

Package ‘phantasus’

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Title Visual and interactive gene expression analysis

Version 1.30.0

Description Phantasus is a web-application for visual and interactive gene expression analysis. Phantasus is based on Morpheus – a web-based software for heatmap visualisation and analysis, which was integrated with an R environment via OpenCPU API. Aside from basic visualization and filtering methods, R-based methods such as k-means clustering, principal component analysis or differential expression analysis with limma package are supported.

URL <https://alserglab.wustl.edu/phantasus>

BugReports <https://github.com/ctlab/phantasus/issues>

Depends R (>= 4.3)

biocViews GeneExpression, GUI, Visualization, DataRepresentation, Transcriptomics, RNASeq, Microarray, Normalization, Clustering, DifferentialExpression, PrincipalComponent, ImmunoOncology

Imports ggplot2, protolite, Biobase, GEOquery, Rook, htmltools, httpuv, jsonlite, limma, edgeR, opencpu, assertthat, methods, httr, rhdf5, utils, parallel, stringr, fgsea (>= 1.9.4), svglite, gtable, stats, Matrix, pheatmap, scales, ccaPP, grid, grDevices, AnnotationDbi, DESeq2, data.table, curl, apeglm, tidyr, config (>= 0.3.2), rhdf5client (>= 1.25.1), yaml, fs, phantasusLite, XML

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adjustDataset	<i>Adjust dataset</i>
---------------	-----------------------

Description

Adjust dataset

Usage

```
adjustDataset(
  es,
  scaleColumnSum = NULL,
  log2 = FALSE,
  onePlusLog2 = FALSE,
  inverseLog2 = FALSE,
  quantileNormalize = FALSE,
  zScore = FALSE,
  robustZScore = FALSE,
  sweep = NULL
)
```

Arguments

es	Expression set to perform adjustment on
scaleColumnSum	perform sum scaling of columns (default FALSE)
log2	perform logarithm2 adjustment (default FALSE)
onePlusLog2	perform $\log_2(1+x)$ adjustment (default FALSE)
inverseLog2	perform 2^x adjustment (default FALSE)
quantileNormalize	perform quantile normalization (default FALSE)
zScore	perform zScore adjustment: subtract mean, divide by std (default FALSE)
robustZScore	perform robustZScore adjustment: subtract median, divide by MAD (default FALSE)
sweep	perform sweep adjustment on rows/columns (default FALSE)

Value

Nothing. Adjusted dataset will be assigned as ES in global environment

Examples

```
## Not run:
es <- gseGSE('GSE53986')[[1]]
adjustDataset(es, log2 = T, quantileNormalize = T)

## End(Not run)
```

annotationDBMeta	<i>Create meta file for AnnotationDB</i>
------------------	--

Description

annotationDBMeta function creates txt files containing meta information of provided sqlite files for AnnotationDB.

Usage

```
annotationDBMeta(annotDir)
```

Arguments

annotDir	path to folder with annotationDB sqlite files
----------	---

Value

nothing

Examples

```
## Not run:  
annotationDBMeta('/var/phantasus/cache')  
  
## End(Not run)
```

calcPCA	<i>Principal Component Analysis.</i>
---------	--------------------------------------

Description

calcPCA calculates PCA-matrix for the given ExpressionSet and returns this matrix encoded to JSON.

Usage

```
calcPCA(es, replacena = "mean")
```

Arguments

es	an ExpressionSet object, should be normalized
replacena	method for replacing NA values (mean by default)

Value

json with full description of the plot for plotly.js

Examples

```
## Not run:  
data(es)  
calcPCA(es)  
  
## End(Not run)
```

calculatedAnnotation *Create calculated annotation*

Description

calculatedAnnotation adds a column calculated by operation

Usage

```
calculatedAnnotation(  
  es,  
  operation,  
  rows = c(),  
  columns = c(),  
  isColumns = FALSE,  
  name = NULL  
)
```

Arguments

es	ExpressionSet object.
operation	Name of the operation to perform calculation
rows	List of specified rows' indices (optional), indices start from 0
columns	List of specified columns' indices (optional), indices start from 0#'
isColumns	Apply fn to columns
name	Name of the new annotation

Value

Nothing. Annotated dataset will be assigned to es in environment

checkGPLsFallback *Check possible annotations for GEO Dataset.*

Description

checkGPLs returns GPL-names for the specified GEO identifier.

Usage

```
checkGPLsFallback(name)
```

Arguments

name String, containing GEO identifier of the dataset.

Value

Vector of filenames serialized in JSON format. If there is only one GPL for that dataset, the function will return name.

Examples

```
## Not run:
checkGPLs('GSE27112')
checkGPLs('GSE14308')

## End(Not run)
```

checkGSEType *Checks GSE to be supported*

Description

Checks GSE to be supported

Usage

```
checkGSEType(name, destDir, combine = any)
```

Arguments

name GSE id, with optional GPL specification
destDir path to cache directory
combine function on how to combine results, when multiple platforms are present

Value

logical vector if the dataset is supported or not

collapseDataset	<i>Collapse dataset</i>
-----------------	-------------------------

Description

collapseDataset performs a collapse action on expression set

Usage

```
collapseDataset(  
  es,  
  isRows = TRUE,  
  selectOne = FALSE,  
  fn,  
  fields,  
  removeEmpty = TRUE  
)
```

Arguments

es	Expression set
isRows	Work with rows. False if columns (default True - row mode)
selectOne	select best match or merge duplicates
fn	select/merge function
fields	fields to unique on
removeEmpty	remove unannotated genes

Value

Nothing. Collapsed dataset will be assigned to es in environment

Examples

```
## Not run:  
es <- getGSE('GSE53986')[[1]]  
collapseDataset(es, isRows = TRUE, selectOne = TRUE,  
  fn = mean, fields = c('Gene ID', 'Gene symbol'))  
  
## End(Not run)
```

colMeansByGroups *Calculate column averages in row groups*

Description

Calculate column averages in row groups

Usage

```
colMeansByGroups(m, groups)
```

Arguments

m	matrix n x m
groups	vector of size n of numbers from 1 to k

Value

matrix k*m of column averages by groups

convertByAnnotationDB *Map indexes using Annotation DB*

Description

convertByAnnotationDB function returns keyType ids from dbName mapped to columnName in es.

Usage

```
convertByAnnotationDB(
  es,
  dbName,
  columnName,
  columnType,
  keyType,
  otherOptions
)
```

Arguments

es	source ExpressionSet
dbName	name of AnnotationDB file
columnName	name of column in featureData of source ExpressionSet
columnType	Type of indexes in columnName
keyType	Type of mapped indexes
otherOptions	additional parameters for conversion. Currently only named boolean delete-DotVersion is not ignored.

Value

JSON object with a vector of converted IDs

createDockerConf	<i>Creates default docker conf file Function creates default docker user configuration file based on provided setup_file or on default parameters if setup_file doesn't exist. If user_conf_file exists function does nothing.</i>
------------------	--

Description

Creates default docker conf file Function creates default docker user configuration file based on provided setup_file or on default parameters if setup_file doesn't exist. If user_conf_file exists function does nothing.

Usage

```
createDockerConf(
  setup_file = confFile("setup.yml"),
  user_conf_file = confFile("user.conf")
)
```

Arguments

setup_file	name of config from file. If unset or not existed, "default".
user_conf_file	Location of the setup.yml file with setup parameters. If not existed use file from package

createES	<i>Create ExpressionSet.</i>
----------	------------------------------

Description

createES function produces an ExpressionSet object from given data, and exports it to global scope.

Usage

```
createES(data, pData, varLabels, fData, fvarLabels, eData)
```

Arguments

data	Gene expression matrix.
pData	Matrix with phenotypical data.
varLabels	Names of phenoData columns.
fData	Matrix with feature data.
fvarLabels	Names of featureData columns.
eData	List with experimentData

Value

produced ExpressionSet object

Examples

```
## Not run:
data <- matrix(1:15, 5, 3)
pData <- c("A", "B", "C")
varLabels <- "cat"
fData <- c("p", "r", "s", "t", "u")
fvarLabels <- "id"
eData <- list(name="", lab="", contact="", title="", url="", other=list(), pubMedIds="")
createES(data, pData, varLabels, fData, fvarLabels, eData)

## End(Not run)
```

es

Example dataset

Description

Small slice from GSE27112-GPL6103 for runnable examples.

Usage

```
data(es)
```

Format

An object of class ExpressionSet with 20 rows and 5 columns.

Examples

```
## Not run:
data(es)
performKmeans(es, k = 2)

## End(Not run)
```

fgseaExample

Example pathway data.frame for fgsea tool

Description

Example pathway data.frame for fgsea tool

```
generatePreloadedSession
```

Generate files for preloaded session from a session link.

Description

Generate files for preloaded session from a session link.

Usage

```
generatePreloadedSession(sessionURL, preloadedName, preloadedDir)
```

Arguments

sessionURL String with session link produced by phantasus.
preloadedName String with name that should be assigned to the session.
preloadedDir Path to the directory with preloaded datasets and sessions.

Value

Function produces two files (preloadedName.rda with ExpressionSet and preloadedName.json with session features) in preloadedDir folder.

Examples

```
## Not run:
sessionURL <- "https://ctlab.itmo.ru/phantasus/?session=x063c1b365b9211" # link from 'Get dataset link...' to
newName <- "my_session" # user defined name
preloadedDir <- "./preloaded" # directory where files will be stored. In order too get access through phantasus
dir.create(preloadedDir, showWarnings = FALSE)
generatePreloadedSession(sessionURL= sessionURL,
                        preloadedName = newName,
                        preloadedDir = preloadedDir)

servePhantasus(preloadedDir=preloadedDir, openInBrowser=FALSE)
# open browser manually at http://0.0.0.0:8000/phantasus/index.html?preloaded=my_session

## End(Not run)
```

```
getArchs4Files
```

Returns list of ARCHS4 hdf5 files with expression data

Description

Returns list of ARCHS4 hdf5 files with expression data

Usage

```
getArchs4Files(cacheDir)
```

Arguments

cacheDir base directory for cache

Value

list of .h5 files

getCountsMetaPart *Create meta-data for single counts collection*

Description

Creates a part of counts collections meta-data

Usage

```
getCountsMetaPart(counts_dir, collection_name, verbose)
```

Arguments

counts_dir path to directory with count collections
collection_name name of collection and collection's directory
verbose logical value which determines a content of the output.

Details

Function assumes that collection_name contains meta.txt which is valid (in sense of [validateCountsCollection](#)). For each row in meta.txt function reads specified sample_id dataset and writes every sample id to the resulting data.table with source file name and collection name.

Value

data.table with meta-data or nothing if destdir does not exist or does not contain files.

See Also

[validateCountsCollection.getCountsMetaPart](#)

Examples

```
## Not run:
collDir <- "/path/to/my/collection"
valid_collection = validateCountsCollection(collectionDir = collDir, verbose = TRUE)
if (valid_collection){
  metaPart = getCountsMetaPart(destdir = collDir, verbose = TRUE)
}

## End(Not run)
```

getES *Load ExpressionSet by GEO identifier*

Description

getES return the ExpressionSet object(s) corresponding to GEO identifier.

Usage

```
getES(  
  name,  
  type = NA,  
  destdir = getPhantasusConf("cache_folders")$geo_path,  
  mirrorPath = getPhantasusConf("geo_mirrors")  
)
```

Arguments

name	String, containing GEO identifier of the dataset. It should start with 'GSE' or 'GDS' and can include exact GPL to annotate dataset, separated with dash ('-') from the identifier.
type	Type of the dataset: 'GSE' or 'GDS'. If not specified, the function will take first three letters of name variable as type.
destdir	Directory for caching loaded Series and GPL files from GEO database.
mirrorPath	URL string which specifies the source of matrices.

Value

List of ExpressionSet objects, that were available by given in name variable GEO identifier.

Examples

```
## Not run:  
  getES('GSE14308', type = 'GSE', destdir = 'cache')  
  getES('GSE27112')  
  getES('GDS4922')  
  
## End(Not run)
```

getGDS *Load ExpressionSet from GEO Datasets*

Description

getGDS return the ExpressionSet object corresponding to GEO Dataset identifier.

Usage

```
getGDS(
  name,
  destdir = getPhantastusConf("cache_folders")$geo_path,
  mirrorPath = getPhantastusConf("geo_mirrors")
)
```

Arguments

name	String, containing GEO identifier of the dataset. It should start with 'GSE' or 'GDS' and can include exact GPL to annotate dataset, separated with dash ('-') from the identifier.
destdir	Directory for caching loaded Series and GPL files from GEO database.
mirrorPath	URL string which specifies the source of matrices.

Value

ExpressionSet object wrapped in list, that was available by given in name variable GEO identifier.

Examples

```
## Not run:
  getGDS('GDS4922', destdir = tempdir(), mirrorPath = "https://ftp.ncbi.nlm.nih.gov")

## End(Not run)
```

getGSE

Load ExpressionSet from GEO Series

Description

getGSE return the ExpressionSet object(s) corresponding to GEO Series Identifier.

Usage

```
getGSE(
  name,
  destdir = getPhantastusConf("cache_folders")$geo_path,
  mirrorPath = getPhantastusConf("geo_mirrors")
)
```

Arguments

name	String, containing GEO identifier of the dataset. It should start with 'GSE' or 'GDS' and can include exact GPL to annotate dataset, separated with dash ('-') from the identifier.
destdir	Directory for caching loaded Series and GPL files from GEO database.
mirrorPath	URL string which specifies the source of matrices.

Value

List of ExpressionSet objects, that were available by given in name variable GEO identifier.

Examples

```
## Not run:
  getGSE('GSE14308', destdir = 'cache')
  getGSE('GSE27112')
  getGSE('GSE53986')

## End(Not run)
```

getPhantasusConf	<i>Read Phantasus Config</i>
------------------	------------------------------

Description

Read Phantasus Config

Usage

```
getPhantasusConf(
  value = NULL,
  configName = Sys.getenv("R_CONFIG_ACTIVE"),
  file = file.path(tools::R_user_dir(package = "phantasus", which = "config"),
    "user.conf")
)
```

Arguments

value	Value to retrieve from the config file.
configName	R_CONFIG_ACTIVE value. If unset, "default".
file	Location of the config file

gseaPlot	<i>Returns path to an svg file with enrichment plot</i>
----------	---

Description

Returns path to an svg file with enrichment plot

Usage

```

gseaPlot(
  es,
  rankBy,
  selectedGenes,
  width,
  height,
  vertical = FALSE,
  addHeatmap = FALSE,
  showAnnotation = NULL,
  annotationColors = NULL,
  pallete = c("blue", "white", "red")
)

```

Arguments

es	ExpressionSet object.
rankBy	name of the numeric column used for gene ranking
selectedGenes	indexes of selected genes (starting from one, in the order of fData)
width	width of the image (in inches)
height	height of the image (in inches)
vertical	whether to use vertical orientation (default: FALSE)
addHeatmap	whether to add an expression heatmap, sorted by rankBy (default: FALSE)
showAnnotation	a name of column annotation to add to the heatmap, default: NULL (no annotation)
annotationColors	a list of colors to use in annotation
pallete	a vector of colors to draw heatmap

Value

path to an svg file

isHSDS	<i>check if url responding as HSDS server TRUE - hsdS FALSE - web link but not working NULL - not web link</i>
--------	--

Description

check if url responding as HSDS server TRUE - hsdS FALSE - web link but not working NULL - not web link

Usage

```
isHSDS(url)
```

Arguments

url	URL to check
-----	--------------

limmaAnalysis *Differential Expression analysis.*

Description

limmaAnalysis performs differential expression analysis from limma package and returns a Protobuf-serialized resulting de-matrix.

Usage

```
limmaAnalysis(  
  es,  
  fieldValues,  
  version = "One-factor design",  
  contrast = list("Comparison", "Target", "Reference"),  
  designData = NULL  
)
```

Arguments

es	ExpressionSet object. It should be normalized for more accurate analysis.
fieldValues	Vector of comparison values, mapping categories' names to columns/samples
version	name of the limma analysis implementation. Should be "One-factor design" or "Advanced design"
contrast	a character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change
designData	data.frame with design matrix

Value

Name of the file containing serialized de-matrix.

Examples

```
## Not run:  
data(es)  
limmaAnalysis(es, fieldValues = c("A", "A", "A", "B", "B"))  
  
## End(Not run)
```

loadCounts	<i>Loads expression data from .h5 count files. Only samples with counted expression are kept. If es already contains expression data it is returned as is.</i>
------------	--

Description

Loads expression data from .h5 count files. Only samples with counted expression are kept. If es already contains expression data it is returned as is.

Usage

```
loadCounts(es, counts_dir)
```

Arguments

es	ExpressionSet from GEO to check for expression in ARCHS4/dee2 or other h5 files
counts_dir	directory with .h5 files collections. There must be meta.rda file in counts_dir and each collection's sub directory must have meta.txt file with description. Also counts_dir must contain counts_priority.txt file.

Value

either original es or an ExpressionSet with loaded count data from ARCHS4

loadFromARCHS4	<i>Loads expression data from ARCHS4 count files. Only samples with counted expression are kept. If es already contains expression data it is returned as is.</i>
----------------	---

Description

Loads expression data from ARCHS4 count files. Only samples with counted expression are kept. If es already contains expression data it is returned as is.

Usage

```
loadFromARCHS4(es, archs4_files)
```

Arguments

es	ExpressionSet from GEO to check for expression in ARCHS4
archs4_files	list of available .h5 files from ARCHS4 project

Value

either original es or an ExpressionSet with loaded count data from ARCHS4

loadGEO	<i>Load GEO Dataset.</i>
---------	--------------------------

Description

loadGEO returns the file with serialized ExpressionSet using ProtoBuf, parsed from data downloaded from GEO by identifier.

Usage

```
loadGEO(name, type = NA)
```

Arguments

name	String, containing GEO identifier of the dataset. It should start with 'GSE' or 'GDS' and can include exact GPL to annotate dataset, separated with dash ('-') from the identifier.
type	Type of the dataset: 'GSE' or 'GDS'. If not specified, the function will take first three letters of name variable as type.

Value

File with ProtoBuf-serialized ExpressionSet-s that were downloaded by this identifier. For GSE-datasets there can be multiple annotations, so in file will be a list mapping name with GPL to ExpressionSet.

Examples

```
## Not run:
  loadGEO("GSE27112")
  loadGEO("GDS4922")

## End(Not run)
```

loadPreloaded	<i>Load GEO Dataset.</i>
---------------	--------------------------

Description

loadPreloaded returns the file with serialized ExpressionSets using ProtoBuf, that were preloaded on server.

Usage

```
loadPreloaded(name)
```

Arguments

name	String, containing filename. Assuming that in the directory with preloaded files preloadedDir exists file filename.rda with list of ExpressionSets ess.
------	---

Value

File with ProtoBuf-serialized ExpressionSet-s that were loaded from specified file.

performKmeans	<i>K-means clusterisation.</i>
---------------	--------------------------------

Description

performKmeans returns a vector of corresponding clusters for each gene from a given ExpressionSet.

Usage

```
performKmeans(es, k, replacena = "mean")
```

Arguments

es	ExpressionSet object.
k	Expected number of clusters.
replacena	Method for replacing NA values in series matrix (mean by default)

Value

Vector of corresponding clusters, serialized to JSON.

Examples

```
## Not run:
data(es)
performKmeans(es, k = 2)

## End(Not run)
```

queryAnnotationDBMeta *Get meta list for annotationDB files*

Description

queryAnnotationDBMeta Function reads txt meta files for provided sqlite annotation databases.

Usage

```
queryAnnotationDBMeta()
```

Value

meta info in JSON

Examples

```
## Not run:
queryAnnotationDBMeta()

## End(Not run)
```

read.gct	<i>Reads ExpressionSet from a GCT file. Function is deprecated, please use phantasusLite:::readGct() instead</i>
----------	--

Description

Reads ExpressionSet from a GCT file. Function is deprecated, please use phantasusLite:::readGct() instead

Usage

```
read.gct(...)
```

Arguments

... parameters for phantasusLite:::readGct() call

Value

ExpressionSet object

reparseCachedESs	<i>Reparse cached expression sets from GEO.</i>
------------------	---

Description

The function should be used on phantasus version updates that change behavior of loading datasets from GEO. It finds all the datasets that were cached and runs 'getES' for them again. The function uses cached Series and other files from GEO.

Usage

```
reparseCachedESs(destdir, mirrorPath = getPhantasusConf("geo_mirrors"))
```

Arguments

destdir Directory used for caching loaded Series files from GEO database.
mirrorPath URL string which specifies the source of matrices.

Value

vector of previously cached GSE IDs

Examples

```
reparseCachedESs(destdir=tempdir(), "https://ftp.ncbi.nlm.nih.gov")
```

reproduceInR	<i>Reproduce session in R code</i>
--------------	------------------------------------

Description

Reproduce session in R code

Usage

```
reproduceInR(sessionName, leaf = T, step = 0, savedEnv = new.env())
```

Arguments

sessionName	String, OCPU session name
leaf	Boolean, is it leaf (default = F)
step	Integer, step of recursion (default = 0)
savedEnv	Environment, where to store complex arguments (default = new.env())

Value

JSON with R code

Examples

```
## Not run:
  setwd(tempdir())
  reproduceInR('x039f1672026678');

## End(Not run)
```

servePhantasus	<i>Serve phantasus.</i>
----------------	-------------------------

Description

servePhantasus starts http server handling phantasus static files and opencpu server.

Usage

```
servePhantasus(
  host = getPhantasusConf("host"),
  port = getPhantasusConf("port"),
  staticRoot = getPhantasusConf("static_root"),
  preloadedDir = getPhantasusConf("preloaded_dir"),
  openInBrowser = TRUE,
  quiet = TRUE,
  background = FALSE
)
```

Arguments

host	Host to listen.
port	Port to listen.
staticRoot	Path to static files with phantasia.js (on local file system).
preloadedDir	Full path to directory with preloaded files.
openInBrowser	Boolean value which states if application will be automatically loaded in default browser.
quiet	Boolean value which states whether the connection log should be hidden (default: TRUE)
background	Boolean value which states whether the server should be started in background (default: FALSE)

Value

A handle to the server as returned by 'httpuv::startServer'

Examples

```
## Not run:
s <- servePhantasia(background=FALSE)
s$stop()

## End(Not run)

httpuv::stopAllServers() # can be used if handle is lost
```

setupPhantasia	<i>Setup phantasia. Read user config file (or create default one) and fill cache_root using sources in file.</i>
----------------	---

Description

Setup phantasia. Read user config file (or create default one) and fill cache_root using sources in file.

Usage

```
setupPhantasia(setup_name = "default", file = confFile("setup.yml"))
```

Arguments

setup_name	name of config from file. If unset or not existed, "default".
file	Location of the setup.yml file with setup parameters. If not existed use file from package

shinyGAMAnalysis	<i>Constructs data frame with gene annotations and submits it into Shiny GAM web-server</i>
------------------	---

Description

Constructs data frame with gene annotations and submits it into Shiny GAM web-server

Usage

```
shinyGAMAnalysis(es)
```

Arguments

es	Expression set object
----	-----------------------

Value

URL for Shiny GAM

subsetES	<i>Subsets es, if rows or columns are not specified, all are retained</i>
----------	---

Description

Subsets es, if rows or columns are not specified, all are retained

Usage

```
subsetES(es, columns = c(), rows = c())
```

Arguments

es	ExpressionSet object.#'
columns	List of specified columns' indices (optional), indices start from 0#'
rows	List of specified rows' indices (optional), indices start from 0

Value

new expression set 'es'

updateARCHS4 *Update archs4 files.*

Description

Download archs4 or archs4zoo counts in cacheDir. If directory does not exist function makes nothing and produce corresponding warnings.

Usage

```
updateARCHS4(
  cacheDir = file.path(getPhantasusConf("cache_folders")$rnaseq_counts, "archs4"),
  organism = c("all"),
  force = FALSE
)
```

Arguments

cacheDir	file path to archs4 cache directory
organism	vector which determines organisms to download: human, mouse, zoo or all as default. Also can be a genus. Possible genus: <ol style="list-style-type: none"> 1. drosophila 2. gallus 3. bos 4. caenorhabditis 5. danio 6. rattus 7. saccharomyces 8. arabidopsis
force	logical value which let function replace current files

updateARCHS4meta *Update ARCHS4 meta files*

Description

Creates meta.txt file, which describes typical archs4 and archs4Zoo files.

Usage

```
updateARCHS4meta(
  archDir = file.path(getPhantasusConf("cache_folders")$rnaseq_counts, "archs4")
)
```

Arguments

archDir	path to directory with arch4 .h5 files.
---------	---

Details

This function produces very specific "hardcoded" meta.txt file for arch4 and archs4ZOO counts collections. See [validateCountsCollection](#) for more common information and meta.txt file structure

See Also

[validateCountsCollection](#)

updateCountsMeta	<i>Update meta-data for counts collections</i>
------------------	--

Description

Creates meta.rda file which contain information about all samples in all collections. Also function checks priority.txt file. This file is used to manage collections with the same samples.

Usage

```
updateCountsMeta(
  counts_dir = getPhantasusConf("cache_folders")$rnaseq_counts,
  force = FALSE,
  verbose = FALSE
)
```

Arguments

counts_dir	path to counts cache directory
force	logical value wich lets function replace existing meta.rda file
verbose	logical value which determines a content of the output.

Details

First of all function checks validity of priority.txt file. **Every** Collection should have **unique** priority. If priority.txt is not valid function creates new one, setting priorities for each subdirectory(=collection) equal to order in list.dir output.

Function updates meta.rda if this file is older than at least one .h5 file in counts files. meta.rda is data.table which is a result of union data.tables produced by [getCountsMetaPart](#) for each collection

See Also

[validateCountsCollection](#), [updateCountsMeta](#)

updateDEE2meta	<i>Update DEE2 meta files</i>
----------------	-------------------------------

Description

Creates meta.txt file, which describes typical dee2 files.

Usage

```
updateDEE2meta(  
  destDir = file.path(getPhantasusConf("cache_folders")$rnaseq_counts, "dee2")  
)
```

Arguments

destDir path to directory with DEE2 .h5 files.

Details

This function produces very specific "hardcoded" meta.txt file for dee2 counts collection. See [validateCountsCollection](#) for more common information and meta.txt file structure

See Also

[validateCountsCollection](#)

validateCountsCollection	<i>Check a counts collection</i>
--------------------------	----------------------------------

Description

Function checks existing and structure of meta.txt file in specified counts folder. Also it checks accessibility of specified datasets in corresponding .h5 files.

Usage

```
validateCountsCollection(collectionDir, verbose = FALSE)
```

Arguments

collectionDir path to directory with collection
verbose logical value which determines a content of the output.

Details

collectionDir should contain a bunch of .h5 files and a single meta.txt. meta.txt is .tsv-like file where for each .h5 exists a row with columns:

file_name name of .h5 file in collectionDir.

sample_id name of dataset in file_name which contains sample IDs (sample_geo_accession for example).

sample_dim which dimension of the expression matrix in file_name corresponds to samples. Should be one of c("rows", "columns")

gene_id name of dataset in file_name which contains ids for genes and the "meaning" for that ids(column name in result ES). For correct work this dataset should contain unique values. Example: ENSEMBLID:/meta/genes/ensembl_gene_id

genes_annot Names of datasets and their meanings to extract gene-related metadata from file_name. Can be empty or gene_id-like values separated with semicolon(;).

write.gct	<i>Saves ExpressionSet to a GCT file (version 1.3). Function is deprecated, please use phantasusLite::writeGct() instead</i>
-----------	--

Description

Saves ExpressionSet to a GCT file (version 1.3). Function is deprecated, please use phantasusLite::writeGct() instead

Usage

```
write.gct(...)
```

Arguments

... parameters for phantasusLite::writeGct() call

Value

Result of the closing file (as in 'close()' function)

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