Package ‘orthos’

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Type Package

Title `orthos` is an R package for variance decomposition using conditional variational auto-encoders

Version 1.2.0

Description `orthos` decomposes RNA-seq contrasts, for example obtained from a gene knock-out or compound treatment experiment, into unspecific and experiment-specific components. Original and decomposed contrasts can be efficiently queried against a large database of contrasts (derived from ARCHS4, https://maayanlab.cloud/archs4/) to identify similar experiments. `orthos` furthermore provides plotting functions to visualize the results of such a search for similar contrasts.

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Depends R (>= 4.3), SummarizedExperiment

Imports AnnotationHub, basilisk, BiocParallel, colorspace, cowplot, DelayedArray, dplyr, ExperimentHub, ggplot2, ggpubr, ggrepel, ggsci, grDevices, grid, HDF5Array, keras, methods, orthosData, parallel, ptyr, reticulate, rlang, S4Vectors, stats, tensorflow, tidyr

Suggests BiocManager, BiocStyle, htmltools, knitr, rmarkdown, testthat (>= 3.0.0)

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biocViews RNASeq, DifferentialExpression, GeneExpression

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Description

orthos decomposes RNA-seq contrasts, for example obtained from a genetic or compound treatment experiments, into non-specific and experiment-specific components. Original and decomposed contrasts can be efficiently queried against a large database of public contrasts (derived from ARCHS4, https://maayanlab.cloud/archs4/) to identify similar experiments. orthos furthermore provides plotting functions to visualize the results of such queries for similar contrasts.

Author(s)

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decomposeVar | Decompose input contrasts to decoded and residual fractions

Description
Decompose input contrasts (gene expression deltas) to decoded (generic) and residual (unique) components according to a contrast encoder-decoder pre-trained on a large corpus of public RNAseq experiments.

Usage
```
decomposeVar(
  M,  # Matrix of raw gene counts.
  MD = NULL,  # Matrix of gene deltas (optional). If MD is specified, M is assumed
treatm = NULL,  # to be a raw gene count matrix specifying context for contrasts
  cntr = NULL,  # specified in MD. MD is then a matrix of gene deltas with the same
  processInput = TRUE,  # dimensions as M. If MD is specified, treatm
  organism = c("Human", "Mouse"),  # and cntr have to be NULL.
  featureType = c("AUTO", "ENSEMBL_GENE_ID", "GENE_SYMBOL", "ENTREZ_GENE_ID",  
                  "ARCHS4_ID"),  
  pseudocount = 4,  # Vectors indicating column indices in M corresponding to
t  verbose = TRUE  # treatments and controls. If treatm and cntr are specified, MD has to be
)
```

Arguments

- **M**: Matrix of raw gene counts.
- **MD**: Matrix of gene deltas (optional). If MD is specified, M is assumed to be a raw gene count matrix specifying context for contrasts specified in MD. MD is then a matrix of gene deltas with the same dimensions as M. If MD is specified, treatm and cntr have to be NULL.
- **treatm, cntr**: Vectors indicating column indices in M corresponding to treatments and controls. If treatm and cntr are specified, MD has to be NULL.
- **processInput**: If set to TRUE (default) the count matrix will be preprocessed (library normalized, log2-transformed after addition of a pseudocount, NA values will be set to 0).
- **organism**: Selects the autoencoder model trained on data from this species. One of "Human" or "Mouse".
- **featureType**: Set to "AUTO" for automatic feature id-type detection. Alternatively specify the type of supplied id features. Current supported types are "ENSEMBL_GENE_ID", "GENE_SYMBOL", "ENTREZ_GENE_ID" and "ARCHS4_ID".
- **pseudocount**: Numerical scalar, added to raw counts in M when preprocessInput = TRUE.
- **verbose**: Logical scalar indicating whether to print messages along the way.
Details

When calling `decomposeVar()`, you may see an `ImportError` on the console. This most likely does not have any negative consequences, rather it means that R and python may not be library compatible and that an automated fallback approach is being used (for more details see `testload` argument of `basiliskStart`).

Value

A `SummarizedExperiment` object with the decomposed contrasts in the assays and the decomposed variance as the `colData`.

Author(s)

Panagiotis Papasaikas

Examples

```r
MKL1_human <- readRDS(system.file("extdata", "GSE215150_MKL1_Human.rds", package = "orthos"))

# Specifying M, treatm and cntr:
dec_MKL1_human <- decomposeVar(M = MKL1_human, treatm = c(2, 3), cntr = c(1, 1), organism = "Human", verbose = FALSE)

# Alternatively by specifying M and MD:
pseudocount <- 4
M <- sweep(MKL1_human, 2, colSums(MKL1_human), FUN = "/") * 1e+06
M <- log2(M + pseudocount)
DeltaM <- M[,c("MKL1","caMKL1")]-M[,"Ctrl"] # Matrix of contrasts
ContextM <- M[,c("Ctrl","Ctrl")]
context <- colnames(ContextM) <- colnames(DeltaM) # M and MD need identical dimnames
RES <- decomposeVar(M = ContextM, MD = DeltaM, processInput = FALSE)
```

---

**loadContrastDatabase**  
*Load contrast database*

**Description**

Load a pre-calculated, organism-specific contrast database and return it as a `SummarizedExperiment`.

**Usage**

```r
loadContrastDatabase(
  organism = c("Human", "Mouse"),
  mode = c("ANALYSIS", "DEMO"),
  mustWork = TRUE
)
```
**plotQueryResultsManh**

**Arguments**

- **organism**
  Character scalar selecting the organism for which to load the contrast database. One of "Human" or "Mouse".

- **mode**
  When in "ANALYSIS" mode (default) the complete contrast DB is queried. "DEMO" mode employs a small "toy" database for the queries. "DEMO" should only be used for testing/demonstration purposes and never for actual analysis purposes.

- **mustWork**
  Logical scalar. If FALSE and the contrast database is not available, return an empty SummarizedExperiment object. If TRUE (the default) and the contrast database is not available, loadContrastDatabase throws an error.

**Details**

Organism-specific databases are compiled in HDF5SummarizedExperiment objects. The first time `loadContrastDatabase()` is called for a database either directly or via `queryWithContrasts()` the required objects will be automatically downloaded from `ExperimentHub` and cached in the user ExperimentHub directory (see `ExperimentHub::getExperimentHubOption("CACHE")`) using the 'orthosData' companion data-package.

**Value**

A SummarizedExperiment with pre-calculated contrasts as assays.

**Author(s)**

Panagiotis Papasaikas, Michael Stadler

**Examples**

```r
# !!!Note!!! mode="DEMO" for demonstration purposes only. Default is mode="ANALYSIS"
SE_mouse_demoDB <- loadContrastDatabase (organism="Mouse", mode="DEMO")
SE_mouse_demoDB

SE_human_demoDB <- loadContrastDatabase (organism="Human", mode="DEMO")
SE_human_demoDB
```

---

**plotQueryResultsManh** Visualize query results as a composite manhattan/density plot.

**Description**

Visualize query results as a composite manhattan/density plot.

**Usage**

`plotQueryResultsManh(queryResults, doPlot = TRUE)`
plotQueryResultsViolin

Visualize query results as violin plots

Description

Visualize query results as violin plots

Usage

plotQueryResultsViolin(queryResults, doPlot = TRUE)
**queryWithContrasts**

**Query the contrast database with a set of contrasts**

**Description**

Query the contrast database with a set of contrasts

**Usage**

```r
queryWithContrasts(
  contrasts,  
  use = c("expressed.in.both", "all.genes"),  
  exprThr = 0.25,  
  organism = c("Human", "Mouse"),
)```

**Arguments**

- `queryResults` A list containing the results of a query performed with `queryWithContrasts`
- `doPlot` Logical scalar specifying if a plot should be generated.

**Value**

A list of ggplot violin plots (one for each dataset) for the scores of queries using different contrast components against the respective contrast DBs.

**Author(s)**

Panagiotis Papasaikas, Michael Stadler

**Examples**

```r
MKL1_human <- readRDS(system.file("extdata", "GSE215150_MKL1_Human.rds", package = "orthos"))

# Decompose contrasts:
dec_MKL1_human <- decomposeVar(M = MKL1_human, treatm = c(2, 3), cntr = c(1, 1),
                                 organism = "Human", verbose = FALSE)

# Perform query against contrast DB with the decomposed fractions.
# !!!Note!!! mode="DEMO" for demonstration purposes only.
params <- BiocParallel::MulticoreParam(workers = 2)
query.res.human <- queryWithContrasts(dec_MKL1_human, organism = "Human",
                                      BPPARAM = params, verbose = FALSE,
                                      mode = "DEMO")

# plot results for individual contrasts using violin plots::
ViolinPlots <- plotQueryResultsViolin(query.res.human, doPlot = FALSE)
ViolinPlots[["caMKL1"]]
```
plotType = c("violin", "manh", "none"),
detailTopn = 10,
verbose = TRUE,
BPPARAM = BiocParallel::bpparam(),
chunk_size = 500,
mode = c("ANALYSIS", "DEMO")
)

Arguments

contrasts A SummarizedExperiment object with assays containing contrasts named INPUT_CONTRASTS, DECODED_CONTRASTS and RESIDUAL_CONTRASTS (at least one should be present) and context information in an assay named CONTEXT. The latter is only required when use="expressed.in.both". This is typically generated using decomposeVar.

use Determines if all.genes or genes expressed in both query and target context will be used. Note that "expressed.in.both", though more accurate, is slower.

exprThr is the quantile in the provided context that determines the expression value above which a gene is considered to be expressed. This same value is then used for thresholding the contrast database. Only applies when use="expressed.in.both".

organism Uses the 'orthosData' contrast database from this species. One of "Human" or "Mouse".

plotType Select the type of visualization for the query results "violin", "manh" or "none" to suppress the plotting.

detailTopn specifies the number of top hits for which metadata will be returned in the TopHits slot of the results.

verbose Logical scalar indicating whether to print messages along the way.

BPPARAM BiocParallelParam object specifying how parallelization is to be performed using e.g. MulticoreParam or SnowParam).

chunk_size Column dimension for the grid used to read blocks from the HDF5 Matrix. Sizes between 250 and 1000 are recommended. Smaller sizes reduce memory usage.

mode When in "ANALYSIS" mode (default) the complete contrast DB is queried. "DEMO" mode employs a small "toy" database for the queries. "DEMO" should only be used for testing/demonstration purposes and never for actual analysis purposes.

Value

A list with three elements called "pearson.rhos", "zscores" and "TopHits", containing raw and z-scored Pearson’s rho correlation coefficients between the query contrast(s) and the contrasts in the database, as well as detailed metadata for the detailTopn best hits.

Author(s)

Panagiotis Papasaikas
Ex
ts

Examples

MKL1_human <- readRDS(system.file("extdata", "GSE215150_MKL1_Human.rds", package = "orthos"))

# Decompose contrasts:
dec_MKL1_human <- decomposeVar(M = MKL1_human, treatm = c(2, 3), cntr = c(1, 1),
                                   organism = "Human", verbose = FALSE)

# Perform query against contrast DB with the decomposed fractions.
# !!!Note!!! mode="DEMO" for demonstration purposes only.
params <- BiocParallel::MulticoreParam(workers = 2)
query.res.human <- queryWithContrasts(dec_MKL1_human, organism = "Human",
                                       BPPARAM = params, verbose = FALSE,
                                       mode = "DEMO")

testOrthosEnv

Test conda environment

Description

Test conda environment

Usage

testOrthosEnv()

Value

A list indicating whether keras is available, and the version of TensorFlow.

Author(s)

Charlotte Soneson

Examples

testOrthosEnv()
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