

Package ‘phantasusLite’

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Type Package

Title Loading and annotation RNA-seq counts matrices

Version 1.4.0

Description PhantasusLite – a lightweight package with helper functions of general interest extracted from phantasus package. In particular it simplifies working with public RNA-seq datasets from GEO by providing access to the remote HSDS repository with the precomputed gene counts from ARCHS4 and DEE2 projects.

Depends R (>= 4.2)

Imports data.table, rhdf5client(>= 1.25.1), httr, stringr, stats, utils, Biobase, methods

biocViews GeneExpression, Transcriptomics, RNASeq

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RoxygenNote 7.3.1

Suggests testthat (>= 3.0.0), knitr, rmarkdown, BiocStyle, rhdf5, GEOquery

VignetteBuilder knitr

Config/testthat/edition 3

URL <https://github.com/ctlab/phantasusLite/>

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

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createH5	<i>Creates metafiles for HDF5-files</i>
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Description

Creates metafiles for HDF5-files

Usage

```
createH5(data, file, dataset_name)
```

Arguments

data	contains metadata
file	contains file name
dataset_name	contains dataset name

Value

Returns NULL

createIndexH5	<i>Writes indexes to the file</i>
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Description

Writes indexes to the file

Usage

```
createIndexH5(data, file)
```

Arguments

data	contains metadata
file	contains the file name

Value

Returns NULL

createIndexH5Remote	<i>Creates HDF5-file with indexes</i>
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Description

Creates HDF5-file with indexes

Usage

```
createIndexH5Remote(  
  url,  
  collections = c("archs4", "dee2"),  
  destfile = "index.h5"  
)
```

Arguments

url	contains URL to the root of counts files
collections	vector of collection names to process
destfile	where to put resulting index file

Value

Returns NULL

createMetaH5	<i>Converts collection meta.txt files to meta.h5, putting them to the respective collection folders</i>
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Description

Converts collection meta.txt files to meta.h5, putting them to the respective collection folders

Usage

```
createMetaH5(counts_dir)
```

Arguments

counts_dir	contains directory name
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Value

Returns NULL

createPriorityH5	<i>Creates HDF5-File with priority</i>
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Description

Creates HDF5-File with priority

Usage

```
createPriorityH5(counts_dir, force = FALSE, verbose = FALSE)
```

Arguments

counts_dir	contains counts directory
force	logical value which lets function replace existing priority file
verbose	logical value which determines a content of the output.

Value

Returns NULL

getCountsMetaPart	<i>Gets list with metadata</i>
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Description

Gets list with metadata

Usage

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

Arguments

counts_dir	contains counts directory
collection_name	contains name of the collection
verbose	logical value which determines a content of the output.

Value

list with metadata

getHSDSFileList	<i>Returns list of all HDF5-files on HSDS-server</i>
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Description

Returns list of all HDF5-files on HSDS-server

Usage

```
getHSDSFileList(  
  url = "https://alserglab.wustl.edu/hsds/?domain=/counts",  
  directory = NULL  
)
```

Arguments

url	containing url of the server and root domain.
directory	containing name of the directory

Value

List of all HDF5-files on the server or all files of the collection

Examples

```
url <- 'https://alserglab.wustl.edu/hsds/?domain=/counts'  
getHSDSFileList(url)
```

getIndexRemote	<i>Creates a data table with indexes and chunks of samples in remote HDF5-files</i>
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Description

Creates a data table with indexes and chunks of samples in remote HDF5-files

Usage

```
getIndexRemote(url, collections)
```

Arguments

url	contains url to the root of counts files
collections	contains names of the collections

Value

table with samples, indexes and chunks in all HDF5-files

gsmToChunk	<i>Gets chunk from GSE identifiers.</i>
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Description

Gets chunk from GSE identifiers.

Usage

```
gsmToChunk(samples)
```

Arguments

samples	containing a list of samples
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Value

list of chunks

inferCondition	<i>Adds condition to the annotation.</i>
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Description

Adds condition to the annotation.

Usage

```
inferCondition(es)
```

Arguments

es contains ExpressionSet object

Value

Annotated ExpressionSet with conditions and replicates

Examples

```
ess <- GEOquery::getGEO("GSE143903")
es <- ess[[1]]
es <- inferCondition(es)
es$condition # contains inferred groups
es$replicate # contains inferred replicate numbers
```

inferConditionImpl	<i>Creates condition from the samples titles</i>
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Description

Creates condition from the samples titles

Usage

```
inferConditionImpl(gse_titles)
```

Arguments

gse_titles contains titles

Value

List of conditions and replicates

loadCountsFromH5FileHSDS

Load count matrix from remote HDF5-file

Description

Load count matrix from remote HDF5-file

Usage

```
loadCountsFromH5FileHSDS(
  es,
  url = "https://alserglab.wustl.edu/hsds/?domain=/counts",
  file,
  sampleIndexes = NULL
)
```

Arguments

es	containing ExpressionSet loaded from GEO. Contains empty expression matrix.
url	containing url of the server and root domain.
file	containing name of the file (relative to the root domain)
sampleIndexes	containing sample indexes list

Value

ExpressionSet object with loaded count matrix

Examples

```
ess <- GEOquery::getGEO("GSE53053")
es <- ess[[1]]
url <- 'https://alserglab.wustl.edu/hsds/?domain=/counts'
file <- "/dee2/mmusculus_star_matrix_20240409.h5"
es <- loadCountsFromH5FileHSDS(es, url, file)
```

loadCountsFromHSDS

Load count matrix from HDF5-files.

Description

Load count matrix from HDF5-files.

Usage

```
loadCountsFromHSDS(
  es,
  url = "https://alserglab.wustl.edu/hsds/?domain=/counts"
)
```


Arguments

`es` containing ExpressionSet loaded from GEO. Contains empty expression matrix.
`url` containing url of the server and root domain.

Value

ExpressionSet with loaded count matrix

Examples

```
ess <- GEOquery::getGEO("GSE85653")
es <- ess[[1]]
url <- 'https://alserglab.wustl.edu/hds/?domain=/counts'
es <- loadCountsFromHSDS(es, url)
```

readGct	<i>Reads ExpressionSet from a GCT file.</i>
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Description

Only versions 1.2 and 1.3 are supported.

Usage

```
readGct(gct)
```

Arguments

`gct` Path to gct file

Value

ExpressionSet object

Examples

```
es <- readGct(system.file("extdata/testdata/gct/test.gct", package="phantasusLite"))
```

removeRepeatWords	<i>Removes repeated words from conditions</i>
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Description

Removes repeated words from conditions

Usage

```
removeRepeatWords(titles)
```

Arguments

titles	contains titles
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Value

titles without repeated words

updateARCHS4meta	<i>Creates meta.txt file, which describes typical archs4 and archs4Zoo files.</i>
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Description

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

Usage

```
updateARCHS4meta(  
  archDir = file.path(getOption("phantasusCacheDir"), "counts/archs4")  
)
```

Arguments

archDir	path to directory with arch4 .h5 files.
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Value

Returns NULL

updateDEE2meta	<i>Creates meta.txt file, which describes typical dee2 files.</i>
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Description

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

Usage

```
updateDEE2meta(
  destDir = file.path(getOption("phantasusCacheDir"), "counts/dee2")
)
```

Arguments

destDir	path to directory with DEE2 .h5 files.
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Value

Returns NULL

updateIndexH5	<i>Updates indexes from HDF5-files</i>
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Description

Updates indexes from HDF5-files

Usage

```
updateIndexH5(counts_dir, force = FALSE, verbose = FALSE)
```

Arguments

counts_dir	contains counts directory
force	logical value which lets function replace existing index file
verbose	logical value which determines a content of the output.

Value

Returns NULL

validateCountsCollection
Validates counts collection

Description

Validates counts collection

Usage

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

Arguments

collectionDir contains directory name
verbose logical value which determines a content of the output.

Value

false if collection is not valid

writeGct *Saves ExpressionSet to a GCT file (version 1.3).*

Description

Saves ExpressionSet to a GCT file (version 1.3).

Usage

```
writeGct(es, file, gzip = FALSE)
```

Arguments

es ExpressionSet object to save
file Path to output gct file
gzip Whether to gzip apply gzip-compression for the output file#'

Value

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

Examples

```
es <- readGct(system.file("extdata/testdata/gct/test.gct", package="phantasusLite"))
out <- tempfile(fileext = ".gct.gz")
writeGct(es, out, gzip=TRUE)
```

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