Package ‘padma’

March 9, 2024

Type Package
Title Individualized Multi-Omic Pathway Deviation Scores Using Multiple Factor Analysis
Version 1.12.0
Depends R (>= 4.1.0), SummarizedExperiment, S4Vectors
Imports FactoMineR, MultiAssayExperiment, methods, graphics, stats, utils
Suggests testthat, BiocStyle, knitr, rmarkdown, KEGGREST, missMDA, ggplot2, ggrepel, car, cowplot
Description Use multiple factor analysis to calculate individualized pathway-centric scores of deviation with respect to the sampled population based on multi-omic assays (e.g., RNA-seq, copy number alterations, methylation, etc). Graphical and numerical outputs are provided to identify highly aberrant individuals for a particular pathway of interest, as well as the gene and omics drivers of aberrant multi-omic profiles.
License GPL (>=3)
URL https://github.com/andreamrau/padma
https://github.com/andreamrau/padma/issues
biocViews Software, StatisticalMethod, PrincipalComponent, GeneExpression, Pathways, RNASeq, BioCarta, MethylSeq
Encoding UTF-8
LazyData TRUE
RoxygenNote 7.2.1
VignetteBuilder knitr
git_url https://git.bioconductor.org/packages/padma
git_branch RELEASE_3_18
git_last_commit d925ab1
git_last_commit_date 2023-10-24
Repository Bioconductor 3.18
Date/Publication 2024-03-08
Author Andrea Rau [cre, aut] (<https://orcid.org/0000-0001-6469-488X>),
Regina Manansala [aut],
Florence Jaffrézic [ctb],
Denis Laloë [aut],
Paul Auer [aut]

Maintainer Andrea Rau <andrea.rau@inrae.fr>

R topics documented:

  * factorMap .................................................. 2
  * LUAD_subset ............................................... 4
  * mirtarbase .................................................. 5
  * msigdb ........................................................ 5
  * omicsContribution .......................................... 6
  * padma .......................................................... 7
  * padmaResults-class ......................................... 10
  * padmaRun ...................................................... 11
  * pathway_name ............................................... 13
  * summary.pdmaResults-method ................................. 15

Index 17

---

factorMap

Plot an MFA factor map for individuals or partial factor map based on padma analysis

Description

Produce an MFA factor map for individuals, or MFA partial factor map for a given individual, for a pair of dimensions provided by the user.

Usage

```r
factorMap(
  padma_obj,
  partial_id = NULL,
  dim_x = 1,
  dim_y = 2,
  plot_ellipse = TRUE,
  ggplot = TRUE,
  repel_labels = ifelse(ggplot == TRUE, TRUE, FALSE)
)
```
factorMap

Arguments

padma_obj Output from running the padma function (with 'full_results = TRUE')
partial_id Index or sample name to be plotted for a partial factor map.
dim_x Dimension number of the MFA to be plotted on the x-axis.
dim_y Dimension number of the MFA to be plotted on the y-axis.
pplot_ellipse If TRUE, superimpose a normal confidence ellipsis on the factor map.
ggplot If TRUE, use ggplot2 for plotting
repe_labels If TRUE, use ggrepel to repel sample labels from each other

Value

Plot, or factor map of class ggplot if ggplot2 = TRUE.

Examples

LUAD_subset <- padma::LUAD_subset
## Create MultiAssayExperiment object with LUAD data
omics_data <-
  list(rnaseq = as.matrix(LUAD_subset$rnaseq),
       methyl = as.matrix(LUAD_subset$methyl),
       mirna = as.matrix(LUAD_subset$mirna),
       cna = as.matrix(LUAD_subset$cna))
pheno_data <-
data.frame(LUAD_subset$clinical,
            row.names = LUAD_subset$clinical$bcr_patient_barcode)
mae <- suppressMessages(
  MultiAssayExperiment::MultiAssayExperiment(
    experiments = omics_data, colData = pheno_data))

## Run padma
run_padma <-
padma(mae, gene_map = padma::mirtarbase,
      pathway_name = "c2-cp_BIOCARTA_D4GDI_PATHWAY", verbose = FALSE)

summary(run_padma)

## padma plots
## Not run:
factorMap(run_padma, dim_x = 1, dim_y = 2)
factorMap(run_padma, dim_x = 1, dim_y = 2,
          partial_id = "TCGA-78-7536")
omicsContrib(run_padma, max_dim = 10)

## End(Not run)
## LUAD_subset

Subset of batch-corrected multi-omic TCGA data in lung adenocarcinoma

### Description

List of multi-omic (RNA-seq, copy number alterations, methylation, and miRNA-seq) and phenotypic data in 144 individuals in the TCGA-LUAD data for the 13 genes in the D4-GDI signaling pathway.

### Usage

```r
data(LUAD_subset)
```

### Format

A named list of five objects of class `data.frame` containing a subset of the batch-corrected multi-omic TCGA data from lung adenocarcinoma, corresponding to the 13 genes in the D4 GDI signaling pathway. `clinical` is of dimension 144 x 55 and contains clinical variables for the 144 individuals. `rnaseq`, `methyl`, `cna`, and `mirna` are of dimension 13 (genes) or 1 (miRNAs) x 144 (samples), where the row names contain the gene symbol or miRNA name.

### Source

TCGA

### References


### Examples

```r
data(LUAD_subset)
head(LUAD_subset)
```
**mirtarbase**  
**Curated miR-target interaction predictions from miRTarBase**

**Description**
Data frame of 10,754 predicted miRNA gene targets from miRTarBase (version 7.0), filtered to include only predictions with the 'Functional MTI' support type.

**Usage**
```r
data(mirtarbase)
```

**Format**
An object of class `data.frame` with two columns: `miRNA`, which provides the miRNA identifier (e.g., 'hsa-miR-20a-5p') and `Target Gene`, which provides the corresponding predicted gene target.

**Source**

**References**

**Examples**
```r
data(mirtarbase)
head(mirtarbase)
```

---

**msigdb**  
**MSigDB canonical pathways and corresponding gene lists**

**Description**
Data frame of 1322 pathways and corresponding gene symbols included in the MSigDB canonical pathways curated gene set catalog, which includes genes whose products are involved in metabolic and signaling pathways reported in curated public databases. This specifically corresponds to the 'C2 curated gene sets' catalog from MSigDB v5.2 available at http://bioinf.wehi.edu.au/software/MSigDB/ as described in the limma Bioconductor package.

**Usage**
```r
data(msigdb)
```
Format

An object of class `data.frame` with two columns: `geneset`, which provides the 1322 MSigDB curated pathway names (e.g., 'c2_cp_BIOCARTA_41BB_PATHWAY') and `symbol`, which provides the comma-separated corresponding list of gene symbols.

Source

MSigDB Gene sets http://bioinf.wehi.edu.au/software/MSigDB/

References


Examples

data(msigdb)
head(msigdb)

| omicsContrib | Plot the omics contribution per MFA axis and the overall weighted contribution |

Description

Plot barplots indicating the percent contribution of each omics to each MFA dimension, as well as the overall weighted (by eigenvalue) percent contribution to the full analysis.

Usage

```r
omicsContrib(
  padma_obj,
  max_dim = min(10, nrow(MFA_results(padma_obj)$eig)),
  ggplot = TRUE
)
```

Arguments

- `padma_obj`: Output from running the `padma` function (with 'full_results = TRUE')
- `max_dim`: Maximum dimension number of the MFA to be plotted
- `ggplot`: If TRUE, use `ggplot2` for plotting (and `cowplot` for combining `ggplots`)

Value

Barplots of percent variance contribution, optionally of class `ggplot`. 

Examples

```r
LUAD_subset <- padma::LUAD_subset
## Create MultiAssayExperiment object with LUAD data
omics_data <-
  list(rnaseq = as.matrix(LUAD_subset$rnaseq),
       methyl = as.matrix(LUAD_subset$methyl),
       mirna = as.matrix(LUAD_subset$mirna),
       cna = as.matrix(LUAD_subset$cna))
pheno_data <-
  data.frame(LUAD_subset$clinical,
             row.names = LUAD_subset$clinical$bcr_patient_barcode)
mae <- suppressMessages(
  MultiAssayExperiment::MultiAssayExperiment(
    experiments = omics_data, colData = pheno_data))

## Run padma
run_padma <-
  padma(mae, gene_map = padma::mirtarbase,
        pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY", verbose = FALSE)
summary(run_padma)

## padma plots
## Not run:
factorMap(run_padma, dim_x = 1, dim_y = 2)
factorMap(run_padma, dim_x = 1, dim_y = 2,
          partial_id = "TCGA-78-7536")
omicsContrib(run_padma, max_dim = 10)
## End(Not run)
```

Description

This is the primary user interface for the `padma` package. Generic S4 methods are implemented to calculate individualized pathway deviation scores on the basis of matched, multi-omic data. The supported classes for input are `list` and `MultiAssayExperiment`. The output of `padma` is an S4 object of class `padmaResults`.

Usage

```r
padma(object, ...)
```

## S4 method for signature 'list'
```r
padma(
  object,
```
colData,  
gene_map = padma::mirtarbase,  
base_ids = NULL,  
supp_ids = NULL,  
pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY",  
impute = FALSE,  
variance_threshold = 1e-04,  
full_results = TRUE,  
verbose = TRUE,  
...  
)  

## S4 method for signature 'MultiAssayExperiment'  

padma(  
  object,  
  gene_map = padma::mirtarbase,  
  base_ids = NULL,  
  supp_ids = NULL,  
  pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY",  
impute = FALSE,  
  variance_threshold = 1e-04,  
  full_results = TRUE,  
  verbose = TRUE,  
...  
)  

Arguments

- **object**: Matched multi-omic data. May be provided as (1) a `MultiAssayExperiment` or (2) a named list, with each element corresponding to a matrix representing an omic, with biological entities in rows. Row names should include unique biological entity IDs (e.g., gene symbols, miRNA names); columns represent individuals. If more than one biological entity is used, a `gene_map` data.frame providing mappings between IDs and gene names should be provided if the default mirtarbase is not sufficient.

- **...**: Optional additional arguments

- **colData**: (optional) A `DataFrame` or `data.frame` of characteristics for all biological units, to be used in creating a `MultiAssayExperiment` from an object of class `list`.

- **gene_map**: (optional) Data frame mapping arbitrary biological entities (e.g. miRNAs) to genes. Contains two columns, where the first provides the IDs of the entity and the second provides the IDs of the corresponding target gene. By default, the miRNA-gene interactions of type 'Functional MTI' from miRTarBase are used (see the preloaded 'mirtarbase' data in the package).

- **base_ids**: (optional) Sample names to be used as reference base data. By default, all samples are used.

- **supp_ids**: (optional) Sample names to be used as supplementary individuals to be projected onto the analysis based on the individuals identified in `base_ids`. By default,
takes the value NULL, but should not overlap with base_ids if provided by the user.

pathway_name Character of either a KEGG pathway identifier or MSigDB pathway names (e.g., see the pathway names in the 'geneset' column of the preloaded msigdb data in the package), or a vector of gene symbols.

impute If TRUE, impute missing values separately in base and supplementary data using MFA as implemented in the missMDA package; otherwise simple mean imputation is used (default).

variance_threshold Minimal variance required across samples to retain a biological entity in the analysis

full_results If TRUE (default), include full MFA results in function output; otherwise, provide concise output to save space.

verbose If TRUE, provide verbose output.

Value

An S4 object of class padmaResults, where individualized pathway deviation scores are stored as the assay data, and the corresponding pathway name, full MFA results, number of genes, and names of imputed or filtered genes are stored as slots that can be retrieved using the appropriate accessor functions.

Author(s)

Andrea Rau

Examples

LUAD_subset <- padma::LUAD_subset
## Create MultiAssayExperiment object with LUAD data
omics_data <-
  list(rnaseq = as.matrix(LUAD_subset$rnaseq),
       methyl = as.matrix(LUAD_subset$methyl),
       mirna = as.matrix(LUAD_subset$mirna),
       cna = as.matrix(LUAD_subset$cna))
pheno_data <-
  data.frame(LUAD_subset$clinical,
             row.names = LUAD_subset$clinical$bcr_patient_barcode)
mae <-
  suppressMessages(
    MultiAssayExperiment::MultiAssayExperiment(,
      experiments = omics_data, colData = pheno_data))

## Run padma
run_padma <-
  padma(mae, gene_map = padma::mirtarbase,
        pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY", verbose = FALSE)

summary(run_padma)
padmaResults-class

**Description**

padmaResults is a subclass of RangedSummarizedExperiment, used to store the individualized pathway deviation scores as well as some additional information useful about the pathway name (pathway_name), the gene-level contributions to each deviation score (pathway_gene_deviation), a full set of outputs related to the MFA (MFA_results, and the number of genes used in the analysis as well as the names of those for which data imputation or filtering was required (ngenes, imputed_genes, and removed_genes, respectively).

**Usage**

```r
padmaResults(
  SummarizedExperiment,
  pathway_name = NULL,
  pathway_gene_deviation = NULL,
  MFA_results = NULL,
  ngenes = NULL,
  imputed_genes = NULL,
  removed_genes = NULL
)
```

**Arguments**

- **SummarizedExperiment**
  a RangedSummarizedExperiment of padma results
- **pathway_name**
  The name of the pathway, if applicable
- **pathway_gene_deviation**
  Per-gene contributions to each individualized pathway deviation score
- **MFA_results**
  List of all detailed results from the MFA
- **ngenes**
  Number of genes used in the pathway deviation score calculation
- **imputed_genes**
  Names of genes, per omic, for which data imputation was used to replace missing values
- **removed_genes**
  Names of genes, per omic, which were filtered from the analysis due to low variation
Details

This constructor function would not typically be used by 'end users'. This simple class extends the RangedSummarizedExperiment class of the SummarizedExperiment package to allow other packages to write methods for results objects from the padma package. It is used by padmaRun to wrap up the results table.

Value

a padmaResults object

padmaRun

Calculate individualized deviation scores from multi-omic data

Description

Calculate individualized deviation scores from multi-omic data

Usage

padmaRun(
  omics_data,
  gene_map = padma::mirtarbase,
  base_ids = NULL,
  supp_ids = NULL,
  pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY",
  impute = FALSE,
  variance_threshold = 1e-04,
  full_results = TRUE,
  verbose = TRUE,
  ...
)

Arguments

omics_data Object of class 'MultiAssayExperiment' containing omics data from n matched individuals.
gene_map (optional) Data frame mapping arbitrary biological entities (e.g. miRNAs) to genes. Contains two columns, where the first provides the IDs of the entity and the second provides the IDs of the corresponding target gene. By default, the miRNA-gene interactions of type 'Functional MTI' from miRTarBase are used (see the preloaded 'mirtarbase' data in the package).
base_ids (optional) Sample names to be used as reference base data. By default, all samples are used.
supp_ids (optional) Sample names to be used as supplementary individuals to be projected onto the analysis based on the individuals identified in base_ids. By default, takes the value NULL, but should not overlap with base_ids if provided by the user.
pathway_name Character of either a KEGG pathway identifier or MSigDB pathway names (e.g., see the pathway names in the 'geneset' column of the preloaded msigdb data in the package), or a vector of gene symbols.

impute If TRUE, impute missing values separately in base and supplementary data using MFA as implemented in the missMDA package; otherwise simple mean imputation is used (default).

variance_threshold Minimal variance required across samples to retain a biological entity in the analysis.

full_results If TRUE (default), include full MFA results in function output; otherwise, provide concise output to save space.

verbose If TRUE, provide verbose output.

Value An S4 object of class padmaResults, where individualized pathway deviation scores are stored as the assay data, and the corresponding pathway name, full MFA results, number of genes, and names of imputed or filtered genes are stored as slots that can be retrieved using the appropriate accessor functions.

Examples

LUAD_subset <- padma::LUAD_subset
## Create MultiAssayExperiment object with LUAD data
omics_data <-
  list(rnaseq = as.matrix(LUAD_subset$rnaseq),
       methyl = as.matrix(LUAD_subset$methyl),
       mirna = as.matrix(LUAD_subset$mirna),
       cna = as.matrix(LUAD_subset$cna))
pheno_data <-
  data.frame(LUAD_subset$clinical,
             row.names = LUAD_subset$clinical$bcr_patient_barcode)

mae <-
  suppressMessages(
    MultiAssayExperiment::MultiAssayExperiment(
      experiments = omics_data, colData = pheno_data))

## Run padma
run_padma <-
  padma(mae, gene_map = padma::mirtarbase,
        pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY", verbose = FALSE)

summary(run_padma)

## padma plots
## Not run:
factorMap(run_padma, dim_x = 1, dim_y = 2)
factorMap(run_padma, dim_x = 1, dim_y = 2,
          partial_id = "TCGA-78-7536")
omicsContribution(run_padma, max_dim = 10)

## End(Not run)

Accessors for a padmaResults object.

**Description**

Accessors for a padmaResults object.

**Usage**

```r
pathway_name(object, ...)
pathway_gene_deviation(object, ...)
MFA_results(object, ...)
ngenes(object, ...)
imputed_genes(object, ...)
removed_genes(object, ...)
```

```r
## S4 method for signature 'padmaResults'
pathway_name(object)
## S4 method for signature 'padmaResults'
MFA_results(object)
## S4 method for signature 'padmaResults'
ngenes(object)
## S4 method for signature 'padmaResults'
imputed_genes(object)
## S4 method for signature 'padmaResults'
removed_genes(object)
## S4 method for signature 'padmaResults'
pathway_gene_deviation(object)
## S4 method for signature 'padmaResults'
show(object)
```
Arguments

object a padmaResults object

... Additional optional parameters

Value

Output varies depending on the method.

Author(s)

Andrea Rau

Examples

LUAD_subset <- padma::LUAD_subset
## Create MultiAssayExperiment object with LUAD data
omics_data <-
  list(rnaseq = as.matrix(LUAD_subset$rnaseq),
       methyl = as.matrix(LUAD_subset$methyl),
       mirna = as.matrix(LUAD_subset$mirna),
       cna = as.matrix(LUAD_subset$cna))
pheno_data <-
data.frame(LUAD_subset$clinical,
           row.names = LUAD_subset$clinical$bcr_patient_barcode)
mae <-
suppressMessages(
  MultiAssayExperiment::MultiAssayExperiment(
    experiments = omics_data, colData = pheno_data))

## Run padma
run_padma <-
  padma(mae, gene_map = padma::mirtarbase,
        pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY", verbose = FALSE)

summary(run_padma)

## padma plots
## Not run:
factorMap(run_padma, dim_x = 1, dim_y = 2)
factorMap(run_padma, dim_x = 1, dim_y = 2,
          partial_id = "TCGA-78-7536")
omicsContrib(run_padma, max_dim = 10)

## End(Not run)
**Summary, padmaResults-method**

*Summarize results from padma*

**Description**

A function to summarize the pathway deviation results from padma, using the quantiles of the calculated multi-omic pathway deviation scores.

**Usage**

```r
## S4 method for signature 'padmaResults'
summary(object, ...)
```

**Arguments**

- `object` An object of class 'padmaResults'
- `...` Additional arguments

**Value**

Summary of the padmaResults object.

**Author(s)**

Andrea Rau

**References**


**See Also**

padma

**Examples**

```r
LUAD_subset <- padma::LUAD_subset
## Create MultiAssayExperiment object with LUAD data
omics_data <-
  list(rnaseq = as.matrix(LUAD_subset$rnaseq),
       methyl = as.matrix(LUAD_subset$methyl),
       mirna = as.matrix(LUAD_subset$mirna),
       cna = as.matrix(LUAD_subset$cna))
pheno_data <-
data.frame(LUAD_subset$clinical,
            row.names = LUAD_subset$clinical$bcr_patient_barcode)
```
mae <-
suppressMessages(
  MultiAssayExperiment::MultiAssayExperiment(
    experiments = omics_data, colData = pheno_data))

## Run padma
run_padma <-
  padma(mae, gene_map = padma::mirtarbase,
        pathway_name = "c2_cp_BIOCARTA_D4GDI_PATHWAY", verbose = FALSE)

summary(run_padma)

## padma plots
## Not run:
factorMap(run_padma, dim_x = 1, dim_y = 2)
factorMap(run_padma, dim_x = 1, dim_y = 2,
          partial_id = "TCGA-78-7536")
omicsContrib(run_padma, max_dim = 10)

## End(Not run)
## Index

**datasets**
- LUAD_subset, 4
- mirtarbase, 5
- msigdb, 5

**methods**
- padma, 7
- summary, padmaResults-method, 15

- factorMap, 2

- imputed_genes (pathway_name), 13
- imputed_genes, padmaResults-method (pathway_name), 13

- LUAD_subset, 4

- MFA_results (pathway_name), 13
- MFA_results, padmaResults-method (pathway_name), 13
- mirtarbase, 5
- msigdb, 5

- ngenes (pathway_name), 13
- ngenes, padmaResults-method (pathway_name), 13

- omicsContrib, 6

- padma, 7, 15
- padma, list-method (padma), 7
- padma, MultiAssayExperiment-method (padma), 7
- padma-methods (padma), 7
- padmaResults (padmaResults-class), 10
- padmaResults-class, 10
- padmaRun, 11, 11
- pathway_gene_deviation (pathway_name), 13
- pathway_gene_deviation, padmaResults-method (pathway_name), 13

- removed_genes (pathway_name), 13
- removed_genes, padmaResults-method (pathway_name), 13

- show (pathway_name), 13
- show, padmaResults-method (pathway_name), 13
- summary (summary, padmaResults-method), 15
- summary, padmaResults-method, 15
- summary-methods (summary, padmaResults-method), 15