

MLExpResso

Cheat Sheet

Aleksandra Dąbrowska [aut, cre]

Alicja Gosiewska [aut]

Przemysław Biecek [aut, ths]



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_comparison_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.

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	AARSD1	AATF	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')
res_exp <- calculate_test(exp, gr_exp, 'lrt')
```

```
> head(res_exp)
```

	id	log2.fold	pval	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.

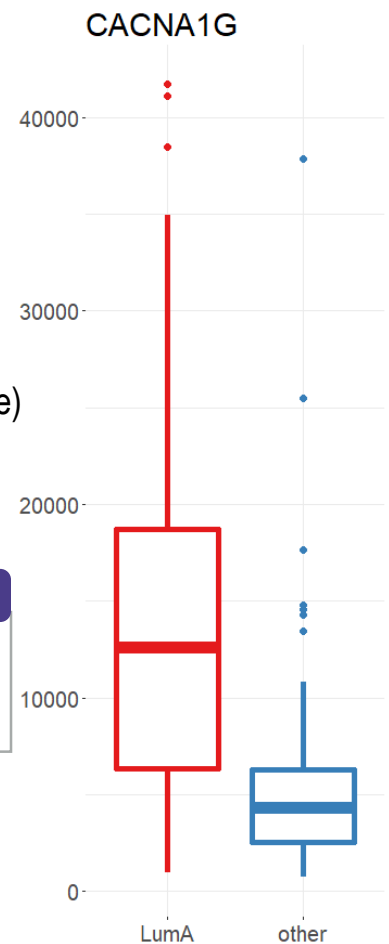
Value	Test
'ttest'	student's t-tets
'nbinom2'	negative binomial test
'lrt'	likelihood-ratio test
'qlf'	quasi-likelihood F-test

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).

Example

```
plot_diff_boxplot(data = exp,
                  condition = gr_exp,
                  gene = 'CACNA1G')
```



MLExpRessoData

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

MLExpRessoData 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
'methanalysis'	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	cg00021527	cg00031162	cg00032227	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)
gr_met <- BRCA_met$SUBTYPE
gr_met <- ifelse(gr_met == 'LumA', 'LumA', 'other')
res_met <- calculate_test(met, gr_met, 'ttest')
> head(res_met)
```

	id	log2.fold	pval	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	KSRL	0.200	2.1e-11	0.66	0.46	0.55

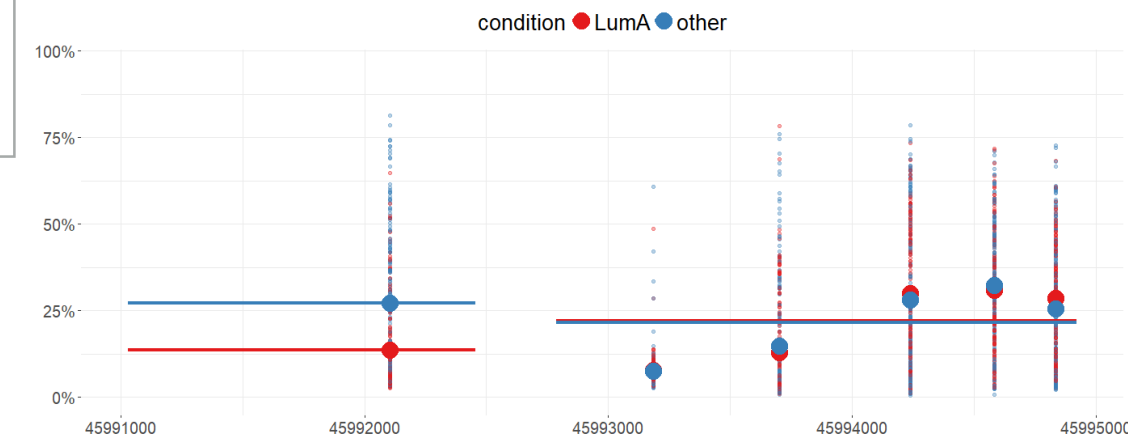
MLExpResso::plot_methylation_path(data, condition, gene)

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.

Example

```
plot_methylation_path(data = BRCA_met,
                     condition = gr_met,
                     gene = 'CACNA1G',
                     observ = T)
```

CACNA1G



Comparing test results

MLEPResso::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	2
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
66	BIRC5	2.0	9.5e-24	-0.00054	5.3e-01	2.2e-12	1
334	KRT16	4.3	4.1e-19	0.04868	1.6e-05	2.6e-12	2

Visualization of identified regions

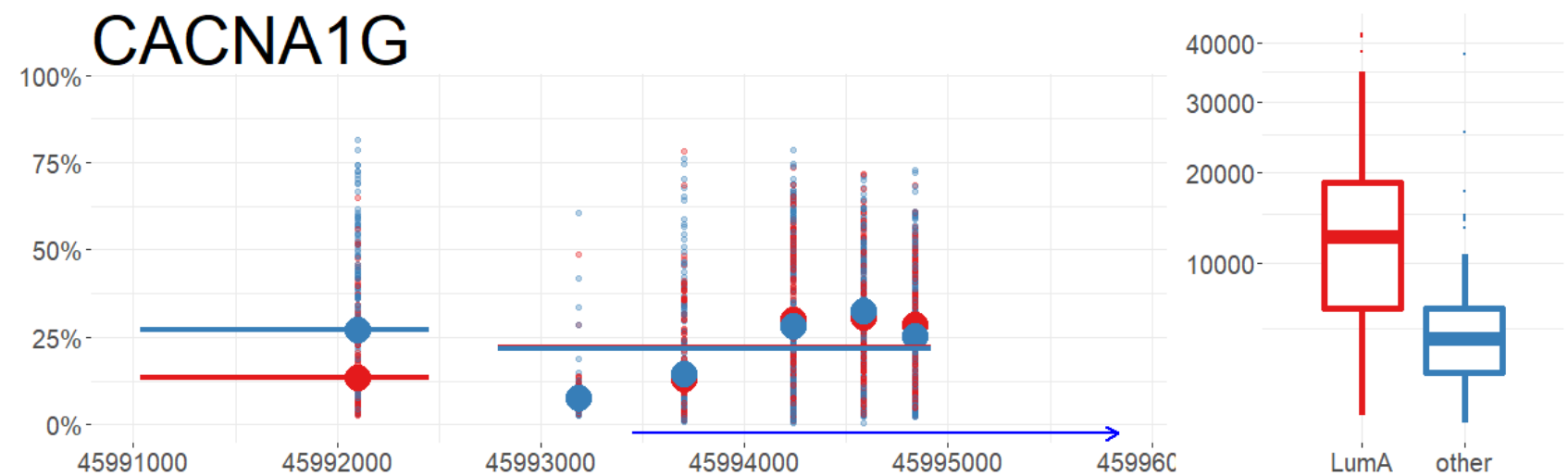
Plot_gene()

MLEPResso::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()

MLEPResso::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.

Example

```
plot_volcanoes(data.m = met,
                data.e = exp,
                condition.m = gr_met,
                condition.e = gr_exp,
                gene = 'CACNA1G',
                test.m = res_met,
                test.e = res_exp)
```

