Package ‘Moonlight2R’

May 30, 2024

Type Package

Title Identify oncogenes and tumor suppressor genes from omics data

Version 1.2.0

Depends R (>= 4.3), doParallel, foreach

Imports parmigene, randomForest, gplots, circlize, RColorBrewer, HiveR, clusterProfiler, DOSE, Biobase, grDevices, graphics, GEOquery, stats, purrr, RISmed, grid, utils, ComplexHeatmap, GenomicRanges, dplyr, fuzzyjoin, rtracklayer, magrittr, qpdf, readr, seqminer, stringr, tibble, tidyHeatmap, tidyR, AnnotationHub, easyPubMed, org.Hs.eg.db, EpiMix, BiocGenerics, ggplot2, ExperimentHub

Description The understanding of cancer mechanism requires the identification of genes playing a role in the development of the pathology and the characterization of their role (notably oncogenes and tumor suppressors). We present an updated version of the R/bioconductor package called MoonlightR, namely Moonlight2R, which returns a list of candidate driver genes for specific cancer types on the basis of omics data integration. The Moonlight framework contains a primary layer where gene expression data and information about biological processes are integrated to predict genes called oncogenic mediators, divided into putative tumor suppressors and putative oncogenes. This is done through functional enrichment analyses, gene regulatory networks and upstream regulator analyses to score the importance of well-known biological processes with respect to the studied cancer type. By evaluating the effect of the oncogenic mediators on biological processes or through random forests, the primary layer predicts two putative roles for the oncogenic mediators: i) tumor suppressor genes (TSGs) and ii) oncogenes (OCGs). As gene expression data alone is not enough to explain the deregulation of the genes, a second layer of evidence is needed. We have automated the integration of a secondary mutational layer through new functionalities in Moonlight2R. These functionalities analyze mutations in the cancer cohort and classifies these into driver and passenger mutations using the driver mutation prediction tool, CScape-somatic. Those oncogenic mediators with at
At least one driver mutation are retained as the driver genes. As a consequence, this methodology does not only identify genes playing a dual role (e.g. TSG in one cancer type and OCG in another) but also helps in elucidating the biological processes underlying their specific roles. In particular, Moonlight2R can be used to discover OCGs and TSGs in the same cancer type. This may for instance help in answering the question whether some genes change role between early stages (I, II) and late stages (III, IV). In the future, this analysis could be useful to determine the causes of different resistances to chemotherapeutic treatments.

License GPL-3

biocViews DNAMethylation, DifferentialMethylation, GeneRegulation, GeneExpression, MethylationArray, DifferentialExpression, Pathways, Network, Survival, GeneSetEnrichment, NetworkEnrichment

Suggests BiocStyle, knitr, rmarkdown, testthat (>= 3.0.0), devtools, roxygen2, png

SystemRequirements C SnapeSomatic

VignetteBuilder knitr

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BugReports https://github.com/ELELAB/Moonlight2R/issues

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Description

This function annotated a confidence level to the score

Usage

confidence(s, type)

Arguments

s  the score

  type  coding or noncoding

Value

returns a confidence level or remark/error message

Examples

remark <- confidence(0.8, type='Coding')
**cscape_somatic_output**  
*Cscape-somatic annotations of TCGA-LUAD*

**Description**
Output from DMA. This contains the cscape-somatic annotations for all differentially expressed genes.

**Usage**
```r
data(cscape_somatic_output)
```

**Format**
A 645x7 matrix.

**Value**
A 645x7 matrix.

---

**dataDMA**  
*Output example from the function Driver Mutation Analysis*

**Description**
The predicted driver genes, which have at least one driver mutation.

**Usage**
```r
data(dataDMA)
```

**Format**
A list of two.

**Value**
A list of two, containing 0 tumor-suppressor and 1 oncogene.
**dataFEA**

*Functional enrichment analysis*

**Description**

The output of the FEA function which does enrichment analysis.

**Usage**

```r
data(dataFEA)
```

**Format**

A dataframe of dimension 101x7

**Details**

The input to the FEA is the differentially expressed genes.

**Value**

A dataframe of dimension 101x7

---

**dataFilt**

*Gene expression data from TCGA-LUAD*

**Description**

A matrix that provides processed gene expression data (obtained from RNA seq) from the TCGA-LUAD project.

**Usage**

```r
data(dataFilt)
```

**Format**

A 3000x20 matrix

**Details**

The matrix contains the genes in rows and samples in columns. The data has been downloaded and processed using TCGAbiolinks.

**Value**

A 3000x20 matrix
dataGLS

**Literature search of driver genes**

**Description**
A tibble containing results of literature search where predicted driver genes stored in dataDMA were queried for their role as drivers in PubMed

**Usage**
data(dataGLS)

**Format**
A 13x8 tibble.

**Details**
The tibble contains PubMed IDs, doi, title, abstract, year of publication, keywords, and total number of publications for the genes.

**Value**
A 13x8 tibble.

dataGMA

**Output example from GMA function**

**Description**
The predicted driver genes based on methylation evidence

**Usage**
data(dataGMA)

**Format**
A list of length two

**Details**
The data is a list of two elements where each element represents predicted oncogenes and tumor suppressors

**Value**
A list of length two
**dataGRN**

*Gene regulatory network*

**Description**

The output of the GRN function which finds connections between genes.

**Usage**

```r
data(dataGRN)
```

**Format**

A list of 2 elements where the first element is a 23x613 matrix and the second element is a vector of length 23

**Details**

The input to the GRN is the differentially expressed genes and the gene expression data.

**Value**

A list of 2 elements where the first element is a 23x613 matrix and the second element is a vector of length 23

**dataGRN_no_noise**

*Gene regulatory network*

**Description**

The output of the GRN function which finds connections between genes where the noise is set to 0 for testing reproducibility purposes.

**Usage**

```r
data(dataGRN_no_noise)
```

**Format**

A list of 2 elements where the first element is a 23x613 matrix and the second element is a vector of length 23

**Details**

The input to the GRN is the differentially expressed genes and the gene expression data.
**dataMAF**  

*Value*  
A list of 2 elements where the first element is a 23x613 matrix and the second element is a vector of length 23

**Description**  
An examplary MAF file from TCGA on lung cancer LUAD. It contains 500 randomly selected mutations.

**Usage**  

data(dataMAF)

**Format**  
A 500x141 matrix.

**Value**  
A 500x141 matrix.

---

**dataMethyl**  

*Description*  
A data matrix containing methylation data from TCGA-LUAD where CpG probes are in rows and samples are in columns.

**Usage**  

data(dataMethyl)

**Format**  
A 73x27 matrix.

**Details**  
The CpG probes are in rows and samples are in columns.

**Value**  
A 73x27 matrix.
Description

The predicted TSGs and OCGs and their moonlight gene z-score based on the small sample TCGA-LUAD data. The PRA() were run with expert-based approach with apoptosis and proliferation of cells.

Usage

data(dataPRA)

Format

A list of two.

Value

A list of two.

Description

The output of the URA function which carries out the upstream regulator analysis

Usage

data(dataURA)

Format

A 23x2 matrix

Details

The input to URA is the output of GRN and a list of biological processes and the differentially expressed genes

Value

A 23x2 matrix
**dataURA_plot**

*Upstream regulator analysis*

**Description**

The output of the URA function which carries out the upstream regulator analysis.

**Usage**

data(dataURA_plot)

**Format**

A 12x2 matrix

**Details**

This URA data is used to showcase some of the visualization functions.

**Value**

A 12x2 matrix

---

**DEGsmatrix**

*Differentially expressed genes*

**Description**

A matrix containing differentially expressed genes between lung cancer and normal samples found using TCGA-LUAD data and TCGAbiolinks.

**Usage**

data(DEGsmatrix)

**Format**

A 3390x5 matrix

**Details**

The matrix contains the differentially expressed genes in rows and log2 fold change and FDR values in columns.

**Value**

A 3390x5 matrix
**DEG_Methylation_Annotations**

*Output example from GMA function*

**Description**

Output file from running GMA function which is a summary of DEGs and associated CpG probes

**Usage**

```r
data(DEG_Methylation_Annotations)
```

**Format**

A 3435x35 tibble

**Details**

The data is a table where each row is a CpG probe in a DEG. Various annotations such as start/end site of CpG probe, promoter/enhancer annotations, NCG annotations are included in the table.

**Value**

A 3435x35 tibble

---

**DEG_Mutations_Annotations**

*Differentially expressed genes’s Mutations*

**Description**

Output from DMA. This contains the differentially expressed genes’s mutations and all annotations generated in DMA() on the TCGA-LUAD project.

**Usage**

```r
data(DEG_Mutations_Annotations)
```

**Format**

A 3561x173 matrix.

**Value**

A 3561x173 matrix.
DiseaseList

Description
A dataset containing information about 101 cancer-related biological processes.

Usage
data(DiseaseList)

Format
A list of 101 elements

Details
The dataset contains a list of the 101 biological processes which includes genes playing a role in each biological processes including literature findings of the genes’ function in the biological processes.

Value
A list of 101 elements

DMA

Description
This function carries out the driver mutation analysis.

Usage
DMA(
dataMAF,
dataDEGs,
dataPRA,
runCscape = TRUE,
coding_file,
noncoding_file,
results_folder = "./DMAresults"
)


Arguments

dataMAF A MAF file rda object. The MAF file must at least contain the following columns:
- Hugo_Symbol eg. BRCA1
- Chromosome eg. chr1
- Start_Position eg. 54402
- End_Position e.g. 54443
- Strand eg. +
- Variant_Classification
- Variant_Type
- Reference_Allele
- Tumor_Seq_Allele1
- Tumor_Seq_Allele2

dataDEGs Output DEA function.
dataPRA Output PRA function.
runCscape Boolean. If FALSE will load CScape output file from results-folder Default = TRUE.
coding_file A character string. Path to and name of CScape-somatic coding file. Can be downloaded at http://cscape-somatic.biocompute.org.uk/#download. The .tbi file must be placed in the same folder.
noncoding_file A character string. Path to and name of CScape-somatic noncoding file. Can be downloaded at http://cscape-somatic.biocompute.org.uk/#download. The .tbi file must be placed in the same folder.
results_folder A character string. Path to the results generated by this function.

Details

For more information about the different annotations added to the mutations please see the documentation as follows: data(NCG), data(EncodePromoters), data(LOC_protein) data(LOC_transcription) and data(LOC_translation).

Value

List of two, containing TSGs and OCGs with at least one driver mutation. Additionally files are saved in results_folder. All output files are in compressed .rda format.

DEG_mutations_annotations.rda All differentially expressed genes’ mutations and their annotations. These annotations include e.g. CScape-somatic assessment, Level of Consequence, overlap with promoter sites and information from Network of Cancer Genes (NCG 7.0). All information from MAF and DEA is contained.

Oncogenic_mediators_annotation_summary.rda All oncogenic mediators and an summarisation of their mutation based on CScape-somatic assessment, Level of Consequences and total number of mutations. If a gene as previously been assessed as a driver in Network of Cancer Genes (7.0), it is annotated in a separate column.

Cscape_somatic_output.rda The file contain the cscape-somatic assessment for every mutation found in the differentially expressed genes. It is formatted exactly as the output of cscape-somatic, as if it was run in the terminal, except it is saved as .rda instead of csv.
**Examples**

```r
dMA(dataMAF = dataMAF,
   dataDEGs = DEGsmatrix,
   dataPRA = dataPRA,
   coding_file = "path/css_coding.vcf.gz",
   noncoding_file = "path/css_noncoding.vcf.gz",
   results_folder = "path/results")
```

#If the cscape-somatic file have already been created
```r
cscape_somatic_output <- read.csv("./results/Cscape_somatic_output.csv")
save(cscape_somatic_output, file = ".//results/Cscape_somatic_output.rda")
```

```r
dMA(dataMAF = dataMAF,
   dataDEGs = DEGsmatrix,
   dataPRA = dataPRA,
   runCscape = FALSE,
   results_folder = ".//results")
```

---

**EAGenes**

**Information about genes**

### Description

A matrix containing information about 20038 genes including their gene description, location and family

### Usage

```r
data(EAGenes)
```

### Format

A 20038x5 matrix

### Details

The matrix contains the genes in rows and description, location and family in columns.

### Value

A 20038x5 matrix
EncodePromoters

Description

Experimentally verified promoter sites by J. Michael Cherry, Stanford. Downloaded from the ENCODE identifier ENCSR294YNI. It contains chromosome, start and end sites of promoters.

Usage

data(EncodePromoters)

Format

A tibble with no columnnames or rownames.

1. The first column is chromosome eg. chr1
2. The second column is start position eg. 10451
3. The third column is end position eg. 10563

Value

A 84738x6 table

Source

https://www.encodeproject.org/

References

ENCODE identifier: ENCSR294YNI

EpiMix_getInfiniumAnnotation


Usage

EpiMix_getInfiniumAnnotation(plat = "EPIC", genome = "hg38")

Arguments

plat A character string representing the methylation platform which can either be HM27, HM450 or EPIC

genome A character string representing the genome build version which can either be hg19 or hg38

Value

Probe annotations

EpiMix_Results_Regular

Output example from GMA function

Description

The object, a list, that was returned from running the EpiMix function and is one of the outputs from the GMA function.

Usage

data(EpiMix_Results_Regular)
### Format

A list of length nine

### Details

The data is a list of nine elements which is outputted from the EpiMix function

### Value

A list of length nine

<table>
<thead>
<tr>
<th>FEA</th>
<th>FEA</th>
</tr>
</thead>
</table>

### Description

This function carries out the functional enrichment analysis (FEA)

### Usage

```
FEA(BPname = NULL, DEGsmatrix)
```

### Arguments

- **BPname**: BPname biological process such as "proliferation of cells", "ALL" (default) if FEA should be carried out for all 101 biological processes
- **DEGsmatrix**: DEGsmatrix output from DEA such as dataDEGs

### Value

matrix from FEA

### Examples

```r
data(DEGsmatrix)
data(DiseaseList)
data(EAGenes)
DEGsmatrix <- DEGsmatrix[seq.int(2), ]
dataFEA <- FEA(DEGsmatrix = DEGsmatrix, BPname = "apoptosis")
```
**GEO_TCGA**

---

**GEO_TCGA**

*Information on GEO and TCGA data*

---

**Description**

A matrix that provides the GEO dataset matched to one of 18 TCGA cancer types.

**Usage**

```r
data(GEO_TCGA)
```

**Format**

A 18x12 matrix

**Details**

The matrix contains the cancer types in rows and information about sample type from both TCGA and GEO in columns.

**Value**

A 18x12 matrix

---

**getDataGEO**

---

**Description**

This function retrieves and prepares GEO data.

**Usage**

```r
getDataGEO(GEOobject = "GSE39004", platform = "GPL6244", TCGAtumor = NULL)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GEOobject</td>
<td>GEO object</td>
</tr>
<tr>
<td>platform</td>
<td>platform</td>
</tr>
<tr>
<td>TCGAtumor</td>
<td>tumor name</td>
</tr>
</tbody>
</table>

**Value**

return GEO gset
GLS

This function carries out gene literature search.

Description

GLS This function carries out gene literature search.

Usage

GLS(genes, query_string = "AND cancer AND driver", max_records = 20)

Arguments

genes A character string containing the genes to search in PubMed database
query_string A character string containing words in query to follow the gene of interest. Default is "AND cancer AND driver" resulting in a final query of "Gene AND cancer AND driver". Standard PubMed syntax can be used in the query. For example Boolean operators AND, OR, NOT can be applied and tags such as [AU], [TITLE/ABSTRACT], [Affiliation] can be used.
max_records An integer containing the maximum number of records to be fetched from PubMed.

Value

A tibble containing results of literature search where PubMed was queried for information of input genes. Each row in the tibble contains a PubMed ID matching the query, doi, title, abstract, year of publication, keywords, and total number of PubMed publications, resulting in a total of eight columns.

Examples

genes_query <- "BRCA1"
dataGLS <- GLS(genes = genes_query,
query_string = "AND cancer AND driver",
max_records = 2)
GMA

This function carries out Gene Methylation Analysis

Description

GMA This function carries out Gene Methylation Analysis

Usage

GMA(
  dataMET,
  dataEXP,
  dataPRA,
  dataDEGs,
  sample_info,
  met_platform = "HM450",
  prevalence_filter = NULL,
  output_dir = "/GMAresults",
  cores = 1,
  roadmap.epigenome.ids = NULL,
  roadmap.epigenome.groups = NULL
)

Arguments

dataMET A data matrix containing the methylation data where the CpG probes are in the rows and samples are in the columns

dataEXP A data matrix containing the gene expression data where the genes are in the rows and the samples are in the columns

dataPRA A table containing the output of the PRA function

dataDEGs A table containing the output of a DEA where gene names are rownames

sample_info A table containing information on the samples. This table needs to contain two columns called primary and sample.type. The primary column contains sample names which should be the same as the column names in dataMET. The sample.type column indicates for each sample if it is a Cancer or Normal sample.

met_platform A character string representing the microarray type that was used to collect the methylation data. This can either be HM27, HM450 or EPIC. Default is HM450.

prevalence_filter A float or NULL representing if a prevalence filter should be applied or not. Default is NULL, meaning a prevalence filter will not be applied. If a float is specified, a prevalence filter will be applied where methylation states of probes will be altered depending on the threshold of prevalence supplied as prevalence_filter. For example, if prevalence_filter = 20, it means that if the prevalence of the hyper- or hypomethylated CpG probe exceeds 20, the methylation state will be unchanged but if the prevalence is lower than 20 the methylation state will be
changed to NA, meaning no methylation state was detected. In case of dual methylated probes, the methylation state will stay dual if both the prevalence of hyper- and hypomethylations exceed 20, but if only one of the prevalences exceed 20 the dual state will be changed to the state exceeding 20. If none of the prevalences exceed 20, the dual state will be changed to NA.

output_dir  Path to where the results will be stored. If this directory does not exist, it will be created by the function. Default is ./GMAresults.

cores  Number of cores to be used. Default is 1.

roadmap.epigenome.ids  A character string representing the epigenome ID that will be used to select enhancers. Since enhancers are tissue-specific, the tissue type needs to be specified in EpiMix. The enhancers are found from the RoadmapEpigenome project and the IDs can be found from Figure 2 in the publication with doi: 10.1038/nature14248. Default is NULL.

roadmap.epigenome.groups  A character string representing the epigenome group that will be used to select enhancers. Details are provided above with the roadmap.epigenome.ids parameter. Default is NULL.

Value

List of two elements, containing predicted oncogenes and tumor suppressors. Additionally, various output files are saved in the specified output directory: DEG_Methylation_Annotations.rda, Oncogenic_mediators_methylation_summary.rda, EpiMix_Results_Enhancer.rds, EpiMix_Results-Regular.rds, FunctionalPairs_Enhancer.csv, FunctionalPairs-Regular.csv, FunctionalProbes-Regular.rds

Examples

data("dataMethyl")
data("dataFilt")
data("dataPRA")
data("DEGsmatrix")
data("LUAD_sample_anno")
data("NCG")
data("EncodePromoters")
data("MetEvidenceDriver")
pattern <- "^\.(4)\.(2)\.(4)\.(2)\.*"
colnames(dataFilt) <- sub(pattern, "\\", colnames(dataFilt))
dataGMA <- GMA(dataMET = dataMethyl, dataEXP = dataFilt,
dataPRA = dataPRA, dataDEGs = DEGsmatrix,
sample_info = LUAD_sample_anno, met_platform = "HM450",
prevalence_filter = NULL,
output_dir = "./GMAresults", cores = 1, roadmap.epigenome.ids = "E096",
roadmap.epigenome.groups = NULL)
**Description**

This function carries out the gene regulatory network inference using parmigene

**Usage**

```r
GRN(
    TFs,
    DEGsmatrix,
    DiffGenes = FALSE,
    normCounts,
    kNearest = 3,
    nGenesPerm = 2000,
    nBoot = 400,
    noise_mi = 1e-12
)
```

**Arguments**

- **TFs**
  a vector of genes.
- **DEGsmatrix**
  DEGsmatrix output from DEA such as dataDEGs
- **DiffGenes**
  if TRUE consider only diff.expr genes in GRN
- **normCounts**
  is a matrix of gene expression with genes in rows and samples in columns.
- **kNearest**
  the number of nearest neighbors to consider to estimate the mutual information. Must be less than the number of columns of normCounts.
- **nGenesPerm**
  nGenesPerm
- **nBoot**
  nBoot
- **noise_mi**
  noise in knnmi.cross function. Default is 1e-12.

**Value**

an adjacent matrix

**Examples**

```r
data('DEGsmatrix')
data('dataFilt')
dataGRN <- GRN(TFs = sample(rownames(DEGsmatrix), 30),
               DEGsmatrix = DEGsmatrix,
               DiffGenes = TRUE,
               normCounts = dataFilt,
               nGenesPerm = 2,
               nBoot = 2)
```
GSEA

Description
This function carries out the GSEA enrichment analysis.

Usage
GSEA(DEGsmatrix, top, plot = FALSE)

Arguments
- DEGsmatrix: DEGsmatrix output from DEA such as dataDEGs
- top: is the number of top BP to plot
- plot: if TRUE return a GSEA's plot

Value
return GSEA result

Examples
data("DEGsmatrix")
DEGsmatrix_example <- DEGsmatrix[1:2,]
dataFEA <- GSEA(DEGsmatrix = DEGsmatrix_example)

knownDriverGenes

Description
A list of known cancer driver genes from COSMIC

Usage
data(knownDriverGenes)

Format
A list containing two elements where the first element is a character vector of 55 and the second element is a character vector of #84

Details
The list contains two elements: a vector of known tumor # suppressors and a vector of known oncogenes
### LiftMAF

**Value**

A list containing two elements where the first element is a character vector of 55 and the second element is a character vector of # 84

**Description**

This function lifts a MAF file to a different genomic build.

**Usage**

```r
LiftMAF(Infile, Current_Build)
```

**Arguments**

- **Infile**: A tibble of MAF.
- **Current_Build**: A character string, either GRCh38 or GRCh37

**Value**

MAF tibble with positions lifted to another build

**Examples**

```r
data(dataMAF)
dataMAF_example <- dataMAF[1,]
LiftMAF(dataMAF_example, Current_Build = 'GRCh38')
```

### listMoonlight

*A list of oncogenic mediators of 5 TCGA cancer types*

**Description**

A list of oncogenic mediators of 5 TCGA cancer types: BLCA, BRCA, LUAD, READ and STAD

**Usage**

```r
data(listMoonlight)
```

**Format**

A list containing 5 elements where each element contains differentially expressed genes and output from the URA and PRA functions of 5 TCGA cancer types
Details

Each element in the list contains differentially expressed genes and output from the URA and PRA functions.

Value

A list containing 5 elements where each element contains differentially expressed genes and output from the URA and PRA functions of 5 TCGA cancer types.

<table>
<thead>
<tr>
<th>LOC_protein</th>
<th>Level of Consequence: Protein</th>
</tr>
</thead>
</table>

Description

A dataset binary dataset describing if a mutation of a certain class and type possibly have an effect on protein structure or function.

Usage

data(LOC_protein)

Format

A 18x7 table

Details

The values are binary: 0 no effect is possible, 1 an effect is possible.

See supplementary material for details.

Value

A 18x7 table

References

paper
**LOC_transcription**  
*Level of Consequence: Transcription*

**Description**  
A dataset describing if a mutation of a certain class and type possibly have an effect on transcript level.

**Usage**  
`data(LOC_transcription)`

**Format**  
A 18x7 table

**Details**  
The values are binary: 0 no effect is possible, 1 an effect is possible.  
See supplementary material for details.

**Value**  
A 18x7 table

**References**  
paper

**LOC_translation**  
*Level of Consequence: Translation*

**Description**  
A dataset describing if a mutation of a certain class and type possibly have an effect on peptide level.

**Usage**  
`data(LOC_translation)`

**Format**  
A 18x7 table
Details

The values are binary: 0 no effect is possible, 1 an effect is possible.
See supplementary material for details.

Value

A 18x7 table

References

paper

Description

This function carries out the literature phenotype analysis (LPA)

Usage

LPA(dataDEGs, BP, BPlist)

Arguments

dataDEGs is output from DEA
BP is biological process
BPlist is list of genes annotated in BP

Value

table with number of pubmed that affects, increase or decrease genes annotated in BP

Examples

data('DEGsmatrix')
data('DiseaseList')
BPselected <- c("apoptosis")
BPAnnotations <- DiseaseList[[match(BPselected, names(DiseaseList))]]$ID
Sample annotations of TCGA-LUAD project

**Description**

A matrix that annotates LUAD samples as either cancer or normal

**Usage**

```r
data(LUAD_sample_anno)
```

**Format**

A 23x2 matrix

**Details**

The matrix contains two columns: "primary" which contains patient barcodes of TCGA-LUAD and "sample.type" which denotes if the sample is either a "Cancer" or "Normal" sample

**Value**

A 23x2 matrix

---

**MAFtoCscape**

**Description**

This function extracts columns from a MAF tibble to fit CScape input format

**Usage**

```r
MAFtoCscape(MAF)
```

**Arguments**

- `MAF` tibble of MAF

**Value**

tibble of cscape-somatic input

**Examples**

```r
data(dataMAF)
MAFtoCscape(dataMAF[seq.int(2),])
```
MetEvidenceDriver  

*Methylation evidence table to define driver genes*

**Description**

A tibble containing combinations of methylation states of probes used to define driver genes.

**Usage**

```r
data(MetEvidenceDriver)
```

**Format**

A 30x6 tibble.

**Details**

The tibble contains a value of 1 if a probe is found that is either hypo-, hyper-, dualmethylated or not methylated. This is compared with Moonlight’s predictions of role of oncogenic mediators to define driver genes based on methylation evidence.

**Value**

A 30x6 tibble.

---

moonlight  

*moonlight pipeline*

**Description**

moonlight is a tool for identification of cancer driver genes. This function wraps the different steps of the complete analysis workflow.

**Usage**

```r
moonlight(
  dataDEGs,
  dataFilt,
  BPname = NULL,
  Genelist = NULL,
  kNearest = 3,
  nGenesPerm = 2000,
  DiffGenes = FALSE,
  nBoot = 400,
  nTF = NULL,
  thres.role = 0,
```
moonlight

```r
  dataMAF,
  path_cscape_coding,
  path_cscape_noncoding
)

Arguments

dataDEGs    table of differentially expressed genes
dataFilt     matrix of gene expression data with genes in rows and samples in columns
BPname      biological processes to use, if NULL: all processes will be used in analysis, RF for candidate; if not NULL the candidates for these processes will be determined (no learning)
Genelist    Genelist
kNearest    kNearest
nGenesPerm  nGenesPerm
DiffGenes   DiffGenes
nBoot       nBoot
nTF          nTF
thres.role  thres.role
dataMAF      A MAF file rda object for DMA
path_cscape_coding    A character string to path of CScape-somatic coding file
path_cscape_noncoding    A character string to path of CScape-somatic non-coding file

Value

table with cancer driver genes TSG and OCG.

Examples

drivers <- moonlight(dataDEGs = DEGsmatrix,
dataFilt = dataFilt,
BPname = c("apoptosis", "proliferation of cells"),
dataMAF = dataMAF,
path_cscape_coding = "css_coding.vcf.gz",
path_cscape_noncoding = "css_noncoding.vcf.gz")
```
**Description**

A dataset retrieved from Network of Cancer Genes 7.0

**Usage**

`data(NCG)`

**Format**

The format have been rearranged from the original. `|<NCG_driver>|<NCG_cgc_annotation>|<NCG_vogelstein_annotation>|<NCG_saito_annotation>|<NCG_pubmed_id>`

**Details**

The NCG_driver is reported as a OCG or TSG when at least one of three three databases have documented it. These are cosmic gene census (cgc), vogelstein et al. 2013 or saito et al. 2020. The NCG_driver is reported as a candidate, when literature support the gene as a cancer driver.

**Value**

A 3347x7 table

**Source**

[http://ncg.kcl.ac.uk/](http://ncg.kcl.ac.uk/)

**References**

Oncogenic mediators methylation summary

Output example from GMA function

Description
Output file from running the GMA function which is a summary of the oncogenic mediators and their sum of methylated CpG probes together with the evidence level of their role as driver gene.

Usage
data(Oncogenic_mediators_methylation_summary)

Format
A 25x7 tibble

Details
The data is a table where each row is an oncogenic mediator and the columns represent the predicted driver role and the sum of hypo-, hyper-, and dualmethylated CpG sites.

Value
A 25x7 tibble

Oncogenic mediators mutation summary

Oncogenic Mediators Mutation Summary

Description
Output from DMA. This contains the oncogenic mediator from the TCGA-LUAD project, and their mutation asesments summarised based on CSCape-somatic and Level of Consequence.

Usage
data(Oncogenic_mediators_mutation_summary)

Format
A 12x15 matrix.

Value
A 12x15 matrix.
plotCircos

Description

This function visualize the plotCircos

Usage

plotCircos(
  listMoonlight,
  listMutation = NULL,
  additionalFilename = NULL,
  intensityColOCG = 0.5,
  intensityColTSG = 0.5,
  intensityColDual = 0.5,
  fontSize = 1
)

Arguments

listMoonlight  output Moonlight function
listMutation   listMutation
additionalFilename
  additionalFilename
intensityColOCG
  intensityColOCG
intensityColTSG
  intensityColTSG
intensityColDual
  intensityColDual
fontSize       fontSize

Value

no return value, plot is saved

Examples

data('listMoonlight')
plotCircos(listMoonlight = listMoonlight, additionalFilename = "_ncancer5")
plotDMA

Description

This function creates one or more heatmaps on the output from DMA. It visualises the CScape-Somatic annotations per oncogenic mediator either in a single heatmap or split into several different ones. It is also possible to provide a personalised genelist to visualise.

Usage

```r
plotDMA(
  DEG_Mutations_Annotations,
  Oncogenic_mediators_mutation_summary,
  type = "split",
  genelist = c(),
  additionalFilename = ""
)
```

Arguments

- **DEG_Mutations_Annotations**
  A tibble, output file from DMA.

- **Oncogenic_mediators_mutation_summary**
  A tibble, output file from DMA.

- **type**
  A character string. It can take the values "split" or "complete". If both type and genelist are NULL, the function will default to "split".
  - "split" will split the entire dataset into sections of 40 genes and create individual plots. These plots will be merged into one pdf. The genes will be sorted alphabetically.
  - "complete" will create one plot, though it will not be possible to see the individual gene names. The heatmap will be clustered hierarchically.

- **genelist**
  A character vector containing HUGO symbols. A single heatmap will be created with only these genes. The heatmap will be hierarchically clustered. This will overwrite type.

- **additionalFilename**
  A character string. Adds prefix or filepath to the filename of the pdf.

Value

No return value. DMA results are plotted.
Examples

data('DEG_Mutations_Annotations')
data('Oncogenic_mediators_mutation_summary')
plotDMAP(DEG_Mutations_Annotations,
        Oncogenic_mediators_mutation_summary,
        genelist = c("ACSS2", "AFAP1L1"), additionalFilename = "myplots_")

Description

This function visualize the functional enrichment analysis (FEA)'s barplot

Usage

plotFEA(
  dataFEA,
  topBP = 10,
  additionalFilename = NULL,
  height,
  width,
  offsetValue = 5,
  angle = 90,
  xleg = 35,
  yleg = 5,
  titleMain = "",
  minY = -5,
  maxY = 10,
  mycols = c("#8DD3C7", "#FFFFB3", "#BEABA")
)

Arguments

dataFEA
  dataFEA
topBP
  topBP
additionalFilename
  additionalFilename
height
  Figure height
width
  Figure width
offsetValue
  offsetValue
angle
  angle
xleg
  xleg
yleg
  yleg
plotGMA

This function plots results of the Gene Methylation Analysis. It visualizes the number of hyp/hy/hy dual methylated CpG sites in oncogenic mediators or in a user-supplied gene list. The results are visualized either in a single heatmap or split into different ones which is specified in the function's three modes: split, complete and genelist.

Description

plotGMA This function plots results of the Gene Methylation Analysis. It visualizes the number of hypo/hyper/dual methylated CpG sites in oncogenic mediators or in a user-supplied gene list. The results are visualized either in a single heatmap or split into different ones which is specified in the function's three modes: split, complete and genelist.

Usage

plotGMA(
  DEG_Methylation_Annotations,
  Oncogenic_mediators_methylation_summary,
  type = "split",
  genelist = NULL,
  additionalFilename = ""
)

Arguments

DEG_Methylation_Annotations
  A tibble which is outputted from the GMA function.

Oncogenic_mediators_methylation_summary
  A tibble which is outputted from the GMA function.

Value

no return value, FEA result is plotted

Examples

data(DEGsmatrix)
data(DiseaseList)
data(EAGenes)
data(dataFEA)
plotFEA(dataFEA = dataFEA[1:10,], additionalFilename = "_example", height = 20, width = 10)
**plotHeatmap**

- **type**
  A character string which can either be split, complete or genelist. If type is set to split, the entire dataset is split into groups of 40 genes and individual heatmaps of groups each containing 40 genes will be created and subsequently merged into one pdf where each page in the pdf is an individual heatmap. The genes will be sorted alphabetically. If type is set to complete, a single heatmap is created where the number of differentially methylated CpG sites are shown for all oncogenic mediators. If type is set to genelist, a single heatmap will be created for genes supplied by the user in the genelist parameter. Default is split.

- **genelist**
  A character string containing HUGO symbols of genes to be visualized in a single heatmap. Default is NULL.

- **additionalFilename**
  A character string that can be used to add a prefix or filepath to the filename of the pdf visualizing the heatmap. Default is an empty string.

**Value**

No value is returned. Visualizations in form of heatmaps are saved.

**Examples**

```r
data("DEG_Methylation_Annotations")
data("Oncogenic_mediators_methylation_summary")
genes <- c("ACAN", "ACE2", "ADAM19", "AFAP1L1")
plotGMA(DEG_Methylation_Annotations = DEG_Methylation_Annotations,
        Oncogenic_mediators_methylation_summary = Oncogenic_mediators_methylation_summary,
        type = "genelist", genelist = genes,
        additionalFilename = "/GMAresults/")
```

---

**Description**

This function creates a unclustered heatmap from the inputted data tibble and saves it

**Usage**

```r
plotHeatmap(df)
```

**Arguments**

- **df**
  a tibble

**Value**

The name of the alphabetically first gene in the tibble
**plotMetExp**

This function visualizes results of EpiMix.

**Usage**

```r
plotMetExp(
  EpiMix_results, 
  probe_name, 
  dataMET, 
  dataEXP, 
  gene_of_interest, 
  additionalFilename = "" 
)
```

**Arguments**

- `EpiMix_results` The object, a list, that was returned from running the EpiMix function and is one of the outputs from the GMA function.
- `probe_name` A character string containing the name of the CpG probe that will be plotted.
- `dataMET` A data matrix containing the methylation data where the CpG probes are in the rows and samples are in the columns.
- `dataEXP` A data matrix containing the gene expression data where the genes are in the rows and the samples are in the columns.
- `gene_of_interest` A character string containing the name of the gene that will be plotted.
- `additionalFilename` A character string that can be used to add a prefix or filepath to the filename of the pdf visualizing the heatmap. Default is an empty string.

**Value**

No value is returned. Visualizations are saved.

**Examples**

```r
data("EpiMix_Results_Regular")
data("dataMethyl")
data("dataFilt")
pattern <- "^(.{4}-.{2}-.{4}-.{2}).*"
colnames(dataFilt) <- sub(pattern, "\\1", colnames(dataFilt))
plotMetExp(EpiMix_results = EpiMix_Results_Regular, 
  probe_name = "cg03035704", 
  dataMET = dataMethyl, 
  dataEXP = dataFilt, 
  gene_of_interest = "CG03035704", 
  additionalFilename = "EpiMix_Results_Regular.png")
```
```
dataEXP = dataFilt,  
gen_of_interest = "ACVRL1",  
additionalFilename = "/GMAresults/")
```

---

**Description**

This function creates a heatmap of Moonlight gene z-scores for selected genes.

**Usage**

```r
plotMoonlight(
  DEG_Mutations_Annotations,
  Oncogenic_mediators_mutation_summary,
  dataURA,
  gene_type = "drivers",
  n = 50,
  genelist = c(),
  BPlist = c(),
  additionalFilename = ""
)
```

**Arguments**

- **DEG_Mutations_Annotations**
  
  A tibble, output file from DMA.

- **Oncogenic_mediators_mutation_summary**
  
  A tibble, output file from DMA.

- **dataURA**
  
  Output URA function.

- **gene_type**
  
  A character string either "mediators" or "drivers".
  
  - If NULL defaults to "drivers".
  
  - "mediators" will show the oncogenic mediators with the highest number of mutations regardless of driver/passenger classification.
  
  - "drivers" will show the driver genes with the highest number of driver mutations.

- **n**
  
  Numeric. The top number of genes to be plotted. If NULL defaults to 50.

- **genelist**
  
  A vector of strings containing Hugo Symbols of genes. Overwrites gene_type argument.

- **BPlist**
  
  A vector of strings. Selection of the biological processes to visualise. If left empty defaults to every BP provided in the URA file.

- **additionalFilename**
  
  A character string. Adds prefix or filepath to the filename of the pdf.
plotMoonlightMet

Value

No return value. Moonlight scores are plotted for selected genes.

Examples

data(DEG_Mutations_Annotations)
data(Oncogenic_mediators_mutation_summary)
data(dataURA_plot)
plotMoonlight(DEG_Mutations_Annotations,
Oncogenic_mediators_mutation_summary,
dataURA_plot, genelist = c("AFAP1L1", "ABCG2"),
additionalFilename = "myplot_")

plotMoonlightMet  plotMoonlightMet  This function visualizes the effect of genes on biological processes and total number of hypo/hyper/dual methylated CpG sites in genes.

Description

plotMoonlightMet This function visualizes the effect of genes on biological processes and total number of hypo/hyper/dual methylated CpG sites in genes.

Usage

plotMoonlightMet(
  DEG_Methylation_Annotations,
  Oncogenic_mediators_methylation_summary,
dataURA,
  genes,
  additionalFilename = ""
)

Arguments

DEG_Methylation_Annotations
  A tibble which is outputted from the GMA function.
Oncogenic_mediators_methylation_summary
  A tibble which is outputted from the GMA function.
dataURA
  Output of the URA function: a table containing the effect of oncogenic mediators on biological processes. This effect is quantified through Moonlight Gene Z-scores.
genesis
  A character string containing the genes to be visualized.
additionalFilename
  A character string that can be used to add a prefix or filepath to the filename of the pdf visualizing the heatmap. Default is an empty string.
plotNetworkHive

Value

No value is returned. Visualization in form of a heatmap is saved.

Examples

```r
data("DEG_Methylation_Annotations")
data("Oncogenic_mediators_methylation_summary")
data("dataURA_plot")
genes <- c("ACAN", "ACE2", "ADAM19", "AFAP1L1")
plotMoonlightMet(DEG_Methylation_Annotations = DEG_Methylation_Annotations,
    Oncogenic_mediators_methylation_summary = Oncogenic_mediators_methylation_summary,
    dataURA = dataURA_plot,
    genes = genes,
    additionalFilename = "./GMAresults/")
```

plotNetworkHive

plotNetworkHive: Hive network plot

Description

This function visualizes the GRN as a hive plot

Usage

```
plotNetworkHive(dataGRN, namesGenes, thres, additionalFilename = NULL)
```

Arguments

- `dataGRN`: output GRN function
- `namesGenes`: list TSG and OCG to define axes
- `thres`: threshold of edges to be included
- `additionalFilename`: additionalFilename

Value

no results Hive plot is executed

Examples

```r
data(knownDriverGenes)
data(dataGRN)
plotNetworkHive(dataGRN = dataGRN, namesGenes = knownDriverGenes, thres = 0.55)
```
**plotURA**  

*plotURA: Upstream regulatory analysis heatmap plot*

---

**Description**

This function visualizes the URA in a heatmap

**Usage**

```r
plotURA(dataURA, additionalFilename = "URAplot")
```

**Arguments**

- `dataURA`: output URA function
- `additionalFilename`: figure name

**Value**

heatmap

**Examples**

```r
data(dataURA)
data(DiseaseList)
data(tabGrowBlock)
data(knowDriverGenes)
dataDual <- PRA(dataURA = dataURA, 
    BPname = c("apoptosis","proliferation of cells"),
    thres.role = 0)
TSGs_genes <- names(dataDual$TSG)
OCGs_genes <- names(dataDual$OCG)
plotURA(dataURA = dataURA[c(TSGs_genes, OCGs_genes),], additionalFilename = ".example")
```

---

**PRA**  

*Pattern Recognition Analysis (PRA)*

---

**Description**

This function carries out the pattern recognition analysis

**Usage**

```r
PRA(dataURA, BPname, thres.role = 0)
```
Arguments

- `dataURA` (output URA function)
- `BPname` (BPname)
- `thres.role` (thres.role)

Value

returns list of TSGs and OCGs when biological processes are provided, otherwise a randomForest based classifier that can be used on new data

Examples

```r
data(dataURA)
data(DiseaseList)
data(tabGrowBlock)
data(knownDriverGenes)
dataPRA <- PRA(dataURA = dataURA[,1:2], BPname = c("apoptosis","proliferation of cells"), thres.role = 0)
```

---

**PRAtoTibble**

**Description**

This function changes the PRA output to tibble format

**Usage**

```r
PRAtoTibble(dataPRA)
```

**Arguments**

- `dataPRA` (RDA object (list of two) from PRA)

**Value**

tibble with drivers

**Examples**

```r
data('dataPRA') PRAtoTibble(dataPRA)
```
**RunCscape_somatic**

**Description**
This function retrieve cscape-scores to SNPs

**Usage**
RunCscape_somatic(input, coding_file, noncoding_file)

**Arguments**
- `input`: Input matching cscape input
- `coding_file`: cscape_table with coding scores
- `noncoding_file`: cscape_table with noncoding scores

**Value**
returns a tibble with a score and remark for each SNP

**Examples**
```r
cscape_out <- RunCscape_somatic(input, coding_file, noncoding_file)
```

---

**tabGrowBlock**

**Information of growing/blocking characteristics of 101 biological processes**

**Description**
A matrix with biological processes in rows and the cancer growing or blocking effect of the process in columns

**Usage**
data(tabGrowBlock)

**Format**
A 101x3 matrix

**Details**
For each biological processes the cancer growing/blocking effect is indicated
Value

A 101x3 matrix

Description

This function retrieves the individual score for a SNP

Usage

tabix_func(Ranges, Reference_Alele, Mutant, file_coding, file_noncoding)

Arguments

Ranges The position
Reference_Alele The reference nucleotide
Mutant The mutant nucleotide
file_coding cescape_table with coding scores
file_noncoding cescape_table with noncoding scores

Value

returns the score

Examples

data <- tabix_func(Ranges, Reference_Alele, Mutant, file_coding, file_noncoding)

URA

URA Upstream Regulator Analysis

Description

This function carries out the upstream regulator analysis

Usage

URA(dataGRN, DEGsmatrix, BPname, nCores = 1)
Arguments

- **dataGRN**: output GNR function
- **DEGsmatrix**: output DPA function
- **BPname**: biological processes
- **nCores**: number of cores to use

Value

- an adjacent matrix

Examples

```r
data(DEGsmatrix)
dataDEGs <- DEGsmatrix
data(dataGRN)
data(DiseaseList)
data(EAGenes)
dataURA <- URA(dataGRN = dataGRN,
DEGsmatrix = dataDEGs,
BPname = c("apoptosis",
"proliferation of cells")))
```
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