MLExpResso

Cheat Sheet

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Introduction

MLExpResso is an R package for integrative analyses and visualization of gene expression and DNA methylation data.

Key functions of this package are:

- identification of genes with affected expression calculate test() function,
- identification of DMR differentially methylated regions - calculate test() function.
- identification of regions with changes in expression and methylation - calculate comparition table() function,
- visualization of identified regions plot gene() and plot volcanoes() functions.

The joint modeling and visualization of genes expression and methylation improve interpretability of identified signals.

Identification of genes with affected expression

MLExpResso::calculate test(data, condition, test)

Function calculate test() computes log folds, p-values and means for chosen test for both, methylation and expression data.

> BRCA_exp[1:3, 1:5]

	SUBTYPE	AANAT	AARSDI	AATE	AATK
TCGA-A1-A0SB-01A-11R-A144-07	Normal	9	2354	2870	317
TCGA-A1-A0SD-01A-11R-A115-07	LumA	2	1846	5656	312
TCGA-A1-A0SE-01A-11R-A084-07	LumA	11	3391	9522	736

Example

```
library("MLExpResso")
library("MLExpRessoData")
exp <- BRCA exp[ ,-1]
gr exp <- BRCA exp$SUBTYPE
gr exp <- ifelse(gr exp == 'LumA', 'LumA', 'other')</pre>
res exp <- calculate test(exp, gr exp, 'lrt')
```

> head(res exp)

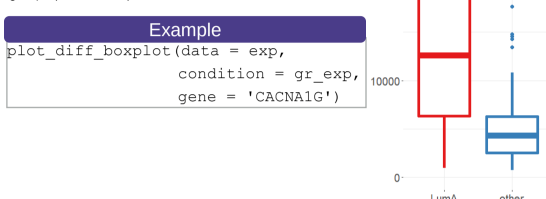
	meaa.	CO_CMP)				
	id	log2.fold	p∨al	mean_LumA	mean_other	mean
1	AURKB	2.3	3.2e-32	539	2324	1485
2	CBX2	2.9	2.8e-26	633	4297	2574
3	KPNA2	1.4	8.6e-24	11547	26427	19434
4	PRR11	3.8	2.3e-22	396	3480	2031
5	BIRC5	2.0	2.0e-21	1957	6658	4449
6	GSG2	1.4	3.5e-21	278	629	464

Argument test allows using many different statistic tests for finding differences in expression. All available values are in the table below.

Test		
student's t-tets		
negative binomial test		
likelihood-ratio test		
quasi-likelihood F-test		

MLExpResso::plot diff boxplot(data, condition, gene)

Function plot diff boxplot() generates a boxplot of values from choosen data frame column with division in groups (two or more).



CACNA1G

40000

30000

20000

MLExpRessoData

The methodology is supplemented with example applications to The Cancer Genome Atlas data.

MLExpRssoData is an R package which contains information from The Cancer Genome Atlas (TCGA) Data Portal. Data sets in this package are based on Bioconductor package RTCGA. In examples, we use both, methylation and expression data.

- BRCA exp It contains information about gene expression; read counts per-gene, computed for genes for 736 patients with breast cancer. Rows of this data set correspond to samples taken from patients. First column SUBTYPE corresponds to a subtype of BRCA cancer, next columns correspond to genes.
- **BRCA_met** It contains information about methylation of CpG probes for patients with breast cancer. Rows of this data set correspond to patients. more precisely, to samples taken from patients. First column SUBTYPE corresponds to a subtype of BRCA cancer, next columns correspond to CpG probes. Values inside the table indicate the percentage methylation level of CpG probe for a specified sample.

For aggregation CpG probes to correspond genes we use the Illumina human methylation data set from TxDb.Hsapiens.UCSC.hg18.knownGene Bioconductor package.

Identification of DMR - differentially methylated regions

MLExpResso::calculate test(data, condition, test)

Argument test allows using two different statistic tests for finding differences in methylation levels. All available values are in the table on the right.

Value	Test
'ttest'	student's t-test
'methyanalysis'	quasi-likelihood F-test

MLExpResso::aggregate probes(data)

Function aggregate probes() aggregates CpG probes to corresponding genes using. by default, the Illumina human methylation data.

> BRCA_met[1:3, 1:5]

	SUBTYPE	cg0002152/	cg00031162	cg0003222/	cg00050312
TCGA-A1-A0SD-01A-11D-A112-05	LumA	0.038	0.79	0.0064	0.024
TCGA-A2-A04N-01A-11D-A112-05	LumA	0.014	0.74	0.0088	0.028
TCGA-A2-A04P-01A-31D-A032-05	Basal	0.014	0.70	0.0094	0.014
, · · ·					

Example

```
met <- aggregate probes(BRCA met)</pre>
gr met <- BRCA met$SUBTYPE
gr met <- ifelse(gr met == 'LumA', 'LumA', 'other')</pre>
res met <- calculate test(met, gr met, 'ttest')
```

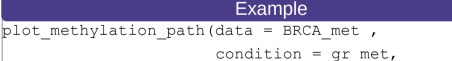
>	head(res	s_met)				
	id	log2.fold	p∨al	mean_LumA	mean_other	mean
1	ICAM2	-0.152	3.8e-17	0.25	0.41	0.33
2	RILP	-0.051	2.6e-13	0.31	0.36	0.33
3	PIPOX	0.115	5.4e-12	0.42	0.31	0.36
4	TNFSF12	-0.134	5.9e-12	0.18	0.31	0.25
5	CD7	0.098	1.6e-11	0.86	0.77	0.81
6	KSR1	0.200	2.1e-11	0.66	0.46	0.55

MLExpResso::plot_methylation_path(data, condition, gene)

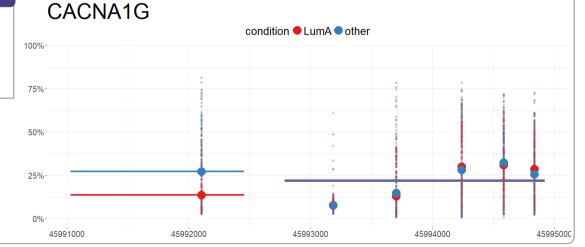
Function plot methylation path() visualizes a chosen gene with marked CpG probes. Y axis describes methylation level.

X axis describes a location of the probe on the chromosome.

Horizontal lines show the mean methylation level for each Island in a division to groups. Groups are defined by colors. Large dots symbolize means of methylation level for CpG probes, small dots symbolize methylation levels for each observation.



gene = 'CACNA1G', observ = T)



Comparing test results

MLExpResso::calculate comparison table(data1, data2, condition1, condition2, test1, test2)

Function calculate_comparison_table() produces a dataset containing p-values and folds from tests evaluated on two datasets e.g methylation or expression. In addition, it produces an importance ranking column, which is the geometric mean of p-values from both tests and a column with a number of probes related to the gene.

```
Example
calculate comparison table (data1 = BRCA exp[,-1], data2 = BRCA met gen,
                              condition1 = condition exp, condition2 = condition met,
                              test1 = "nbinom2", test2 = "ttest")
> head(genes_comparison)
       id nbinom2.log2.fold nbinom2.pval ttest.log2.fold ttest.pval geom.mean.rank no.probes
59
   AURKB
                        2.4
                                1.7e-37
                                                 0.00174
                                                            2.1e-01
                                                                           1.9e-19
102
   CBX2
                        2.9
                                5.4e-31
                                                 0.05847
                                                           1.2e-06
                                                                           8.1e-19
                                                                                           2
327
   KPNA2
                        1.5
                                3.4e-26
                                                0.00121
                                                           7.5e-01
                                                                           1.6e-13
                                                                                           1
277
    GSG2
                        1.4
                                3.3e-25
                                                -0.00186
                                                            2.4e-01
                                                                           2.8e-13
                                                                                           2
```

-0.00054

0.04868

5.3e-01

1.6e-05

Expression (cpm)

LumA 977 6351 12628 14302 18716 41717

min 1st Q med mean 3rd Q max count

2.2e-12

2.6e-12

1

2

Visualization of identified regions

2.0

4.3

9.5e-24

4.1e-19

BIRC5

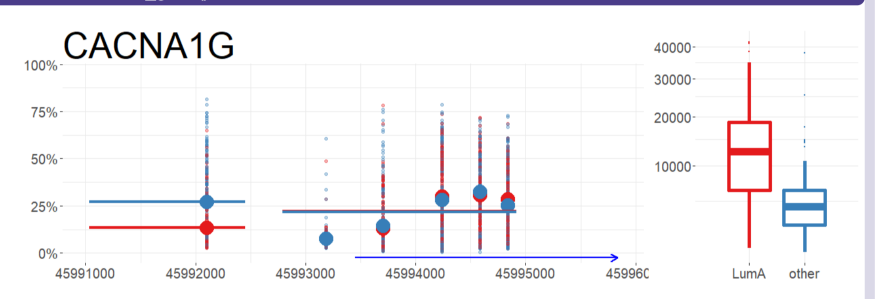
334 KRT16

Plot_gene()

MLExpResso::plot_genes(data.m, data.e, condition.m, condition.e, gene)

Function plot_gene() generates a dashboard with methylation path for methylation and boxplots for groups for chosen gene.

Example plot gene (data.m = BRCA met, data.e = BRCA exp,condition.m = gr met, condition.e = gr exp, gene = "CACNA1G", observ = TRUE, show.gene = TRUE,islands = TRUE)



Plot_volcanoes()

Methylation

min 1st Q med mean 3rd Q max count

MLExpResso::plot_volcanoes(data.m, data.e, condition.m, condition.e, gene, test.m, test.e)

Function **plot_volcanoes()** generates a dashboard with volcano plots for expression and methylation. Also it adds a tables with basic statistics for chosen gene.

