Package 'AMARETTO'

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Туре	Package
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R to	opics documented:
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	AMARETTO_HTMLreport
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AMARETTO_Download

AMARETTO_Download

Description

Downloading TCGA dataset for AMARETTO analysis

Usage

AMARETTO_Download(CancerSite, TargetDirectory, downloadData = TRUE)

Arguments

CancerSite TCGA cancer code for data download TargetDirectory

Directory path to download data

downloadData

Examples

```
TargetDirectory <- file.path(getwd(),"Downloads/") # path to data download directory
CancerSite <- "LIHC"
DataSetDirectories <- AMARETTO_Download(CancerSite, TargetDirectory)</pre>
```

AMARETTO_EvaluateTestSet

AMARETTO_EvaluateTestSet

Description

Code to evaluate AMARETTO on a new gene expression test set. Uses output from AMARETTO_Run() and CreateRegulatorData().

Usage

```
AMARETTO_EvaluateTestSet(AMARETTOresults = AMARETTOresults,
    MA_Data_TestSet = MA_Data_TestSet,
    RegulatorData_TestSet = RegulatorData_TestSet)
```

Arguments

```
AMARETTOresults
```

AMARETTO output from AMARETTO_Run().

MA_Data_TestSet

Gene expression matrix from a test set (that was not used in AMARETTO_Run()).

RegulatorData_TestSet

Test regulator data from CreateRegulatorData().

Examples

AMARETTO_ExportResults

AMARETTO_ExportResults

Description

Retrieve a download of all the data linked with the run (including heatmaps)

Usage

```
AMARETTO_ExportResults(AMARETTOinit, AMARETTOresults, data_address, Heatmaps = TRUE, CNV_matrix = NULL, MET_matrix = NULL)
```

Arguments

AMARETTO initialize output

AMARETTOresults

AMARETTO results output

data_address Directory to save data folder
Heatmaps Output heatmaps as pdf

Examples

```
AMARETTOresults <- AMARETTO_Run(AMARETTOinit)

AMARETTO_ExportResults(AMARETTOinit, AMARETTOresults, "./")
```

AMARETTO_HTMLreport AMARETTO_HTMLreport

Description

Retrieve an interactive html report, including gene set enrichment analysis if asked for.

Usage

```
AMARETTO_HTMLreport(AMARETTOinit, AMARETTOresults, CNV_matrix = NULL, MET_matrix = NULL, SAMPLE_annotation = NULL, ID = NULL, VarPercentage, hyper_geo_test_bool = FALSE, hyper_geo_reference = NULL, output_address = "./", MSIGDB = FALSE, GMTURL = FALSE)
```

Arguments

AMARETTO initialize output

AMARETTOresults

AMARETTO results output

CNV_matrix Processed CNV matrix (\$CNV_matrix element from Preprocess_CancerSite()

list output).

MET_matrix Processed methylation matrix (\$MET matrix element from Preprocess CancerSite()

list output).

SAMPLE_annotation

SAMPLE annotation will be added to heatmap

ID ID column of the SAMPLE annotation data frame

VarPercentage Original Var Percentage used

hyper_geo_test_bool

Boolean if a hyper geometric test needs to be performed. If TRUE provide a

GMT file in the hyper_geo_reference parameter.

hyper_geo_reference

GMT file with gene sets to compare with.

 $\hbox{\tt output_address} \quad \hbox{\tt Output directory for the html files}.$

MSIGDB TRUE if gene sets were retrieved from MSIGDB. Links will be created in the

report.

GMTURL TRUE if second column of gmt contains URLs to gene set, FALSE if it contains

a description

Examples

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Description

Code used to initialize the seed clusters for an AMARETTO run. Requires processed gene expressiosn (rna-seq or microarray), CNV (usually from a GISTIC run), and methylation (from MethylMix, provided in this package) data. Uses the function CreateRegulatorData() and results are fed into the function AMARETTO_Run().

Usage

```
AMARETTO_Initialize(MA_matrix = MA_matrix, CNV_matrix = NULL, MET_matrix = NULL, Driver_list = NULL, NrModules, VarPercentage, PvalueThreshold = 0.001, RsquareThreshold = 0.1, pmax = 10, NrCores = 1, OneRunStop = 0, method = "union")
```

Arguments

Expression matrix, with genes in rows and samples in columns.
CNV matrix, with genes in rows and samples in columns.
Methylation matrix, with genes in rows and samples in columns.
Custom list of driver genes to be considered in analysis
How many gene co-expression modules should AMARETTO search for? Usually around 100 is acceptable, given the large number of possible driver-passenger gene combinations.
Minimum percentage by variance for filtering of genes; for example, 75% would indicate that the CreateRegulatorData() function only analyses genes that have a variance above the 75th percentile across all samples.

PvalueThreshold

Threshold used to find relevant driver genes with CNV alterations: maximal p-value.

RsquareThreshold

Threshold used to find relevant driver genes with CNV alterations: minimal R-square value between CNV and gene expression data.

pmax "pmax" variable for glmnet function from glmnet package; the maximum num-

ber of variables aver to be nonzero. Should not be changed by user unless she/he fully understands the AMARETTO algorithm and how its parameters choices

affect model output.

NrCores A numeric variable indicating the number of computer/server cores to use for

paralellelization. Default is 1, i.e. no parallelization. Please check your computer or server's computing capacities before increasing this number. Parallelization is done via the RParallel package. Mac vs. Windows environments

may behave differently when using parallelization.

method Perform union or intersection of the driver genes evaluated from the input data

matrices and custom driver gene list provided.

Examples

AMARETTO_Preprocess AMARETTO_Preprocess

Description

Wrapper code that analyzes process TCGA GISTIC (CNV) and gene expression (rna-seq or mi-croarray) data via one call

Usage

```
AMARETTO_Preprocess(CancerSite, DataSetDirectories)
```

Arguments

DataSetDirectories

Examples

```
data('BatchData')
TargetDirectory <- file.path(getwd(),"Downloads/") # path to data download directory
CancerSite <- "LIHC"
DataSetDirectories <- AMARETTO_Download(CancerSite,TargetDirectory)
ProcessedData <- AMARETTO_Preprocess(CancerSite,DataSetDirectories)</pre>
```

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AMARETTO_Run AMARETTO_Run

Description

Function to run AMARETTO, a statistical algorithm to identify cancer drivers by integrating a variety of omics data from cancer and normal tissue.

Usage

```
AMARETTO_Run(AMARETTOinit)
```

Arguments

```
AMARETTOinit List output from AMARETTO_Initialize().
```

Examples

AMARETTO_VisualizeModule

AMARETTO_VisualizeModule

Description

Function to visualize the gene modules

Usage

```
AMARETTO_VisualizeModule(AMARETTOinit, AMARETTOresults,
   CNV_matrix = NULL, MET_matrix = NULL, ModuleNr,
   SAMPLE_annotation = NULL, ID = NULL, order_samples = NULL)
```

Arguments

AMARETTOinit List output from AMARETTO_Initialize().

AMARETTOresults

List output from AMARETTO_Run().

CNV_matrix CNV matrix, with genes in rows and samples in columns.

MET_matrix Methylation matrix, with genes in rows and samples in columns.

Module number to visualize.

SAMPLE_annotation

Matrix or Dataframe with sample annotation.

ID Column used as sample name.

order_samples Order samples in heatmap by mean or by clustering.

Driver_Genes

Examples

BatchData

BatchData

Description

A dataset for conducting batch corerction in TCGA samples

Usage

BatchData

Format

An object of class data. frame with 23263 rows and 3 columns.

Source

AMARETTO

Driver_Genes

Driver_Genes

Description

A list of cancer driver genes described in literature.

Usage

Driver_Genes

Format

An object of class list of length 10.

Source

AMARETTO

MethylStates 9

MethylStates

MethylStates

Description

A list of dataframes of Methyl States for cancer types obtained from MethylMix processing of TCGA methylation data

Usage

MethylStates

Format

An object of class list of length 25.

Source

AMARETTO

 ${\tt ProcessedDataLIHC}$

ProcessedDataLIHC

Description

A list of dataframes of processed toy example dataset from TCGA-LIHC.

Usage

ProcessedDataLIHC

Format

An object of class list of length 3.

Source

AMARETTO

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