst 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(MetExpR)
head(BRCA_methylation_chr17)[1:5,1:4]
                                SUBTYPE cg00021527 cg00031162 cg00032227
##
## TCGA-A1-A0SD-01A-11D-A112-05
                                   LumA 0.03781858
                                                   0.7910348 0.006391233
## TCGA-A2-A04N-01A-11D-A112-05
                                   LumA 0.01437552
                                                    0.7359370 0.008752293
## TCGA-A2-A04P-01A-31D-A032-05
                                  Basal 0.01360124
                                                    0.6967802 0.009442039
## TCGA-A2-A04Q-01A-21D-A032-05
                                  Basal 0.01525656
                                                    0.5341244 0.014674247
## TCGA-A2-A04T-01A-21D-A032-05
                                  Basal 0.01167384 0.7378100 0.012251559
```

#### Data preparation

In this analysis we would like to find genes with different methylation level. At first we need to use function map\_to\_gene, which generates new data frame with CpG islands mapped to genes.

```
BRCA_methylation_chr17_gen <- map_to_gene(BRCA_methylation_chr17[,-1])
head(BRCA_methylation_chr17_gen)[1:5,1:4]</pre>
```

```
## TCGA-A2-A04N-01A-11D-A112-05 0.7148533 0.8625816 0.24294092 0.7835302 ## TCGA-A2-A04N-01A-11D-A112-05 0.5850106 0.8355825 0.21367129 0.8466190 ## TCGA-A2-A04P-01A-31D-A032-05 0.4495537 0.8786166 0.03277413 0.3417919 ## TCGA-A2-A04Q-01A-21D-A032-05 0.7120650 0.8819490 0.03460160 0.7264985 ## TCGA-A2-A04T-01A-21D-A032-05 0.6010397 0.7739978 0.02501599 0.6276399
```

Function test\_diff allows us to test for differences between the base means for two or more conditions.

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

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 vector, which each element corresponds to a sample. Our division into this two group relies on numbers of occurences of each subtype. The LumA subtype is the most common, in case of breast cancer.

```
condition <- ifelse(BRCA_methylation_chr17$SUBTYPE=="LumA","LumA", "other")
head(condition,8)</pre>
```

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

#### T-test

One of the most used tools for testing differences between values is t-test. The null hypothesis we have consider, is that means in two groups are equal. To use it in test\_diff function, we set value of parameter test on "ttest".

```
test.mety <- test_diff(BRCA_methylation_chr17_gen, condition, test="ttest")</pre>
```

As a result we obtain a data frame with columns corresponds to: id of gene, mean, logarithm of fold change, p-value for t-test, adjusted p-value (BH method). For more information about customizing this function see the help page for test\_diff.

#### head(test.mety)

```
##
                id
                                log.fold
                                                               padi
                                                 pval
                        mean
## ICAM2
             ICAM2 0.3330801 -0.15151320 3.754116e-17 3.063359e-14
## RILP
              RILP 0.3341447 -0.05073691 2.575168e-13 1.050668e-10
## PIPOX
             PIPOX 0.3647812 0.11505558 5.360053e-12 1.196885e-09
## TNFSF12 TNFSF12 0.2485025 -0.13412855 5.867083e-12 1.196885e-09
## CD7
               CD7 0.8127112 0.09822690 1.641919e-11 2.679612e-09
## KSR1
              KSR1 0.5549808 0.19973400 2.054467e-11 2.794075e-09
```

#### Expression

Gene expression is the process by which information from a gene is used in the synthesis of proteins. The process of gene expression is used by all known life.

#### BRCA\_mRNAseq\_chr17 data set

In this section we will use data set BRCA\_mRNAseq\_chr17, which contains information about gene expression. This data set contains per-gene read counts computed for genes from 17th chromosome for 100 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\_chr17[1:5,1:5]

```
##
                                 SUBTYPE AANAT AARSD1 AATF AATK
## TCGA-A1-A0SB-01A-11R-A144-07
                                 Normal
                                             9
                                                 2354 2870
                                                            317
## TCGA-A1-AOSD-O1A-11R-A115-07
                                    LumA
                                             2
                                                 1846 5656
                                                            312
## TCGA-A1-A0SE-01A-11R-A084-07
                                    LumA
                                            11
                                                 3391 9522
                                                            736
## TCGA-A1-A0SF-01A-11R-A144-07
                                                            169
                                    LumA
                                             0
                                                 2169 4625
## TCGA-A1-A0SG-01A-11R-A144-07
                                    LumA
                                                 2273 3473
```

#### Negative binomial test

Negative binomial test, which uses negative binomial distribution is an another tool for finding differencial expression between our conditions.

As in the t-test we also need a description of the samples, which we keep in a vector, whose elements correspond to different gorups.

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<-ifelse(BRCA_mRNAseq_chr17$SUBTYPE=="LumA","LumA","other")
head(condition,8)</pre>
```

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

For using negative binomial test, in function test\_diff we set value "nbinom" for parameter test. Evaluation for nbinorm may take a few minutes.

```
test.expr <- test_diff(BRCA_mRNAseq_chr17[,-1], condition, test="nbinom")</pre>
```

As a result we obtain the following data frame:

```
head(test.expr)
```

```
## id mean log.fold pval padj

## 1 AANAT 3.455436 -0.40216187 0.9453634 0.9920531

## 2 AARSD1 2779.448414 0.07427334 0.6627296 0.8992428

## 3 AATF 6750.269650 -0.13313947 0.6864598 0.9111193

## 4 AATK 352.805108 -0.02272566 0.8051408 0.9585881

## 5 ABCA5 1933.257431 0.07031881 0.5837682 0.8489059

## 6 ABCA6 689.547294 0.44680041 0.1289441 0.3857753
```

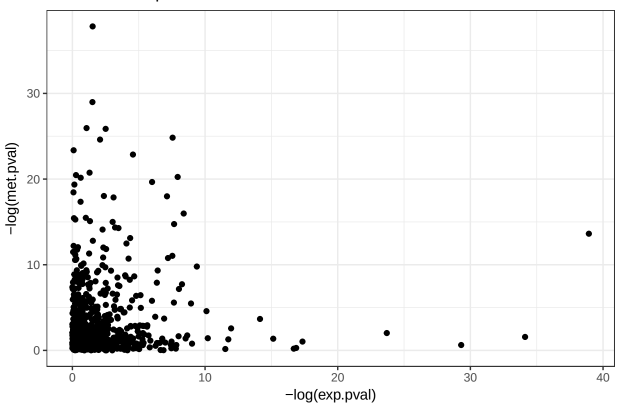
#### Visualization

#### log-log p-value

Firstly, we want to visualise the p-values for expression and methylation from negative binomial test and t-test respectively.

p\_values\_plot(test.expr, test.mety)

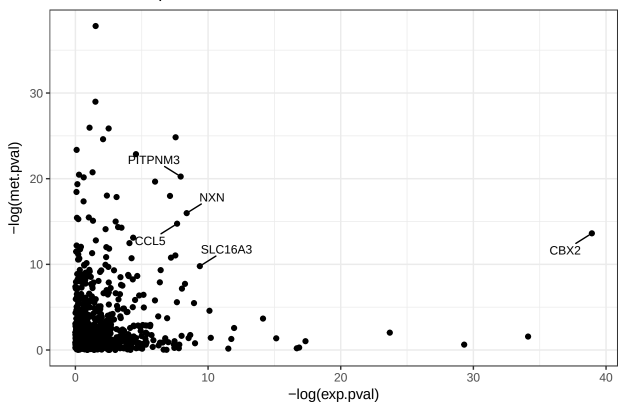
### P-values comparison



Additionally, names parameter allows to mark genes with sum of p-values for methylation and expression, lower than given value. Value of parameter names defines, number of genes to label.

```
p_values_plot(test.expr, test.mety, names = 5)
```

## P-values comparison



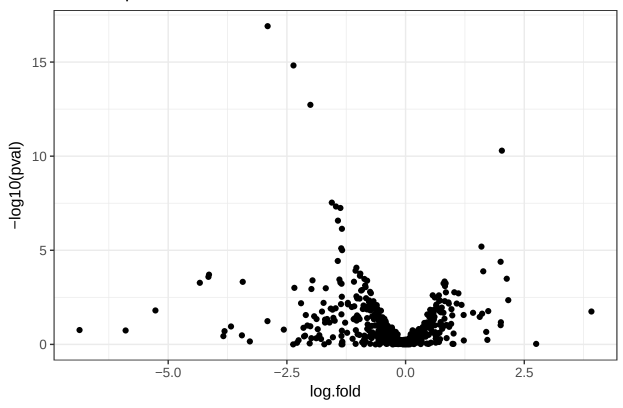
To read more about p\_values\_plot (e.g other ways to labeling genes) see help page for that function.

## Volcano plot

For identify changes in our data sets we use a volcano plot - some type of scatter-plot. It plots logarithm of p-value versus logarithm of fold-change on the y and x axes, respectively.

volcano\_plot(test.expr)

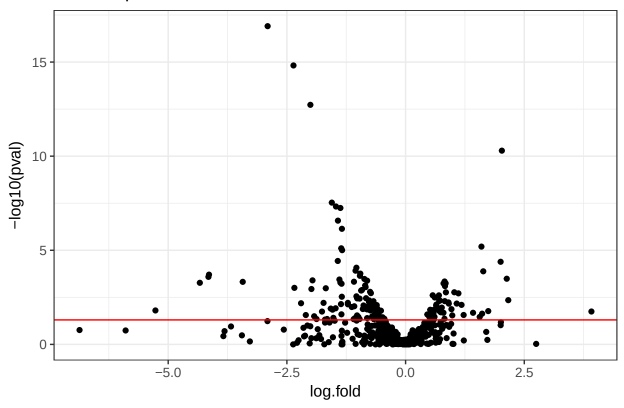
# Volcano plot



Function volcano\_plot has parameters that allow to better analyze the results: line and names. The line parameter allows to set the horizontal line on plot on selected value. The names parameter signs choosen number of genes with the lowest p-value.

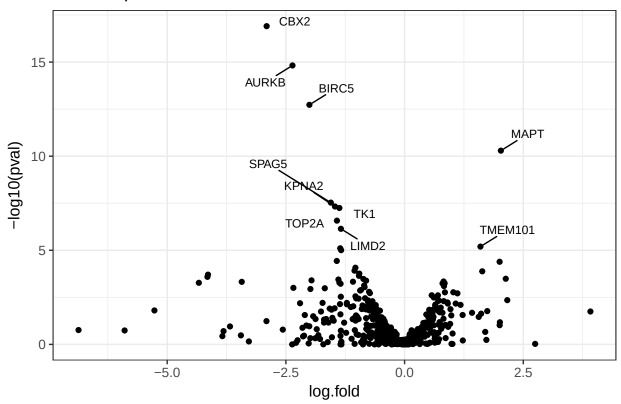
volcano\_plot(test.expr, line = 0.05)

# Volcano plot



volcano\_plot(test.expr, names = 10)

## Volcano plot



#### Methylation and expression for one gene.

In the end we want to present the distribution of methylation and expression for choosen genes.

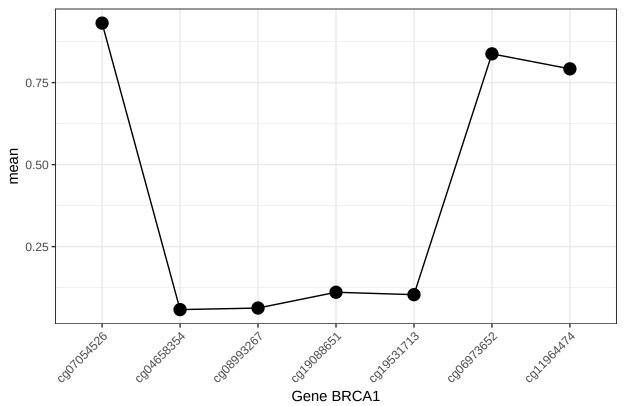
Function CpG mean computes methylation means of CpG islands for choosen gene. In this case: "BRCA1"

```
BRCA1_gene <- CpG_mean(BRCA_methylation_chr17, "BRCA1")
head(BRCA1_gene)</pre>
```

```
##
               Name Symbol CPG_ISLAND_LOCATIONS
## 7047
        cg07054526 BRCA1 17:38525979-38526990 0.93138624
## 4712
         cg04658354
                    BRCA1 17:38530194-38531162 0.05802403
## 8929
        cg08993267
                    BRCA1 17:38530194-38531162 0.06280625
## 19075 cg19088651
                    BRCA1 17:38530194-38531162 0.11090400
## 19527 cg19531713
                    BRCA1 17:38530194-38531162 0.10361921
        cg06973652
                    BRCA1 17:38531525-38532730 0.83772148
```

First we visualize the mean value of methylation for each CpG island on this gene using function genereg\_vs\_met.

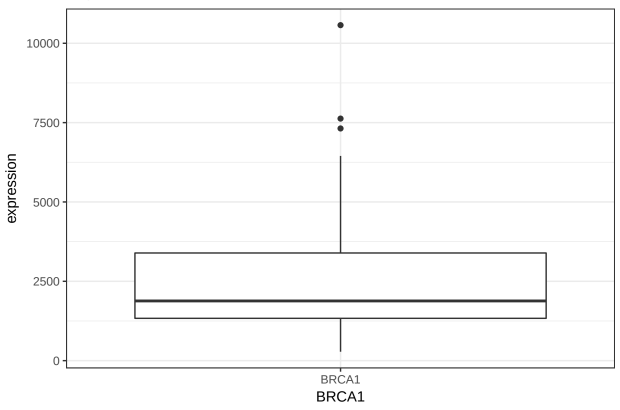
# Methylation of gene BRCA1



With function boxplot\_gene\_expr, we consider the distribution of expression from 100 probes.

boxplot\_gene\_expr(BRCA\_mRNAseq\_chr17,"BRCA1")

# Expression of BRCA1



It is easy to set both plots together.

