

Package ‘TCGAbiolinks’

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Type Package

Title TCGAbiolinks: An R/Bioconductor package for integrative analysis with GDC data

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Depends R (>= 4.0)

Imports downloader (>= 0.4), grDevices, biomaRt, dplyr, graphics, tibble, GenomicRanges, XML (>= 3.98.0), data.table, jsonlite (>= 1.0.0), plyr, knitr, methods, ggplot2, stringr (>= 1.0.0), IRanges, rvest (>= 0.3.0), stats, utils, S4Vectors, R.utils, SummarizedExperiment (>= 1.4.0), TCGAbiolinksGUI.data (>= 1.15.1), readr, tools, tidyr, purrr, xml2, httr (>= 1.2.1)

Description The aim of TCGAbiolinks is : i) facilitate the GDC open-access data retrieval, ii) prepare the data using the appropriate pre-processing strategies, iii) provide the means to carry out different standard analyses and iv) to easily reproduce earlier research results. In more detail, the package provides multiple methods for analysis (e.g., differential expression analysis, identifying differentially methylated regions) and methods for visualization (e.g., survival plots, volcano plots, starburst plots) in order to easily

develop complete analysis pipelines.

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Suggests jpeg, png, BiocStyle, rmarkdown, devtools, maftools, parmigene, c3net, minet, dnet, Biobase, affy, testthat, sesame, AnnotationHub, ExperimentHub, pathview, clusterProfiler, Seurat, ComplexHeatmap, circlize, ConsensusClusterPlus, igraph, supraHex, limma, edgeR, sva, EDASeq, survminer, genefilter, gridExtra, survival, doParallel, parallel, ggrepel (>= 0.6.3), scales, grid

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BugReports <https://github.com/BioinformaticsFMRP/TCGAbiolinks/issues>

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colDataPrepare	Create samples information matrix for GDC samples
----------------	---

Description

Create samples information matrix for GDC samples add subtype information

Usage

colDataPrepare(barcode)

Arguments

barcode TCGA or TARGET barcode

Examples

```
metadata <- colDataPrepare(c("TCGA-OR-A5K3-01A", "C3N-00321-01"))
metadata <- colDataPrepare(c("BLGSP-71-06-00157-01A",
                             "BLGSP-71-22-00332-01A"))
```

dmc.non.parametric	Perform non-parametrix wilcoxon test
--------------------	--------------------------------------

Description

Perform non-parametrix wilcoxon test

Usage

```
dmc.non.parametric(
  matrix,
  idx1 = NULL,
  idx2 = NULL,
  paired = FALSE,
  adj.method = "BH",
  alternative = "two.sided",
  cores = 1
)
```

Arguments

matrix	A matrix
idx1	Index columns group1
idx2	Index columns group2
paired	Do a paired wilcoxon test? Default: True
adj.method	P-value adjustment method. Default:"BH" Benjamini-Hochberg
alternative	wilcoxon test alternative
cores	Number of cores to be used

Value

Data frame with p-values and diff mean

Examples

```
nrows <- 200; ncols <- 20
counts <- matrix(
  runif(nrows * ncols, 1, 1e4), nrows,
  dimnames = list(paste0("cg",1:200),paste0("S",1:20))
)
TCGAbiolinks::dmc.non.parametric(counts,1:10,11:20)
```

gaiaCNVplot	<i>Creates a plot for GAIA output (all significant aberrant regions.)</i>
-------------	---

Description

This function is a auxiliary function to visualize GAIA output (all significant aberrant regions.)

Usage

```
gaiaCNVplot(calls, threshold = 0.01)
```

Arguments

calls	A matrix with the following columns: Chromosome, Aberration Kind Region Start, Region End, Region Size and score
threshold	Score threshold (orange horizontal line in the plot)

Value

A plot with all significant aberrant regions.

Examples

```
call <- data.frame("Chromosome" = rep(9,100),
                  "Aberration Kind" = rep(c(-2,-1,0,1,2),20),
                  "Region Start [bp]" = 18259823:18259922,
                  "Region End [bp]" = 18259823:18259922,
                  "score" = rep(c(1,2,3,4),25))
gaiaCNVplot(call,threshold = 0.01)
call <- data.frame("Chromosome" = rep(c(1,9),50),
                  "Aberration Kind" = rep(c(-2,-1,0,1,2),20),
                  "Region Start [bp]" = 18259823:18259922,
                  "Region End [bp]" = 18259823:18259922,
                  "score" = rep(c(1,2,3,4),25))
gaiaCNVplot(call,threshold = 0.01)
```

GDCdownload

Download GDC data

Description

Uses GDC API or GDC transfer tool to download gdc data The user can use query argument The data from query will be save in a folder: project/data.category

Usage

```
GDCdownload(
  query,
  token.file,
  method = "api",
  directory = "GDCdata",
  files.per.chunk = NULL
)
```

Arguments

query	A query for GDCquery function
token.file	Token file to download controlled data (only for method = "client")
method	Uses the API (POST method) or gdc client tool. Options "api", "client". API is faster, but the data might get corrupted in the download, and it might need to be executed again
directory	Directory/Folder where the data was downloaded. Default: GDCdata
files.per.chunk	This will make the API method only download n (files.per.chunk) files at a time. This may reduce the download problems when the data size is too large. Expected a integer number (example files.per.chunk = 6)

Value

Shows the output from the GDC transfer tools

Author(s)

Tiago Chedraoui Silva

Examples

```

query <- GDCQuery(project = "TCGA-ACC",
                  data.category = "Copy number variation",
                  legacy = TRUE,
                  file.type = "hg19.seg",
                  barcode = c("TCGA-OR-A5LR-01A-11D-A29H-01", "TCGA-OR-A5LJ-10A-01D-A29K-01"))
# data will be saved in GDCdata/TCGA-ACC/legacy/Copy_number_variation/Copy_number_segmentation
GDCdownload(query, method = "api")
## Not run:
# Download clinical data from XML
query <- GDCQuery(project = "TCGA-COAD", data.category = "Clinical")
GDCdownload(query, files.per.chunk = 200)
query <- GDCQuery(
  project = "TARGET-AML",
  data.category = "Transcriptome Profiling",
  data.type = "miRNA Expression Quantification",
  workflow.type = "BCGSC miRNA Profiling",
  barcode = c("TARGET-20-PARUDL-03A-01R", "TARGET-20-PASRRB-03A-01R")
)
# data will be saved in:
# example_data_dir/TARGET-AML/harmonized/Transcriptome_Profiling/miRNA_Expression_Quantification
GDCdownload(query, method = "client", directory = "example_data_dir")
acc.gbm <- GDCQuery(
  project = c("TCGA-ACC", "TCGA-GBM"),
  data.category = "Transcriptome Profiling",
  data.type = "Gene Expression Quantification",
  workflow.type = "STAR - Counts"
)
GDCdownload(
  query = acc.gbm,
  method = "api",
  directory = "example",
  files.per.chunk = 50
)

## End(Not run)

```

GDCprepare

*Prepare GDC data***Description**

Reads the data downloaded and prepare it into an R object

Usage

```
GDCprepare(
  query,
  save = FALSE,
  save.filename,
  directory = "GDCdata",
  summarizedExperiment = TRUE,
  remove.files.prepared = FALSE,
  add.gistic2.mut = NULL,
  mutant_variant_classification = c("Frame_Shift_Del", "Frame_Shift_Ins",
    "Missense_Mutation", "Nonsense_Mutation", "Splice_Site", "In_Frame_Del",
    "In_Frame_Ins", "Translation_Start_Site", "Nonstop_Mutation")
)
```

Arguments

query	A query for GDCquery function
save	Save result as RData object?
save.filename	Name of the file to be save if empty an automatic will be created
directory	Directory/Folder where the data was downloaded. Default: GDCdata
summarizedExperiment	Create a summarizedExperiment? Default TRUE (if possible)
remove.files.prepared	Remove the files read? Default: FALSE This argument will be considered only if save argument is set to true
add.gistic2.mut	If a list of genes (gene symbol) is given, columns with gistic2 results from GDAC firehose (hg19) and a column indicating if there is or not mutation in that gene (hg38) (TRUE or FALSE - use the MAF file for more information) will be added to the sample matrix in the summarized Experiment object.
mutant_variant_classification	List of mutant_variant_classification that will be consider a sample mutant or not. Default: "Frame_Shift_Del", "Frame_Shift_Ins", "Missense_Mutation", "Nonsense_Mutation", "Splice_Site", "In_Frame_Del", "In_Frame_Ins", "Translation_Start_Site", "Nonstop_Mutation"

Value

A summarizedExperiment or a data.frame

Author(s)

Tiago Chedraoui Silva

Examples

```
## Not run:
query <- GDCquery(
  project = "TCGA-KIRP",
  data.category = "Simple Nucleotide Variation",
  data.type = "Masked Somatic Mutation"
)
GDCdownload(query, method = "api", directory = "maf")
maf <- GDCprepare(query, directory = "maf")

## End(Not run)
```

GDCprepare_clinic	<i>Parsing clinical xml files</i>
-------------------	-----------------------------------

Description

This function receives the query argument and parses the clinical xml files based on the desired information

Usage

```
GDCprepare_clinic(query, clinical.info, directory = "GDCdata")
```

Arguments

query	Result from GDCquery, with data.category set to Clinical
clinical.info	Which information should be retrieved. Options Clinical: drug, admin, follow_up, radiation, patient, stage_event or new_tumor_event Options Biospecimen: protocol, admin, aliquot, analyte, bio_patient, sample, portion, slide
directory	Directory/Folder where the data was downloaded. Default: GDCdata

Value

A data frame with the parsed values from the XML

Examples

```
query <- GDCquery(project = "TCGA-COAD",
  data.category = "Clinical",
  file.type = "xml",
  barcode = c("TCGA-RU-A8FL", "TCGA-AA-3972"))
GDCdownload(query)
clinical <- GDCprepare_clinic(query, "patient")
clinical.drug <- GDCprepare_clinic(query, "drug")
clinical.radiation <- GDCprepare_clinic(query, "radiation")
clinical.admin <- GDCprepare_clinic(query, "admin")
```

```

query <- GDCquery(
  project = "TCGA-COAD",
  data.category = "Biospecimen",
  file.type = "xml",
  data.type = "Biospecimen Supplement",
  barcode = c("TCGA-RU-A8FL", "TCGA-AA-3972")
)
GDCdownload(query)
clinical <- GDCprepare_clinic(query, "admin")
clinical.drug <- GDCprepare_clinic(query, "sample")
clinical.radiation <- GDCprepare_clinic(query, "portion")
clinical.admin <- GDCprepare_clinic(query, "slide")

```

GDCquery

Query GDC data

Description

Uses GDC API to search for search, it searches for both controlled and open-access data. For GDC data arguments project, data.category, data.type and workflow.type should be used For the legacy data arguments project, data.category, platform and/or file.extension should be used. Please, see the vignette for a table with the possibilities.

Usage

```

GDCquery(
  project,
  data.category,
  data.type,
  workflow.type,
  legacy = FALSE,
  access,
  platform,
  file.type,
  barcode,
  data.format,
  experimental.strategy,
  sample.type
)

```

Arguments

project	A list of valid project (see list with <code>TCGAbiolinks::getGDCprojects()\$project_id</code>)
---------	--

- BEATAML1.0-COHORT
- BEATAML1.0-CRENOLANIB
- CGCI-BLGSP
- CPTAC-2

- CPTAC-3
- CTSP-DLBCL1
- FM-AD
- HCM1-CMDC
- MMRF-COMMPASS
- NCICCR-DLBCL
- OHSU-CNL
- ORGANOID-PANCREATIC
- TARGET-ALL-P1
- TARGET-ALL-P2
- TARGET-ALL-P3
- TARGET-AML
- TARGET-CCSK
- TARGET-NBL
- TARGET-OS
- TARGET-RT
- TARGET-WT
- TCGA-ACC
- TCGA-BLCA
- TCGA-BRCA
- TCGA-CESC
- TCGA-CHOL
- TCGA-COAD
- TCGA-DLBC
- TCGA-ESCA
- TCGA-GBM
- TCGA-HNSC
- TCGA-KICH
- TCGA-KIRC
- TCGA-KIRP
- TCGA-LAML
- TCGA-LGG
- TCGA-LIHC
- TCGA-LUAD
- TCGA-LUSC
- TCGA-MESO
- TCGA-OV
- TCGA-PAAD
- TCGA-PCPG
- TCGA-PRAD
- TCGA-READ
- TCGA-SARC

	• TCGA-SKCM • TCGA-STAD • TCGA-TGCT • TCGA-THCA • TCGA-THYM • TCGA-UCEC • TCGA-UCS • TCGA-UVM • VAREPOP-APOLLO														
data.category	A valid project (see list with <code>TCGAbiolinks::getProjectSummary(project)</code>) For the complete list please check the vignette. List for harmonized database: • Biospecimen • Clinical • Copy Number Variation • DNA Methylation • Sequencing Reads • Simple Nucleotide Variation • Transcriptome Profiling List for legacy archive • Biospecimen • Clinical • Copy number variation • DNA methylation • Gene expression • Protein expression • Raw microarray data • Raw sequencing data • Simple nucleotide variation														
data.type	A data type to filter the files to download For the complete list please check the vignette.														
workflow.type	GDC workflow type														
legacy	Search in the legacy repository														
access	Filter by access type. Possible values: controlled, open														
platform	Example:														
	<table> <tr> <td>CGH- 1x1M_G4447A</td><td>IlluminaGA_RNASeqV2</td></tr> <tr> <td>AgilentG4502A_07</td><td>IlluminaGA_mRNA_DGE</td></tr> <tr> <td>Human1MDuo</td><td>HumanMethylation450</td></tr> <tr> <td>HG-CGH-415K_G4124A</td><td>IlluminaGA_miRNASeq</td></tr> <tr> <td>HumanHap550</td><td>IlluminaHiSeq_miRNASeq</td></tr> <tr> <td>ABI</td><td>H-miRNA_8x15K</td></tr> <tr> <td>HG-CGH-244A</td><td>SOLiD_DNASeq</td></tr> </table>	CGH- 1x1M_G4447A	IlluminaGA_RNASeqV2	AgilentG4502A_07	IlluminaGA_mRNA_DGE	Human1MDuo	HumanMethylation450	HG-CGH-415K_G4124A	IlluminaGA_miRNASeq	HumanHap550	IlluminaHiSeq_miRNASeq	ABI	H-miRNA_8x15K	HG-CGH-244A	SOLiD_DNASeq
CGH- 1x1M_G4447A	IlluminaGA_RNASeqV2														
AgilentG4502A_07	IlluminaGA_mRNA_DGE														
Human1MDuo	HumanMethylation450														
HG-CGH-415K_G4124A	IlluminaGA_miRNASeq														
HumanHap550	IlluminaHiSeq_miRNASeq														
ABI	H-miRNA_8x15K														
HG-CGH-244A	SOLiD_DNASeq														

IlluminaDNAMethylation_OMA003_CPI	IlluminaGA_DNASeq_automated
IlluminaDNAMethylation_OMA002_CPI	HG-U133_Plus_2
HuEx- 1_0-st-v2	Mixed_DNASeq
H-miRNA_8x15Kv2	IlluminaGA_DNASeq_curated
MDA_RPPA_Core	IlluminaHiSeq_TotalRNASeqV2
HT_HG-U133A	IlluminaHiSeq_DNASeq_automated
diagnostic_images	microsat_i
IlluminaHiSeq_RNASeq	SOLiD_DNASeq_curated
IlluminaHiSeq_DNASeqC	Mixed_DNASeq_curated
IlluminaGA_RNASeq	IlluminaGA_DNASeq_Cont_automated
IlluminaGA_DNASeq	IlluminaHiSeq_WGBS
pathology_reports	IlluminaHiSeq_DNASeq_Cont_automated
Genome_Wide_SNP_6	bio
tissue_images	Mixed_DNASeq_automated
HumanMethylation27	Mixed_DNASeq_Cont_curated
IlluminaHiSeq_RNASeqV2	Mixed_DNASeq_Cont
file.type	To be used in the legacy database for some platforms, to define which file types to be used.
barcode	A list of barcodes to filter the files to download
data.format	Data format filter ("VCF", "TXT", "BAM", "SVS", "BCR XML", "BCR SSF XML", "TSV", "BCR Auxiliary XML", "BCR OMF XML", "BCR Biotab", "MAF", "BCR PPS XML", "XLSX")
experimental.strategy	Filter to experimental strategy. Harmonized: WXS, RNA-Seq, miRNA-Seq, Genotyping Array. Legacy: WXS, RNA-Seq, miRNA-Seq, Genotyping Array, DNA-Seq, Methylation array, Protein expression array, WXS,CGH array, VALIDATION, Gene expression array,WGS, MSI-Mono-Dinucleotide Assay, miRNA expression array, Mixed strategies, AMPLICON, Exon array, Total RNA-Seq, Capillary sequencing, Bisulfite-Seq
sample.type	A sample type to filter the files to download

Value

A data frame with the results and the parameters used

Author(s)

Tiago Chedraoui Silva

Examples

```
query <- GDCquery(
  project = "TCGA-ACC",
  data.category = "Copy Number Variation",
  data.type = "Copy Number Segment"
)
## Not run:
```

```

query <- GDCquery(
  project = "TARGET-AML",
  data.category = "Transcriptome Profiling",
  data.type = "miRNA Expression Quantification",
  workflow.type = "BCGSC miRNA Profiling",
  barcode = c("TARGET-20-PARUDL-03A-01R", "TARGET-20-PASRRB-03A-01R")
)
query <- GDCquery(
  project = "TARGET-AML",
  data.category = "Transcriptome Profiling",
  data.type = "Gene Expression Quantification",
  workflow.type = "STAR - Counts",
  barcode = c("TARGET-20-PADZCG-04A-01R", "TARGET-20-PARJCR-09A-01R")
)
query <- GDCquery(
  project = "TCGA-ACC",
  data.category = "Copy Number Variation",
  data.type = "Masked Copy Number Segment",
  sample.type = c("Primary Tumor")
)
query.met <- GDCquery(
  project = c("TCGA-GBM", "TCGA-LGG"),
  legacy = TRUE,
  data.category = "DNA methylation",
  platform = "Illumina Human Methylation 450"
)
query <- GDCquery(
  project = "TCGA-ACC",
  data.category = "Copy number variation",
  legacy = TRUE,
  file.type = "hg19.seg",
  barcode = c("TCGA-OR-A5LR-01A-11D-A29H-01")
)

## End(Not run)

```

GDCquery_ATAC_seq

Retrieve open access ATAC-seq files from GDC server

Description

Retrieve open access ATAC-seq files from GDC server [https://gdc.cancer.gov/about-data/publications/ATACseq-AWG Manifest](https://gdc.cancer.gov/about-data/publications/ATACseq-AWG_Manifest) available at: https://gdc.cancer.gov/files/public/file/ATACseq-AWG_Open_GDC-Manifest.txt

Usage

```
GDCquery_ATAC_seq(tumor = NULL, file.type = NULL)
```

Arguments

tumor	a valid tumor
file.type	Write maf file into a csv document

Value

A data frame with the maf file information

Examples

```
query <- GDCquery_ATAC_seq(file.type = "txt")
## Not run:
  GDCdownload(query)

## End(Not run)
query <- GDCquery_ATAC_seq(tumor = "BRCA",file.type = "bigWigs")
## Not run:
  GDCdownload(query,method = "client")

## End(Not run)
```

GDCquery_clinic	<i>Get GDC clinical data</i>
-----------------	------------------------------

Description

GDCquery_clinic will download all clinical information from the API as the one with using the button from each project

Usage

```
GDCquery_clinic(project, type = "clinical", save.csv = FALSE)
```

Arguments

project	A valid project (see list with <code>getGDCprojects()\$project_id</code>) • BEATAML1.0-COHORT • BEATAML1.0-CRENOLANIB • CGCI-BLGSP • CPTAC-2 • CPTAC-3 • CTSP-DLBCL1 • FM-AD • HCM1-CMDC • MMRF-COMMPASS • NCICCR-DLBCL
---------	---

- OHSU-CNL
- ORGANOID-PANCREATIC
- TARGET-ALL-P1
- TARGET-ALL-P2
- TARGET-ALL-P3
- TARGET-AML
- TARGET-CCSK
- TARGET-NBL
- TARGET-OS
- TARGET-RT
- TARGET-WT
- TCGA-ACC
- TCGA-BLCA
- TCGA-BRCA
- TCGA-CESC
- TCGA-CHOL
- TCGA-COAD
- TCGA-DLBC
- TCGA-ESCA
- TCGA-GBM
- TCGA-HNSC
- TCGA-KICH
- TCGA-KIRC
- TCGA-KIRP
- TCGA-LAML
- TCGA-LGG
- TCGA-LIHC
- TCGA-LUAD
- TCGA-LUSC
- TCGA-MESO
- TCGA-OV
- TCGA-PAAD
- TCGA-PCPG
- TCGA-PRAD
- TCGA-READ
- TCGA-SARC
- TCGA-SKCM
- TCGA-STAD
- TCGA-TGCT
- TCGA-THCA
- TCGA-THYM
- TCGA-UCEC

	• TCGA-UCS • TCGA-UVM • VAREPOP-APOLLO
type	A valid type. Options "clinical", "Biospecimen" (see list with getGDCprojects()\$project_id)]
save.csv	Write clinical information into a csv document

Value

A data frame with the clinical information

Author(s)

Tiago Chedraoui Silva

Examples

```
clinical <- GDCquery_clinic(project = "TCGA-ACC", type = "clinical", save.csv = TRUE)
clinical <- GDCquery_clinic(project = "TCGA-ACC", type = "biospecimen", save.csv = TRUE)
clinical.cptac2 <- GDCquery_clinic(project = "CPTAC-2", type = "clinical")
clinical.TARGET_ALL_P1 <- GDCquery_clinic(project = "TARGET-ALL-P1", type = "clinical")
clinicaln.fm_ad <- GDCquery_clinic(project = "FM-AD", type = "clinical")
## Not run:
clinical.cptac_3 <- GDCquery_clinic(project = "CPTAC-3", type = "clinical")
clinical.cptac_2 <- GDCquery_clinic(project = "CPTAC-2", type = "clinical")
clinical.HCMI_CMDC <- GDCquery_clinic(project = "HCMI-CMDC", type = "clinical")
clinical.CGCI-HTMCP-CC <- GDCquery_clinic(project = "CGCI-HTMCP-CC", type = "clinical")
clinical <- GDCquery_clinic(project = "NCICCR-DLBCL", type = "clinical")
clinical <- GDCquery_clinic(project = "ORGANOID-PANCREATIC", type = "clinical")

## End(Not run)
```

get.GRCh.bioMart

Get hg19 gene annotation or hg38 (gencode v36)

Description

Get hg19 (from biomaRt) or hg38 (gencode v36) gene annotation

Usage

```
get.GRCh.bioMart(genome = c("hg19", "hg38"), as.granges = FALSE)
```

Arguments

genome	hg38 or hg19
as.granges	Output as GRanges or data.frame

getAdjacencyBiogrid	<i>Get a matrix of interactions of genes from biogrid</i>
---------------------	---

Description

Using biogrid database, it will create a matrix of gene interactions. If columns A and row B has value 1, it means the gene A and gene B interacts.

Usage

```
getAdjacencyBiogrid(tmp.biogrid, names.genes = NULL)
```

Arguments

tmp.biogrid	Biogrid table
names.genes	List of genes to filter from output. Default: consider all genes

Value

A matrix with 1 for genes that interacts, 0 for no interaction.

Examples

```
names.genes.de <- c("PLCB1","MCL1","PRDX4","TTF2","TACC3", "PARP4","LSM1")
tmp.biogrid <- data.frame("Official.Symbol.Interactor.A" = names.genes.de,
                          "Official.Symbol.Interactor.B" = rev(names.genes.de))
net.biogrid.de <- getAdjacencyBiogrid(tmp.biogrid, names.genes.de)
## Not run:
file <- paste0("http://thebiogrid.org/downloads/archives/",
               "Release%20Archive/BIOGRID-3.4.133/BIOGRID-ALL-3.4.133.tab2.zip")
downloader::download(file,basename(file))
unzip(basename(file),junkpaths =TRUE)
tmp.biogrid <- read.csv(gsub("zip","txt",basename(file)),
                      header=TRUE, sep="\t", stringsAsFactors=FALSE)
names.genes.de <- c("PLCB1","MCL1","PRDX4","TTF2","TACC3", "PARP4","LSM1")
net.biogrid.de <- getAdjacencyBiogrid(tmp.biogrid, names.genes.de)

## End(Not run)
```

getDataCategorySummary

Create a Summary table for each sample in a project saying if it contains or not files for a certain data category

Description

Create a Summary table for each sample in a project saying if it contains or not files for a certain data category

Usage

```
getDataCategorySummary(project, legacy = FALSE)
```

Arguments

project	A GDC project
legacy	Access legacy (hg19) or harmonized database (hg38).

Value

A data frame

Author(s)

Tiago Chedraoui Silva

Examples

```
summary <- getDataCategorySummary("TCGA-ACC", legacy = TRUE)
```

getGDCInfo

Check GDC server status

Description

Check GDC server status using the api <https://api.gdc.cancer.gov/status>

Usage

```
getGDCInfo()
```

Value

Return true all status

Examples

```
info <- getGDCInfo()
```

getGDCprojects	<i>Retrieve all GDC projects</i>
----------------	----------------------------------

Description

getGDCprojects uses the following api to get projects <https://api.gdc.cancer.gov/projects>

Usage

```
getGDCprojects()
```

Value

A data frame with last GDC projects

Examples

```
projects <- getGDCprojects()
```

getGistic	<i>Download GISTIC data from firehose</i>
-----------	---

Description

Download GISTIC data from firehose from http://gdac.broadinstitute.org/runs/analyses__latest/data/

Usage

```
getGistic(disease, type = "thresholded")
```

Arguments

disease	TCGA disease. Option available in http://gdac.broadinstitute.org/runs/analyses__latest/data/
type	Results type: thresholded or data

getLinkedOmicsData	<i>Retrieve linkedOmics data</i>
--------------------	----------------------------------

Description

Retrieve linkedOmics data from <http://linkedomics.org/>

Usage

```
getLinkedOmicsData(project, dataset)
```

Arguments

project	A linkedOmics project: • TCGA-ACC • TCGA-BLCA • TCGA-BRCA • TCGA-CESC • TCGA-CHOL • TCGA-COADREAD • TCGA-DLBC • TCGA-ESCA • TCGA-GBM • TCGA-GBMLGG • TCGA-HNSC • TCGA-KICH • TCGA-KIPAN • TCGA-KIRC • TCGA-KIRP • TCGA-LAML • TCGA-LGG • TCGA-LIHC • TCGA-LUAD • TCGA-LUSC • TCGA-MESO • TCGA-OV • TCGA-PAAD • TCGA-PCPG • TCGA-PRAD • TCGA-SARC • TCGA-SKCM • TCGA-STAD
---------	--

- TCGA-STES
- TCGA-TGCT
- TCGA-THCA
- TCGA-THYM
- TCGA-UCEC
- TCGA-UCS
- TCGA-UVM
- CPTAC-COAD

dataset

A dataset from the list below

- Annotated mutation
- Clinical
- Glycoproteome (Gene level)
- Glycoproteome (Site level)
- Methylation (CpG-site level, HM27)
- Methylation (CpG-site level, HM450K)
- Methylation (Gene level, HM27)
- Methylation (Gene level, HM450K)
- miRNA (GA, Gene level)
- miRNA (GA, Isoform level)
- miRNA (GA, miRgene level)
- miRNA (Gene level)
- miRNA (HiSeq, Gene level)
- miRNA (HiSeq, miRgene level)
- miRNA (isoform level)
- miRNA (miRgene level)
- Mutation (Gene level)
- Mutation (Site level)
- Mutation raw file (Somatic and MSIIndel)
- Phosphoproteome (Gene level)
- Phosphoproteome (Site level)
- Phosphoproteomics (Normal)
- Phosphoproteomics (Tumor)
- Proteome (Gene level)
- Proteome (Gene Level)
- Proteome (JHU, Gene level)
- Proteome (PNNL, Gene level, Normal TMT Unshared Log Ratio)
- Proteome (PNNL, Gene level, Tumor TMT Unshared Log Ratio)
- Proteome (PNNL, Gene level)
- Proteome (VU, Gene level, Label-free Unshared Counts)
- RNAseq (GA, Gene level)
- RNAseq (HiSeq, Gene level)
- RPPA (Analyte level)

- RPPA (Analyte Level)
- RPPA (Gene level)
- RPPA (Gene Level)
- SCNV (Focal level, log-ratio)
- SCNV (Focal level, Thresholded)
- SCNV (Gene level, log ratio)
- SCNV (Gene level, log-ratio)
- SCNV (Gene level, Thresholded)
- SCNV (Segment level)

Value

A matrix with the data

Examples

```
## Not run:
TCGA_COAD_protein <- getLinkedOmicsData(
  project = "TCGA-COADREAD",
  dataset = "Proteome (Gene level)"
)
TCGA_COAD_RNASeq_hiseq <- getLinkedOmicsData(
  project = "TCGA-COADREAD",
  dataset = "RNAseq (HiSeq, Gene level)"
)
TCGA_COAD_RNASeq_ga <- getLinkedOmicsData(
  project = "TCGA-COADREAD",
  dataset = "RNAseq (GA, Gene level)"
)
TCGA_COAD_RPPA <- getLinkedOmicsData(
  project = "TCGA-COADREAD",
  dataset = "RPPA (Gene level)"
)

## End(Not run)
```

getManifest

Get a Manifest from GDCquery output that can be used with GDC-client

Description

Get a Manifest from GDCquery output that can be used with GDC-client

Usage

```
getManifest(query, save = FALSE)
```

Arguments

query	A query for GDCquery function
save	Write Manifest to a txt file (tab separated)

Examples

```
query <- GDCquery(
  project = "TARGET-AML",
  data.category = "Transcriptome Profiling",
  data.type = "Gene Expression Quantification",
  workflow.type = "STAR - Counts",
  barcode = c("TARGET-20-PADZCG-04A-01R", "TARGET-20-PARJCR-09A-01R")
)
getManifest(query)
```

getMC3MAF

*Retrieve open access mc3 MAF file from GDC server***Description**

Download data from <https://gdc.cancer.gov/about-data/publications/mc3-2017> https://gdc-docs.nci.nih.gov/Data/Release_No

Usage

```
getMC3MAF()
```

Value

A data frame with the MAF file information from <https://gdc.cancer.gov/about-data/publications/mc3-2017>

Examples

```
## Not run:
  maf <- getMC3MAF()

## End(Not run)
```

getNbCases	<i>Get Number of cases in GDC for a project</i>
------------	---

Description

Get Number of cases in GDC for a project

Usage

```
getNbCases(project, data.category, legacy = FALSE)
```

Arguments

project	A GDC project
data.category	A GDC project data category
legacy	Select between Harmonized or Legacy database

Author(s)

Tiago Chedraoui Silva

Examples

```
## Not run:  
getNbCases("TCGA-ACC", "Clinical")  
getNbCases("CPTAC-2", "Clinical")  
  
## End(Not run)
```

getNbFiles	<i>Get Number of files in GDC for a project</i>
------------	---

Description

Get Number of files in GDC for a project

Usage

```
getNbFiles(project, data.category, legacy = FALSE)
```

Arguments

project	A GDC project
data.category	A GDC project data category
legacy	Select between Harmonized or Legacy database

Author(s)

Tiago Chedraoui Silva

Examples

```
## Not run:  
getNbFiles("TCGA-ACC","Clinical")  
getNbFiles("CPTAC-2","Clinical")  
  
## End(Not run)
```

getProjectSummary	<i>Get Project Summary from GDC</i>
-------------------	-------------------------------------

Description

Get Project Summary from GDC

Usage

```
getProjectSummary(project, legacy = FALSE)
```

Arguments

project	A GDC project
legacy	Select between Harmonized or Legacy database

Author(s)

Tiago Chedraoui Silva

Examples

```
getProjectSummary("TCGA-ACC")  
## Not run:  
getProjectSummary("CPTAC-2")  
  
## End(Not run)
```

getResults	<i>Get the results table from query</i>
------------	---

Description

Get the results table from query, it can select columns with cols argument and return a number of rows using rows argument.

Usage

```
getResults(query, rows, cols)
```

Arguments

query	A object from GDCquery
rows	Rows identifiers (row numbers)
cols	Columns identifiers (col names)

Value

Table with query results

Examples

```
query <- GDCquery(  
  project = "TCGA-GBM",  
  data.category = "Transcriptome Profiling",  
  data.type = "Gene Expression Quantification",  
  workflow.type = "STAR - Counts",  
  barcode = c("TCGA-14-0736-02A-01R-2005-01", "TCGA-06-0211-02A-02R-2005-01")  
)  
results <- getResults(query)
```

getSampleFilesSummary	<i>Retrieve summary of files per sample in a project</i>
-----------------------	--

Description

Retrieve the numner of files under each data_category + data_type + experimental_strategy + platform Almost like <https://portal.gdc.cancer.gov/exploration>

Usage

```
getSampleFilesSummary(project, legacy = FALSE, files.access = NA)
```

Arguments

project	A GDC project
legacy	Access legacy database ? Deafult: FALSE
files.access	Filter by file access ("open" or "controlled"). Default: no filter

Value

A data frame with the maf file information

Author(s)

Tiago Chedraoui Silva

Examples

```
summary <- getSampleFilesSummary("TCGA-UCS")
## Not run:
summary <- getSampleFilesSummary(c("TCGA-OV", "TCGA-ACC"))

## End(Not run)
```

getTSS	<i>getTSS to fetch GENCODE gene annotation (transcripts level) from Bioconductor package biomaRt If upstream and downstream are specified in TSS list, promoter regions of GENCODE gene will be generated.</i>
--------	--

Description

getTSS to fetch GENCODE gene annotation (transcripts level) from Bioconductor package biomaRt
If upstream and downstream are specified in TSS list, promoter regions of GENCODE gene will be generated.

Usage

```
getTSS(
  genome = c("hg38", "hg19"),
  TSS = list(upstream = NULL, downstream = NULL)
)
```

Arguments

genome	Which genome build will be used: hg38 (default) or hg19.
TSS	A list. Contains upstream and downstream like TSS=list(upstream, downstream). When upstream and downstream is specified, coordinates of promoter regions with gene annotation will be generated.

Value

GENCODE gene annotation if TSS is not specified. Coordinates of GENCODE gene promoter regions if TSS is specified.

Examples

```
# get GENCODE gene annotation (transcripts level)
## Not run:
  getTSS <- getTSS()
  getTSS <- getTSS(genome.build = "hg38", TSS=list(upstream=1000, downstream=1000))

## End(Not run)
```

get_IDs

*Extract information from TCGA barcodes.***Description**

get_IDs allows user to extract metadata from barcodes. The dataframe returned has columns for 'project', 'tss', 'participant', 'sample', "portion", "plate", and "center"

Usage

```
get_IDs(data)
```

Arguments

data numeric matrix, each row represents a gene, each column represents a sample

Value

data frame with columns 'project', 'tss', 'participant', 'sample', "portion", "plate", "center", "condition"

gliomaClassifier

*Glioma classifier***Description**

Classify DNA methylation gliomas using data from <https://doi.org/10.1016/j.cell.2015.12.028>

Usage

```
gliomaClassifier(data)
```

Arguments

data DNA methylation matrix or Summarized Experiments with samples on columns and probes on the rows

Value

A list of 3 data frames: 1) Sample final classification 2) Each model final classification 3) Each class probability of classification

Author(s)

Tiago Chedraoui Silva, Tathiane Malta, Houtan Noushmehr

Examples

```
## Not run:
query <- GDCquery(project= "TCGA-GBM",
                  data.category = "DNA methylation",
                  barcode = c("TCGA-06-0122", "TCGA-14-1456"),
                  platform = "Illumina Human Methylation 27",
                  legacy = TRUE)
GDCdownload(query)
data.hg19 <- GDCprepare(query)
classification <- gliomaClassifier(data.hg19)

# Comparing results
TCGAquery_subtype("GBM") %>%
dplyr::filter(patient %in% c("TCGA-06-0122", "TCGA-14-1456")) %>%
dplyr::select("patient", "Supervised.DNA.Methylation.Cluster")

## End(Not run)
```

isServeOK

Check GDC server status is OK

Description

Check GDC server status using the api <https://api.gdc.cancer.gov/status>

Usage

```
isServeOK()
```

Value

Return true if status is ok

Examples

```
status <- isServeOK()
```

matchedMetExp	<i>Get GDC samples with both DNA methylation (HM450K) and Gene expression data from GDC database</i>
---------------	--

Description

For a given TCGA project it gets the samples (barcode) with both DNA methylation and Gene expression data from GDC database

Usage

```
matchedMetExp(project, legacy = FALSE, n = NULL)
```

Arguments

project	A GDC project
legacy	Access legacy (hg19) or harmonized database (hg38).
n	Number of samples to return. If NULL return all (default)

Value

A vector of barcodes

Examples

```
# Get ACC samples with both DNA methylation (HM450K) and gene expression aligned to hg19
samples <- matchedMetExp("TCGA-UCS", legacy = TRUE)
```

PanCancerAtlas_subtypes	<i>Retrieve table with TCGA molecular subtypes</i>
-------------------------	--

Description

PanCancerAtlas_subtypes is a curated table with molecular subtypes for 24 TCGA cancer types

Usage

```
PanCancerAtlas_subtypes()
```

Value

a data.frame with barcode and molecular subtypes for 24 cancer types

Examples

```
molecular.subtypes <- PanCancerAtlas_subtypes()
```

splitAPICall	<i>internal function to break a huge API call into smaller ones so it respects the max character limit of a string</i>
--------------	--

Description

internal function to break a huge API call into smaller ones so it respects the max character limit of a string

Usage

```
splitAPICall(FUN, step = 20, items)
```

Arguments

FUN	function that calls the API
step	How many items to be evaluated per API call
items	vector of items to be using within the function (list of barcodes, aliquot ids, etc)

Author(s)

Tiago Chedraoui Silva

TabSubtypesCol_merged	<i>TCGA samples with their Pam50 subtypes</i>
-----------------------	---

Description

A dataset containing the Sample Ids from TCGA and PAM50 subtyping attributes of 4768 tumor patients

Usage

```
TabSubtypesCol_merged
```

Format

A data frame with 4768 rows and 3 variables:

samples Sample ID from TCGA barcodes, character string

subtype Pam50 classification, character string

color color, character string ...

`TCGAanalyze_analyseGRN`*Generate network*

Description

TCGAanalyze_analyseGRN perform gene regulatory network.

Usage

```
TCGAanalyze_analyseGRN(TFs, normCounts, kNum)
```

Arguments

TFs	a vector of genes.
normCounts	is a matrix of gene expression with genes in rows and samples in columns.
kNum	the number of nearest neighbors to consider to estimate the mutual information. Must be less than the number of columns of normCounts.

Value

an adjacent matrix

`TCGAanalyze_Clustering`*Hierarchical cluster analysis*

Description

Hierarchical cluster analysis using several methods such as ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC).

Usage

```
TCGAanalyze_Clustering(tabDF, method, methodHC = "ward.D2")
```

Arguments

tabDF	is a dataframe or numeric matrix, each row represents a gene, each column represents a sample come from TCGAPrepare.
method	is method to be used for generic cluster such as 'hclust' or 'consensus'
methodHC	is method to be used for Hierarchical cluster.

Value

object of class `hclust` if method selected is `'hclust'`. If method selected is `'Consensus'` returns a list of length `maxK` (maximum cluster number to evaluate.). Each element is a list containing `consensusMatrix` (numerical matrix), `consensusTree` (`hclust`), `consensusClass` (consensus class assignments). `ConsensusClusterPlus` also produces images.

TCGAanalyze_DEA

Differential expression analysis (DEA) using edgeR or limma package.

Description

TCGAanalyze_DEA allows user to perform Differentially expression analysis (DEA), using `edgeR` package or `limma` to identify differentially expressed genes (DEGs). It is possible to do a two-class analysis.

TCGAanalyze_DEA performs DEA using following functions from `edgeR`:

1. `edgeR::DGEList` converts the count matrix into an `edgeR` object.
2. `edgeR::estimateCommonDisp` each gene gets assigned the same dispersion estimate.
3. `edgeR::exactTest` performs pair-wise tests for differential expression between two groups.
4. `edgeR::topTags` takes the output from `exactTest()`, adjusts the raw p-values using the False Discovery Rate (FDR) correction, and returns the top differentially expressed genes.

TCGAanalyze_DEA performs DEA using following functions from `limma`:

1. `limma::makeContrasts` construct matrix of custom contrasts.
2. `limma::lmFit` Fit linear model for each gene given a series of arrays.
3. `limma::contrasts.fit` Given a linear model fit to microarray data, compute estimated coefficients and standard errors for a given set of contrasts.
4. `limma::eBayes` Given a microarray linear model fit, compute moderated t-statistics, moderated F-statistic, and log-odds of differential expression by empirical Bayes moderation of the standard errors towards a common value.
5. `limma::topTable` Extract a table of the top-ranked genes from a linear model fit.

Usage

```
TCGAanalyze_DEA(
  mat1,
  mat2,
  metadata = TRUE,
  Cond1type,
  Cond2type,
  pipeline = "edgeR",
  method = "exactTest",
  fdr.cut = 1,
  logFC.cut = 0,
```

```

batch.factors = NULL,
ClinicalDF = data.frame(),
paired = FALSE,
log.trans = FALSE,
voom = FALSE,
trend = FALSE,
MAT = data.frame(),
contrast.formula = "",
Condtypes = c()
)

```

Arguments

mat1	numeric matrix, each row represents a gene, each column represents a sample with Cond1type
mat2	numeric matrix, each row represents a gene, each column represents a sample with Cond2type
metadata	Add metadata
Cond1type	a string containing the class label of the samples in mat1 (e.g., control group)
Cond2type	a string containing the class label of the samples in mat2 (e.g., case group)
pipeline	a string to specify which package to use ("limma" or "edgeR")
method	is 'glmLRT' (1) or 'exactTest' (2) used for edgeR (1) Fit a negative binomial generalized log-linear model to the read counts for each gene (2) Compute gene-wise exact tests for differences in the means between two groups of negative-binomially distributed counts.
fdr.cut	is a threshold to filter DEGs according their p-value corrected
logFC.cut	is a threshold to filter DEGs according their logFC
batch.factors	a vector containing strings to specify options for batch correction. Options are "Plate", "TSS", "Year", "Portion", "Center", and "Patients"
ClinicalDF	a dataframe returned by GDCquery_clinic() to be used to extract year data
paired	boolean to account for paired or non-paired samples. Set to TRUE for paired case
log.trans	boolean to perform log cpm transformation. Set to TRUE for log transformation
voom	boolean to perform voom transformation for limma-voom pipeline. Set to TRUE for voom transformation
trend	boolean to perform limma-trend pipeline. Set to TRUE to go through limma-trend
MAT	matrix containing expression set as all samples in columns and genes as rows. Do not provide if mat1 and mat2 are used
contrast.formula	string input to determine coefficients and to design contrasts in a customized way
Condtypes	vector of grouping for samples in MAT

Value

table with DEGs containing for each gene logFC, logCPM, pValue, and FDR, also for each contrast

Examples

```
dataNorm <- TCGAanalyze_Normalization(dataBRCA, geneInfo)
dataFilt <- TCGAanalyze_Filtering(tabDF = dataBRCA, method = "quantile", qnt.cut = 0.25)
samplesNT <- TCGAquery_SampleTypes(colnames(dataFilt), typesample = c("NT"))
samplesTP <- TCGAquery_SampleTypes(colnames(dataFilt), typesample = c("TP"))
dataDEGs <- TCGAanalyze_DEA(
  mat1 = dataFilt[,samplesNT],
  mat2 = dataFilt[,samplesTP],
  Cond1type = "Normal",
  Cond2type = "Tumor"
)
```

TCGAanalyze_DEA_Affy *Differentially expression analysis (DEA) using limma package.*

Description

Differentially expression analysis (DEA) using limma package.

Usage

```
TCGAanalyze_DEA_Affy(AffySet, FC.cut = 0.01)
```

Arguments

AffySet	A matrix-like data object containing log-ratios or log-expression values for a series of arrays, with rows corresponding to genes and columns to samples
FC.cut	write

Value

List of list with tables in 2 by 2 comparison of the top-ranked genes from a linear model fitted by DEA's limma

Examples

```
## Not run:
to add example

## End(Not run)
```

Description

This function will search for differentially methylated CpG sites, which are regarded as possible functional regions involved in gene transcriptional regulation.

In order to find these regions we use the beta-values (methylation values ranging from 0.0 to 1.0) to compare two groups.

Firstly, it calculates the difference between the mean methylation of each group for each probes. Secondly, it calculates the p-value using the wilcoxon test using the Benjamini-Hochberg adjustment method. The default parameters will require a minimum absolute beta values delta of 0.2 and a false discovery rate (FDR)-adjusted Wilcoxon rank-sum P-value of < 0.01 for the difference.

After these analysis, we save a volcano plot (x-axis:diff mean methylation, y-axis: significance) that will help the user identify the differentially methylated CpG sites and return the object with the calculus in the rowRanges.

If the calculus already exists in the object it will not recalculated. You should set overwrite parameter to TRUE to force it, or remove the columns with the results from the object.

Usage

```
TCGAanalyze_DMC(
  data,
  groupCol = NULL,
  group1 = NULL,
  group2 = NULL,
  alternative = "two.sided",
  diffmean.cut = 0.2,
  paired = FALSE,
  adj.method = "BH",
  plot.filename = "methylation_volcano.pdf",
  ylab = expression(paste(-Log[10], " (FDR corrected P-values)")),
  xlab = expression(paste("DNA Methylation difference (", beta, "-values)")),
  title = NULL,
  legend = "Legend",
  color = c("black", "red", "darkgreen"),
  label = NULL,
  xlim = NULL,
  ylim = NULL,
  p.cut = 0.01,
  probe.names = FALSE,
  cores = 1,
  save = TRUE,
  save.directory = ".",
  filename = NULL
)
```

Arguments

<code>data</code>	SummarizedExperiment obtained from the TCGAPrepare
<code>groupCol</code>	Columns with the groups inside the SummarizedExperiment object. (This will be obtained by the function <code>colData(data)</code>)
<code>group1</code>	In case our object has more than 2 groups, you should set the name of the group
<code>group2</code>	In case our object has more than 2 groups, you should set the name of the group
<code>alternative</code>	wilcoxon test alternative
<code>diffmean.cut</code>	diffmean threshold. Default: 0.2
<code>paired</code>	Wilcoxon paired parameter. Default: FALSE
<code>adj.method</code>	Adjusted method for the p-value calculation
<code>plot.filename</code>	Filename. Default: volcano.pdf, volcano.svg, volcano.png. If set to FALSE, there will be no plot.
<code>ylab</code>	y axis text
<code>xlab</code>	x axis text
<code>title</code>	main title. If not specified it will be "Volcano plot (group1 vs group2)"
<code>legend</code>	Legend title
<code>color</code>	vector of colors to be used in graph
<code>label</code>	vector of labels to be used in the figure. Example: <code>c("Not Significant", "Hypermethylated in group1", "Hypomethylated in group1")</code>
<code>xlim</code>	x limits to cut image
<code>ylim</code>	y limits to cut image
<code>p.cut</code>	p values threshold. Default: 0.01
<code>probe.names</code>	is probe.names
<code>cores</code>	Number of cores to be used in the non-parametric test Default = <code>groupCol.group1.group2.rda</code>
<code>save</code>	Save object with results? Default: TRUE
<code>save.directory</code>	Directory to save the files. Default: working directory
<code>filename</code>	Name of the file to save the object.

Value

Volcano plot saved and the given data with the results (`diffmean.group1.group2`, `p.value.group1.group2`, `p.value.adj.group1.group2`, `status.group1.group2`) in the `rowRanges` where `group1` and `group2` are the names of the groups

Examples

```
nrows <- 200; ncols <- 20
counts <- matrix(
  runif(nrows * ncols, 1, 1e4), nrows,
  dimnames = list(paste0("cg", 1:200), paste0("S", 1:20))
)
rowRanges <- GenomicRanges::GRanges(
```

```

    rep(c("chr1", "chr2"), c(50, 150)),
    IRanges::IRanges(floor(runif(200, 1e5, 1e6)), width = 100),
    strand = sample(c("+", "-"), 200, TRUE),
    feature_id = sprintf("ID%03d", 1:200)
  )
  names(rowRanges) <- paste0("cg", 1:200)
  colData <- S4Vectors::DataFrame(
    Treatment = rep(c("ChIP", "Input"), 5),
    row.names = paste0("S", 1:20),
    group = rep(c("group1", "group2"), c(10, 10))
  )
  data <- SummarizedExperiment::SummarizedExperiment(
    assays=S4Vectors::SimpleList(counts=counts),
    rowRanges = rowRanges,
    colData = colData
  )
  SummarizedExperiment::colData(data)$group <- c(rep("group 1", ncol(data)/2),
    rep("group 2", ncol(data)/2))
  hypo.hyper <- TCGAanalyze_DMC(data, p.cut = 0.85, "group", "group 1", "group 2")
  SummarizedExperiment::colData(data)$group2 <- c(rep("group_1", ncol(data)/2),
    rep("group_2", ncol(data)/2))
  hypo.hyper <- TCGAanalyze_DMC(
    data = data,
    p.cut = 0.85,
    groupCol = "group2",
    group1 = "group_1",
    group2 = "group_2"
  )

```

TCGAanalyze_EA

Enrichment analysis of a gene-set with GO [BP, MF, CC] and pathways.

Description

The rationale behind an enrichment analysis (gene-set, pathway etc) is to compute statistics of whether the overlap between the focus list (signature) and the gene-set is significant, i.e. the confidence that overlap between the list is not due to chance. The Gene Ontology project describes genes (gene products) using terms from three structured vocabularies: biological process, cellular component and molecular function. The Gene Ontology Enrichment component, also referred to as the "GO Terms" component, allows the genes in any such "changed-gene" list to be characterized using the Gene Ontology terms annotated to them. It asks, whether for any particular GO term, the fraction of genes assigned to it in the "changed-gene" list is higher than expected by chance (is over-represented), relative to the fraction of genes assigned to that term in the reference set. In statistical terms it performs the analysis tests the null hypothesis that, for any particular ontology term, there is no difference in the proportion of genes annotated to it in the reference list and the proportion annotated to it in the test list. We adopted a Fisher Exact Test to perform the EA.

Usage

```
TCGAanalyze_EA(
  GeneName,
  RegulonList,
  TableEnrichment,
  EAGenes,
  GOtype,
  FDRThresh = 0.01,
  GeneSymbolsTable = FALSE
)
```

Arguments

GeneName	is the name of gene signatures list
RegulonList	is a gene signature (list of genes) in which perform EA.
TableEnrichment	is a table related to annotations of gene symbols such as GO[BP,MF,CC] and Pathways. It was created from DAVID gene ontology on-line.
EAGenes	is a table with informations about genes such as ID, Gene, Description, Location and Family.
GOtype	is type of gene ontology Biological process (BP), Molecular Function (MF), Cellular component (CC)
FDRThresh	pvalue corrected (FDR) as threshold to selected significant BP, MF,CC, or pathways. (default FDR < 0.01)
GeneSymbolsTable	if it is TRUE will return a table with GeneSymbols in common GO or pathways.

Value

Table with enriched GO or pathways by selected gene signature.

Examples

```
## Not run:
EAGenes <- get("EAGenes")
RegulonList <- rownames(dataDEGsFiltLevel)
ResBP <- TCGAanalyze_EA(GeneName="DEA genes Normal Vs Tumor",
                        RegulonList,DAVID_BP_matrix,
                        EAGenes,GOtype = "DavidBP")

## End(Not run)
```

`TCGAanalyze_EAcomplete`*Enrichment analysis for Gene Ontology (GO) [BP,MF,CC] and Pathways*

Description

Researchers, in order to better understand the underlying biological processes, often want to retrieve a functional profile of a set of genes that might have an important role. This can be done by performing an enrichment analysis.

We will perform an enrichment analysis on gene sets using the `TCGAanalyze_EAcomplete` function. Given a set of genes that are up-regulated under certain conditions, an enrichment analysis will find identify classes of genes or proteins that are #’over-represented using annotations for that gene set.

Usage

```
TCGAanalyze_EAcomplete(TFname, RegulonList)
```

Arguments

<code>TFname</code>	is the name of the list of genes or TF’s regulon.
<code>RegulonList</code>	List of genes such as TF’s regulon or DEGs where to find enrichment.

Value

Enrichment analysis GO[BP,MF,CC] and Pathways complete table enriched by genelist.

Examples

```
Genelist <- c("FN1","COL1A1")
ansEA <- TCGAanalyze_EAcomplete(TFname="DEA genes Normal Vs Tumor",Genelist)
## Not run:
Genelist <- rownames(dataDEGsFiltLevel)
system.time(ansEA <- TCGAanalyze_EAcomplete(TFname="DEA genes Normal Vs Tumor",Genelist))

## End(Not run)
```

TCGAanalyze_Filtering *Filtering mRNA transcripts and miRNA selecting a threshold.*

Description

TCGAanalyze_Filtering allows user to filter mRNA transcripts and miRNA, samples, higher than the threshold defined quantile mean across all samples.

Usage

```
TCGAanalyze_Filtering(
  tabDF,
  method,
  qnt.cut = 0.25,
  var.func = IQR,
  var.cutoff = 0.75,
  eta = 0.05,
  foldChange = 1
)
```

Arguments

tabDF	is a dataframe or numeric matrix, each row represents a gene, each column represents a sample come from TCGAPrepare
method	is method of filtering such as 'quantile', 'varFilter', 'filter1', 'filter2'
qnt.cut	is threshold selected as mean for filtering
var.func	is function used as the per-feature filtering statistic. See genefilter documentation
var.cutoff	is a numeric value. See genefilter documentation
eta	is a parameter for filter1. default eta = 0.05.
foldChange	is a parameter for filter2. default foldChange = 1.

Value

A filtered dataframe or numeric matrix where each row represents a gene, each column represents a sample

Examples

```
dataNorm <- TCGAbiolinks::TCGAanalyze_Normalization(dataBRCA, geneInfo)
dataNorm <- TCGAanalyze_Normalization(tabDF = dataBRCA,
  geneInfo = geneInfo,
  method = "geneLength")
dataFilt <- TCGAanalyze_Filtering(tabDF = dataNorm, method = "quantile", qnt.cut = 0.25)
```

TCGAanalyze_LevelTab	<i>Adding information related to DEGs genes from DEA as mean values in two conditions.</i>
----------------------	--

Description

TCGAanalyze_LevelTab allows user to add information related to DEGs genes from Differentially expression analysis (DEA) such as mean values and in two conditions.

Usage

```
TCGAanalyze_LevelTab(
  FC_FDR_table_mRNA,
  typeCond1,
  typeCond2,
  TableCond1,
  TableCond2,
  typeOrder = TRUE
)
```

Arguments

FC_FDR_table_mRNA	Output of dataDEGs filter by $\text{abs}(\text{LogFC}) \geq 1$
typeCond1	a string containing the class label of the samples in TableCond1 (e.g., control group)
typeCond2	a string containing the class label of the samples in TableCond2 (e.g., case group)
TableCond1	numeric matrix, each row represents a gene, each column represents a sample with Cond1type
TableCond2	numeric matrix, each row represents a gene, each column represents a sample with Cond2type
typeOrder	typeOrder

Value

table with DEGs, log Fold Change (FC), false discovery rate (FDR), the gene expression level for samples in Cond1type, and Cond2type, and Delta value (the difference of gene expression between the two conditions multiplied logFC)

Examples

```
dataNorm <- TCGAanalyze_Normalization(dataBRCA, geneInfo)
dataFilt <- TCGAanalyze_Filtering(tabDF = dataBRCA, method = "quantile", qnt.cut = 0.25)
samplesNT <- TCGAquery_SampleTypes(colnames(dataFilt), typesample = c("NT"))
samplesTP <- TCGAquery_SampleTypes(colnames(dataFilt), typesample = c("TP"))
```

```

dataDEGs <- TCGAanalyze_DEA(
  dataFilt[,samplesNT],
  dataFilt[,samplesTP],
  Cond1type = "Normal",
  Cond2type = "Tumor"
)
dataDEGsFilt <- dataDEGs[abs(dataDEGs$logFC) >= 1,]
dataTP <- dataFilt[,samplesTP]
dataTN <- dataFilt[,samplesNT]
dataDEGsFiltLevel <- TCGAanalyze_LevelTab(
  FC_FDR_table_mRNA = dataDEGsFilt,
  typeCond1 = "Tumor",
  typeCond2 = "Normal",
  TableCond1 = dataTP,
  TableCond2 = dataTN
)

```

TCGAanalyze_networkInference

infer gene regulatory networks

Description

TCGAanalyze_networkInference taking expression data as input, this will return an adjacency matrix of interactions

Usage

```
TCGAanalyze_networkInference(data, optionMethod = "clr")
```

Arguments

data	expression data, genes in columns, samples in rows
optionMethod	inference method, chose from aracne, c3net, clr and mrnet

Value

an adjacent matrix

TCGAanalyze_Normalization

normalization mRNA transcripts and miRNA using EDASeq package.

Description

TCGAanalyze_Normalization allows user to normalize mRNA transcripts and miRNA, using EDASeq package.

Normalization for RNA-Seq Numerical and graphical summaries of RNA-Seq read data. Within-lane normalization procedures to adjust for GC-content effect (or other gene-level effects) on read counts: loess robust local regression, global-scaling, and full-quantile normalization (Risso et al., 2011). Between-lane normalization procedures to adjust for distributional differences between lanes (e.g., sequencing depth): global-scaling and full-quantile normalization (Bullard et al., 2010).

For instance returns all mRNA or miRNA with mean across all samples, higher than the threshold defined quantile mean across all samples.

TCGAanalyze_Normalization performs normalization using following functions from EDASeq

1. EDASeq::newSeqExpressionSet
2. EDASeq::withinLaneNormalization
3. EDASeq::betweenLaneNormalization
4. EDASeq::counts

Usage

```
TCGAanalyze_Normalization(tabDF, geneInfo, method = "geneLength")
```

Arguments

tabDF	Rnaseq numeric matrix, each row represents a gene, each column represents a sample
geneInfo	Information matrix of 20531 genes about geneLength and gcContent. Two objects are provided: TCGAbiolinks::geneInfoHT,TCGAbiolinks::geneInfo
method	is method of normalization such as 'gcContent' or 'geneLength'

Value

Rnaseq matrix normalized with counts slot holds the count data as a matrix of non-negative integer count values, one row for each observational unit (gene or the like), and one column for each sample.

Examples

```
dataNorm <- TCGAbiolinks::TCGAanalyze_Normalization(dataBRCA, geneInfo)
```

TCGAanalyze_Pathview *Generate pathview graph*

Description

TCGAanalyze_Pathview pathway based data integration and visualization.

Usage

```
TCGAanalyze_Pathview(dataDEGs, pathwayKEGG = "hsa05200")
```

Arguments

dataDEGs	dataDEGs
pathwayKEGG	pathwayKEGG

Value

an adjacent matrix

Examples

```
## Not run:
dataDEGs <- data.frame(mRNA = c("TP53", "TP63", "TP73"), logFC = c(1,2,3))
TCGAanalyze_Pathview(dataDEGs)

## End(Not run)
```

TCGAanalyze_Preprocessing
*Array Array Intensity correlation (AAIC) and correlation boxplot to
define outlier*

Description

TCGAanalyze_Preprocessing perform Array Array Intensity correlation (AAIC). It defines a square symmetric matrix of spearman correlation among samples. According this matrix and boxplot of correlation samples by samples it is possible to find samples with low correlation that can be identified as possible outliers.

Usage

```
TCGAanalyze_Preprocessing(  
  object,  
  cor.cut = 0,  
  datatype = names(assays(object))[1],  
  filename = NULL,  
  width = 1000,  
  height = 1000  
)
```

Arguments

object	gene expression of class RangedSummarizedExperiment from TCGAprepare
cor.cut	is a threshold to filter samples according their spearman correlation in samples by samples. default cor.cut is 0
datatype	is a string from RangedSummarizedExperiment assay
filename	Filename of the image file
width	Image width
height	Image height

Value

Plot with array array intensity correlation and boxplot of correlation samples by samples

TCGAanalyze_Stemness	<i>Generate Stemness Score based on RNASeq (mRNAsi stemness index)</i> <i>Malta et al., Cell, 2018</i>
----------------------	---

Description

TCGAanalyze_Stemness generate the mRNAsi score

Usage

```
TCGAanalyze_Stemness(stemSig, dataGE, colname.score = "stemness_score")
```

Arguments

stemSig	is a vector of the stemness Signature generated using gelnets package. Please check the data from https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5902191/ • SC_PCBC_stemSig - Stemness Score • DE_PCBC_stemSig - endoderm score • EB_PCBC_stemSig - embryoid bodies score • ECTO_PCBC_stemSig - ectoderm score • MESO_PCBC_stemSig - mesoderm score
---------	---

`dataGE` is a matrix of Gene expression (genes in rows, samples in cols) from TCGAprepare

`colname.score` Column name of the output. Default "stemness_score"

Value

table with samples and selected score

Examples

```
# Selecting TCGA breast cancer (10 samples) for example stored in dataBRCA
dataNorm <- TCGAanalyze_Normalization(
  tabDF = dataBRCA,
  geneInfo = geneInfo
)

# quantile filter of genes
dataFilt <- TCGAanalyze_Filtering(
  tabDF = dataNorm,
  method = "quantile",
  qnt.cut = 0.25
)
Stemness_score <- TCGAanalyze_Stemness(
  stemSig = SC_PCBC_stemSig,
  dataGE = dataFilt,
  colname.score = "SC_PCBC_stem_score"
)
ECTO_score <- TCGAanalyze_Stemness(
  stemSig = ECTO_PCBC_stemSig,
  dataGE = dataFilt,
  colname.score = "ECTO_PCBC_stem_score"
)
MESO_score <- TCGAanalyze_Stemness(
  stemSig = MESO_PCBC_stemSig,
  dataGE = dataFilt,
  colname.score = "MESO_PCBC_stem_score"
)
```

TCGAanalyze_survival *Creates survival analysis*

Description

Creates a survival plot from TCGA patient clinical data using survival library. It uses the fields `days_to_death` and `vital`, plus a columns for groups.

Usage

```
TCGAanalyze_survival(
  data,
  clusterCol = NULL,
  legend = "Legend",
  labels = NULL,
  risk.table = TRUE,
  xlim = NULL,
  main = "Kaplan-Meier Overall Survival Curves",
  ylab = "Probability of survival",
  xlab = "Time since diagnosis (days)",
  filename = "survival.pdf",
  color = NULL,
  height = 8,
  width = 12,
  dpi = 300,
  pvalue = TRUE,
  conf.int = TRUE,
  ...
)
```

Arguments

<code>data</code>	TCGA Clinical patient with the information <code>days_to_death</code>
<code>clusterCol</code>	Column with groups to plot. This is a mandatory field, the caption will be based in this column
<code>legend</code>	Legend title of the figure
<code>labels</code>	labels of the plot
<code>risk.table</code>	show or not the risk table
<code>xlim</code>	x axis limits e.g. <code>xlim = c(0, 1000)</code> . Present narrower X axis, but not affect survival estimates.
<code>main</code>	main title of the plot
<code>ylab</code>	y axis text of the plot
<code>xlab</code>	x axis text of the plot
<code>filename</code>	The name of the pdf file.
<code>color</code>	Define the colors/Palette for lines.
<code>height</code>	Image height
<code>width</code>	Image width
<code>dpi</code>	Figure quality
<code>pvalue</code>	show p-value of log-rank test
<code>conf.int</code>	show confidence intervals for point estimates of survival curves.
<code>...</code>	Further arguments passed to ggsurvplot .

Value

Survival plot

Examples

```
# clin <- GDCquery_clinic("TCGA-BRCA","clinical")
clin <- data.frame(
  vital_status = c("alive","alive","alive","dead","alive",
    "alive","dead","alive","dead","alive"),
  days_to_death = c(NA,NA,NA,172,NA,NA,3472,NA,786,NA),
  days_to_last_follow_up = c(3011,965,718,NA,1914,423,NA,5,656,1417),
  gender = c(rep("male",5),rep("female",5))
)
TCGAanalyze_survival(clin, clusterCol="gender")
TCGAanalyze_survival(clin, clusterCol="gender", xlim = 1000)
TCGAanalyze_survival(clin,
  clusterCol="gender",
  risk.table = FALSE,
  conf.int = FALSE,
  color = c("pink","blue"))
TCGAanalyze_survival(clin,
  clusterCol="gender",
  risk.table = FALSE,
  xlim = c(100,1000),
  conf.int = FALSE,
  color = c("Dark2"))
```

TCGAanalyze_SurvivalKM

survival analysis (SA) univariate with Kaplan-Meier (KM) method.

Description

TCGAanalyze_SurvivalKM perform an univariate Kaplan-Meier (KM) survival analysis (SA). It performed Kaplan-Meier survival univariate using complete follow up with all days taking one gene a time from Genelist of gene symbols. For each gene according its level of mean expression in cancer samples, defining two thresholds for quantile expression of that gene in all samples (default ThreshTop=0.67,ThreshDown=0.33) it is possible to define a threshold of intensity of gene expression to divide the samples in 3 groups (High, intermediate, low). TCGAanalyze_SurvivalKM performs SA between High and low groups using following functions from survival package

1. survival::Surv
2. survival::survdiff
3. survival::survfit

Usage

```
TCGAanalyze_SurvivalKM(
  clinical_patient,
  dataGE,
  Genelist,
  Survresult = FALSE,
  ThreshTop = 0.67,
  ThreshDown = 0.33,
  p.cut = 0.05,
  group1,
  group2
)
```

Arguments

<code>clinical_patient</code>	is a data.frame using function 'clinic' with information related to barcode / samples such as bcr_patient_barcode, days_to_death , days_to_last_follow_up , vital_status, etc
<code>dataGE</code>	is a matrix of Gene expression (genes in rows, samples in cols) from TCGAprepare
<code>Genelist</code>	is a list of gene symbols where perform survival KM.
<code>Survresult</code>	is a parameter (default = FALSE) if is TRUE will show KM plot and results.
<code>ThreshTop</code>	is a quantile threshold to identify samples with high expression of a gene
<code>ThreshDown</code>	is a quantile threshold to identify samples with low expression of a gene
<code>p.cut</code>	p.values threshold. Default: 0.05
<code>group1</code>	a string containing the barcode list of the samples in in control group
<code>group2</code>	a string containing the barcode list of the samples in in disease group

Value

table with survival genes pvalues from KM.

Examples

```
# Selecting only 20 genes for example
dataBRCAcomplete <- log2(dataBRCA[1:20,] + 1)

# clinical_patient_Cancer <- GDCquery_clinic("TCGA-BRCA","clinical")
clinical_patient_Cancer <- data.frame(
  bcr_patient_barcode = substr(colnames(dataBRCAcomplete),1,12),
  vital_status = c(rep("alive",3),"dead",rep("alive",2),rep(c("dead","alive"),2)),
  days_to_death = c(NA,NA,NA,172,NA,NA,3472,NA,786,NA),
  days_to_last_follow_up = c(3011,965,718,NA,1914,423,NA,5,656,1417)
)

group1 <- TCGAquery_SampleTypes(colnames(dataBRCAcomplete), typesample = c("NT"))
```

```

group2 <- TCGAquery_SampleTypes(colnames(dataBRCAcomplete), typesample = c("TP"))

tabSurvKM <- TCGAanalyze_SurvivalKM(clinical_patient_Cancer,
                                   dataBRCAcomplete,
                                   Genelist = rownames(dataBRCAcomplete),
                                   Survresult = FALSE,
                                   p.cut = 0.4,
                                   ThreshTop = 0.67,
                                   ThreshDown = 0.33,
                                   group1 = group1, # Control group
                                   group2 = group2) # Disease group

# If the groups are not specified group1 == group2 and all samples are used
## Not run:
tabSurvKM <- TCGAanalyze_SurvivalKM(clinical_patient_Cancer,
                                   dataBRCAcomplete,
                                   Genelist = rownames(dataBRCAcomplete),
                                   Survresult = TRUE,
                                   p.cut = 0.2,
                                   ThreshTop = 0.67,
                                   ThreshDown = 0.33)

## End(Not run)

```

TCGAbatch_Correction *Batch correction using ComBat and Voom transformation using limma package.*

Description

TCGAbatch_correction allows user to perform a Voom correction on gene expression data and have it ready for DEA. One can also use ComBat for batch correction for exploratory analysis. If batch.factor or adjustment argument is "Year" please provide clinical data. If no batch factor is provided, the data will be voom corrected only

TCGAanalyze_DEA performs DEA using following functions from sva and limma:

1. limma::voom Transform RNA-Seq Data Ready for Linear Modelling.
2. sva::ComBat Adjust for batch effects using an empirical Bayes framework.

Usage

```

TCGAbatch_Correction(
  tabDF,
  batch.factor = NULL,
  adjustment = NULL,
  ClinicalDF = data.frame(),
  UnpublishedData = FALSE,
  AnnotationDF = data.frame()
)

```

Arguments

tabDF	numeric matrix, each row represents a gene, each column represents a sample
batch.factor	a string containing the batch factor to use for correction. Options are "Plate", "TSS", "Year", "Portion", "Center"
adjustment	vector containing strings for factors to adjust for using ComBat. Options are "Plate", "TSS", "Year", "Portion", "Center"
ClinicalDF	a dataframe returned by GDCquery_clinic() to be used to extract year data
UnpublishedData	if TRUE perform a batch correction after adding new data
AnnotationDF	a dataframe with column Batch indicating different batches of the samples in the tabDF

Value

data frame with ComBat batch correction applied

TCGAbiolinks	<i>The aim of TCGAbiolinks is : i) facilitate the TCGA open-access data retrieval, ii) prepare the data using the appropriate pre-processing strategies, iii) provide the means to carry out different standard analyses and iv) allow the user to download a specific version of the data and thus to easily reproduce earlier research results. In more detail, the package provides multiple methods for analysis (e.g., differential expression analysis, identifying differentially methylated regions) and methods for visualization (e.g., survival plots, volcano plots, starburst plots) in order to easily develop complete analysis pipelines.</i>
--------------	---

Description

The functions you’re likely to need from **TCGAbiolinks** is [GDCdownload](#), [GDCquery](#). Otherwise refer to the vignettes to see how to format the documentation.

TCGApprepare_Affy	<i>Prepare CEL files into an AffyBatch.</i>
-------------------	---

Description

Prepare CEL files into an AffyBatch.

Usage

TCGApprepare_Affy(ClinData, PathFolder, TabCel)

Arguments

ClinData	write
PathFolder	write
TabCel	write

Value

Normalized Expression data from Affy eSets

Examples

```
## Not run:
to add example

## End(Not run)
```

TCGAquery_MatchedCoupledSampleTypes

Retrieve multiple tissue types from the same patients.

Description

TCGAquery_MatchedCoupledSampleTypes

Usage

```
TCGAquery_MatchedCoupledSampleTypes(barcode, typesample)
```

Arguments

barcode	barcode
typesample	typesample

Value

a list of samples / barcode filtered by type sample selected

Examples

```
TCGAquery_MatchedCoupledSampleTypes(c("TCGA-B0-4698-01Z-00-DX1",
                                         "TCGA-B0-4698-02Z-00-DX1"),
                                         c("TP", "TR"))
barcode <- c("TARGET-20-PANSBH-02A-02D", "TARGET-20-PANSBH-01A-02D",
            "TCGA-B0-4698-01Z-00-DX1", "TCGA-CZ-4863-02Z-00-DX1",
            "TARGET-20-PANSZZ-02A-02D", "TARGET-20-PANSZZ-11A-02D",
            "TCGA-B0-4699-01Z-00-DX1", "TCGA-B0-4699-02Z-00-DX1"
            )
TCGAquery_MatchedCoupledSampleTypes(barcode, c("TR", "TP"))
```

TCGAquery_recount2 *Query gene counts of TCGA and GTEx data from the Recount2 project*

Description

TCGARecount2_query queries and downloads data produced by the Recount2 project. User can specify which project and which tissue to query

Usage

```
TCGAquery_recount2(project, tissue = c())
```

Arguments

project	is a string denoting which project the user wants. Options are "tcga" and "gtex"
tissue	a vector of tissue(s) to download. Options are "adipose tissue", "adrenal gland", "bladder", "blood", "blood vessel", "bone marrow", "brain", "breast", "cervix uteri", "colon", "esophagus", "fallopian tube", "heart", "kidney", "liver", "lung", "muscle", "nerve", "ovary", "pancreas", "pituitary", "prostate", "salivary gland", "skin", "small intestine", "spleen", "stomach", "testis", "thyroid", "uterus", "vagina"

Value

List with \$subtypes attribute as a dataframe with barcodes, samples, subtypes, and colors. The \$filtered attribute is returned as filtered samples with no subtype info

Examples

```
## Not run:
brain.rec<-TCGAquery_recount2(project = "gtex", tissue = "brain")

## End(Not run)
```

TCGAquery_SampleTypes *Retrieve multiple tissue types not from the same patients.*

Description

TCGAquery_SampleTypes for a given list of samples and types, return the union of samples that are from theses type.

Usage

```
TCGAquery_SampleTypes(barcode, typesample)
```

Arguments

barcode is a list of samples as TCGA barcodes
typesample a character vector indicating tissue type to query. Example:

TP	PRIMARY SOLID TUMOR
TR	RECURRENT SOLID TUMOR
TB	Primary Blood Derived Cancer-Peripheral Blood
TRBM	Recurrent Blood Derived Cancer-Bone Marrow
TAP	Additional-New Primary
TM	Metastatic
TAM	Additional Metastatic
THOC	Human Tumor Original Cells
TBM	Primary Blood Derived Cancer-Bone Marrow
NB	Blood Derived Normal
NT	Solid Tissue Normal
NBC	Buccal Cell Normal
NEBV	EBV Immortalized Normal
NBM	Bone Marrow Normal

Value

a list of samples / barcode filtered by type sample selected

Examples

```
# selection of normal samples "NT"
barcode <- c("TCGA-B0-4698-01Z-00-DX1", "TCGA-CZ-4863-02Z-00-DX1")
# Returns the second barcode
TCGAquery_SampleTypes(barcode, "TR")
# Returns both barcode
TCGAquery_SampleTypes(barcode, c("TR", "TP"))
barcode <- c("TARGET-20-PANSBH-14A-02D", "TARGET-20-PANSBH-01A-02D",
            "TCGA-B0-4698-01Z-00-DX1", "TCGA-CZ-4863-02Z-00-DX1")
TCGAquery_SampleTypes(barcode, c("TR", "TP"))
```

TCGAquery_subtype	<i>Retrieve molecular subtypes for a given tumor</i>
-------------------	--

Description

TCGAquery_subtype Retrieve molecular subtypes for a given tumor

Usage

```
TCGAquery_subtype(tumor)
```

Arguments

tumor is a cancer Examples:
lgg gbm luad stad brca
coad read

Value

a data.frame with barcode and molecular subtypes

Examples

```
dataSubt <- TCGAquery_subtype(tumor = "lgg")
```

TCGAumor_purity	<i>Filters TCGA barcodes according to purity parameters</i>
-----------------	---

Description

TCGAumor_purity Filters TCGA samples using 5 estimates from 5 methods as thresholds.

Usage

```
TCGAumor_purity(barcodes, estimate, absolute, lump, ihc, cpe)
```

Arguments

barcodes is a vector of TCGA barcodes

estimate uses gene expression profiles of 141 immune genes and 141 stromal genes

absolute which uses somatic copy-number data (estimations were available for only 11 cancer types)

lump (leukocytes unmethylation for purity), which averages 44 non-methylated immune-specific CpG sites

ihc as estimated by image analysis of haematoxylin and eosin stain slides produced by the Nationwide Childrens Hospital Biospecimen Core Resource

cpe CPE is a derived consensus measurement as the median purity level after normalizing levels from all methods to give them equal means and s.ds

Value

List with \$pure_barcodes attribute as a vector of pure samples and \$filtered attribute as filtered samples with no purity info

Examples

```
dataTableSubt <- TCGAtumor_purity("TCGA-60-2721-01A-01R-0851-07",
                                   estimate = 0.6,
                                   absolute = 0.6,
                                   ihc = 0.8,
                                   lump = 0.8,
                                   cpe = 0.7)
```

TCGAvisualize_BarPlot *Barplot of subtypes and clinical info in groups of gene expression clustered.*

Description

Barplot of subtypes and clinical info in groups of gene expression clustered.

Usage

```
TCGAvisualize_BarPlot(
  DFfilt,
  DFclin,
  DFsubt,
  data_Hc2,
  Subtype,
  cbPalette,
  filename,
  width,
  height,
  dpi
)
```

Arguments

DFfilt	write
DFclin	write
DFsubt	write
data_Hc2	write
Subtype	write
cbPalette	Define the colors of the bar.
filename	The name of the pdf file
width	Image width
height	Image height
dpi	Image dpi

Value

barplot image in pdf or png file

TCGAvisualize_EAbarplot

barPlot for a complete Enrichment Analysis

Description

The figure shows canonical pathways significantly overrepresented (enriched) by the DEGs (differentially expressed genes). The most statistically significant canonical pathways identified in DEGs list are listed according to their p value corrected FDR (-Log) (colored bars) and the ratio of list genes found in each pathway over the total number of genes in that pathway (Ratio, red line).

Usage

```
TCGAvisualize_EAbarplot(
  tf,
  GOMFTab,
  GOBPTab,
  GOCCTab,
  PathTab,
  nBar,
  nRGTAB,
  filename = "TCGAvisualize_EAbarplot_Output.pdf",
  text.size = 1,
  mfrow = c(2, 2),
  xlim = NULL,
  fig.width = 30,
  fig.height = 15,
  color = c("orange", "cyan", "green", "yellow")
)
```

Arguments

tf	is a list of gene symbols
GOMFTab	is results from TCGAanalyze_EAcomplete related to Molecular Function (MF)
GOBPTab	is results from TCGAanalyze_EAcomplete related to Biological Process (BP)
GOCCTab	is results from TCGAanalyze_EAcomplete related to Cellular Component (CC)
PathTab	is results from TCGAanalyze_EAcomplete related to Pathways EA
nBar	is the number of bar histogram selected to show (default = 10)
nRGTAB	is the gene signature list with gene symbols.
filename	Name for the pdf. If null it will return the plot.
text.size	Text size
mfrow	Vector with number of rows/columns of the plot. Default 2 rows/2 columns "c(2,2)"
xlim	Upper limit of the x-axis.

fig.width	Default 30
fig.height	Default 15
color	A vector of colors for each barplot. Deafult: c("orange", "cyan", "green", "yellow")

Value

Complete barPlot from Enrichment Analysis showing significant (default FDR < 0.01) BP,CC,MF and pathways enriched by list of genes.

Examples

```
Genelist <- c("FN1", "COL1A1")
ansEA <- TCGAanalyze_EAcomplete(TFname="DEA genes Normal Vs Tumor", Genelist)
TCGAvsualize_EAbarplot(tf = rownames(ansEA$ResBP),
  GOBPTab = ansEA$ResBP,
  GOCCTab = ansEA$ResCC,
  GOMFTab = ansEA$ResMF,
  PathTab = ansEA$ResPat,
  nRGTab = Genelist,
  nBar = 10,
  filename="a.pdf")
## Not run:
Genelist <- rownames(dataDEGsFiltLevel)
system.time(ansEA <- TCGAanalyze_EAcomplete(TFname="DEA genes Normal Vs Tumor", Genelist))
# Enrichment Analysis EA (TCGAvsualize)
# Gene Ontology (GO) and Pathway enrichment barPlot
TCGAvsualize_EAbarplot(tf = rownames(ansEA$ResBP),
  GOBPTab = ansEA$ResBP,
  GOCCTab = ansEA$ResCC,
  GOMFTab = ansEA$ResMF,
  PathTab = ansEA$ResPat,
  nRGTab = Genelist,
  nBar = 10)

## End(Not run)
```

TCGAvsualize_Heatmap *Heatmap with more sensible behavior using heatmap.plus*

Description

Heatmap with more sensible behavior using heatmap.plus

Usage

```
TCGAvsualize_Heatmap(
  data,
  col.metadata,
  row.metadata,
```

```

col.colors = NULL,
row.colors = NULL,
show_column_names = FALSE,
show_row_names = FALSE,
cluster_rows = FALSE,
cluster_columns = FALSE,
sortCol,
extremes = NULL,
rownames.size = 12,
title = NULL,
color.levels = NULL,
values.label = NULL,
filename = "heatmap.pdf",
width = 10,
height = 10,
type = "expression",
scale = "none",
heatmap.legend.color.bar = "continuous"
)

```

Arguments

data	The object to with the heatmap data (expression, methylation)
col.metadata	Metadata for the columns (samples). It should have on of the following columns: barcode (28 characters) column to match with the samples. It will also work with "bcr_patient_barcode"(12 chars),"patient"(12 chars),"sample"(16 chars) columns but as one patient might have more than one sample, this could lead to errors in the annotation. The code will throw a warning in case two samples are from the same patient.
row.metadata	Metadata for the rows genes (expression) or probes (methylation)
col.colors	A list of names colors
row.colors	A list of named colors
show_column_names	Show column names names? Default: FALSE
show_row_names	Show row names? Default: FALSE
cluster_rows	Cluster rows ? Default: FALSE
cluster_columns	Cluster columns ? Default: FALSE
sortCol	Name of the column to be used to sort the columns
extremes	Extremes of colors (vector of 3 values)
rownames.size	Rownames size
title	Title of the plot
color.levels	A vector with the colors (low level, middle level, high level)
values.label	Text of the levels in the heatmap
filename	Filename to save the heatmap. Default: heatmap.png


```

                                "IDHMut-cod"="tomato"
                                , "IDHMut-noncod"="gold")),
type = "methylation",
show_row_names=TRUE)

```

TCGAvsualize_meanMethylation

Mean methylation boxplot

Description

Creates a mean methylation boxplot for groups (groupCol), subgroups will be highlighted as shapes if the subgroupCol was set.

Observation: Data is a summarizedExperiment.

Usage

```

TCGAvsualize_meanMethylation(
  data,
  groupCol = NULL,
  subgroupCol = NULL,
  shapes = NULL,
  print.pvalue = FALSE,
  plot.jitter = TRUE,
  jitter.size = 3,
  filename = "groupMeanMet.pdf",
  ylab = expression(paste("Mean DNA methylation (", beta, "-values)")),
  xlab = NULL,
  title = "Mean DNA methylation",
  labels = NULL,
  group.legend = NULL,
  subgroup.legend = NULL,
  color = NULL,
  y.limits = NULL,
  sort,
  order,
  legend.position = "top",
  legend.title.position = "top",
  legend.ncols = 3,
  add.axis.x.text = TRUE,
  width = 10,
  height = 10,
  dpi = 600,
  axis.text.x.angle = 90
)

```

Arguments

<code>data</code>	SummarizedExperiment object obtained from TCGAPrep
<code>groupCol</code>	Columns in <code>colData(data)</code> that defines the groups. If no columns defined a columns called "Patients" will be used
<code>subgroupCol</code>	Columns in <code>colData(data)</code> that defines the subgroups.
<code>shapes</code>	Shape vector of the subgroups. It must have the size of the levels of the subgroups. Example: <code>shapes = c(21,23)</code> if for two levels
<code>print.pvalue</code>	Print p-value for two groups
<code>plot.jitter</code>	Plot jitter? Default TRUE
<code>jitter.size</code>	Plot jitter size? Default 3
<code>filename</code>	The name of the pdf that will be saved
<code>ylab</code>	y axis text in the plot
<code>xlab</code>	x axis text in the plot
<code>title</code>	main title in the plot
<code>labels</code>	Labels of the groups
<code>group.legend</code>	Name of the group legend. DEFAULT: <code>groupCol</code>
<code>subgroup.legend</code>	Name of the subgroup legend. DEFAULT: <code>subgroupCol</code>
<code>color</code>	vector of colors to be used in graph
<code>y.limits</code>	Change lower/upper y-axis limit
<code>sort</code>	Sort boxplot by mean or median. Possible values: <code>mean.asc</code> , <code>mean.desc</code> , <code>median.asc</code> , <code>median.desc</code>
<code>order</code>	Order of the boxplots
<code>legend.position</code>	Legend position ("top", "right", "left", "bottom")
<code>legend.title.position</code>	Legend title position ("top", "right", "left", "bottom")
<code>legend.ncols</code>	Number of columns of the legend
<code>add.axis.x.text</code>	Add text to x-axis? Default: FALSE
<code>width</code>	Plot width default:10
<code>height</code>	Plot height default:10
<code>dpi</code>	Pdf dpi default:600
<code>axis.text.x.angle</code>	Angle of text in the x axis

Value

Save the pdf survival plot

Examples

```
nrows <- 200; ncols <- 21
counts <- matrix(runif(nrows * ncols, 0, 1), nrows)
rowRanges <- GenomicRanges::GRanges(rep(c("chr1", "chr2"), c(50, 150)),
                                     IRanges::IRanges(floor(runif(200, 1e5, 1e6)), width=100),
                                     strand=sample(c("+", "-"), 200, TRUE),
                                     feature_id=sprintf("ID%03d", 1:200))
colData <- S4Vectors::DataFrame(Treatment=rep(c("ChIP", "Input", "Other"), 7),
                                row.names=LETTERS[1:21],
                                group=rep(c("group1", "group2", "group3"), c(7,7,7)),
                                subgroup=rep(c("subgroup1", "subgroup2", "subgroup3"), 7))
data <- SummarizedExperiment::SummarizedExperiment(
  assays=S4Vectors::SimpleList(counts=counts),
  rowRanges=rowRanges,
  colData=colData)
TCGAvsualize_meanMethylation(data, groupCol = "group")
# change lower/upper y-axis limit
TCGAvsualize_meanMethylation(data, groupCol = "group", y.limits = c(0,1))
# change lower y-axis limit
TCGAvsualize_meanMethylation(data, groupCol = "group", y.limits = 0)
TCGAvsualize_meanMethylation(data, groupCol = "group", subgroupCol="subgroup")
TCGAvsualize_meanMethylation(data, groupCol = "group")
TCGAvsualize_meanMethylation(data, groupCol = "group", sort="mean.desc", filename="meandesc.pdf")
TCGAvsualize_meanMethylation(data, groupCol = "group", sort="mean.asc", filename="meanasc.pdf")
TCGAvsualize_meanMethylation(data, groupCol = "group", sort="median.asc", filename="medianasc.pdf")
TCGAvsualize_meanMethylation(data, groupCol = "group", sort="median.desc", filename="mediandesc.pdf")
```

TCGAvsualize_oncoprint

Creating a oncoprint

Description

Creating a oncoprint

Usage

```
TCGAvsualize_oncoprint(
  mut,
  genes,
  filename,
  color,
  annotation.position = "bottom",
  annotation,
  height,
  width = 10,
  rm.empty.columns = FALSE,
  show.column.names = FALSE,
```

```

show.row.barplot = TRUE,
label.title = "Mutation",
column.names.size = 8,
label.font.size = 16,
rows.font.size = 16,
dist.col = 0.5,
dist.row = 0.5,
information = "Variant_Type",
row.order = TRUE,
col.order = TRUE,
heatmap.legend.side = "bottom",
annotation.legend.side = "bottom"
)

```

Arguments

mut	A dataframe from the mutation annotation file (see TCGAquery_maf from TCGA Biobioinformatics)
genes	Gene list
filename	name of the pdf
color	named vector for the plot
annotation.position	Position of the annotation "bottom" or "top"
annotation	Matrix or data frame with the annotation. Should have a column bcr_patient_barcode with the same ID of the mutation object
height	pdf height
width	pdf width
rm.empty.columns	If there is no alteration in that sample, whether remove it on the oncoprint
show.column.names	Show column names? Default: FALSE
show.row.barplot	Show barplot annotation on rows?
label.title	Title of the label
column.names.size	Size of the fonts of the columns names
label.font.size	Size of the fonts
rows.font.size	Size of the fonts
dist.col	distance between columns in the plot
dist.row	distance between rows in the plot
information	Which column to use as information from MAF. Options: 1) "Variant_Classification" (The information will be "Frame_Shift_Del", "Frame_Shift_Ins", "In_Frame_Del", "In_Frame_Ins", "Missense_Mutation", "Nonsense_Mutation", "Nonstop_Mutation", "RNA", "Silent", "Splice_Site", "Targeted_Region", "Translation_Start_Site") 2) "Variant_Type" (The information will be INS,DEL,SNP)

row.order	Order the genes (rows) Default:TRUE. Genes with more mutations will be in the first rows
col.order	Order columns. Default:TRUE.
heatmap.legend.side	Position of the heatmap legend
annotation.legend.side	Position of the annotation legend

Value

A oncoprint plot

Examples

```
## Not run:
library(maftools)
library(dplyr)
query <- GDCquery(
  project = "TCGA-CHOL",
  data.category = "Simple Nucleotide Variation",
  access = "open",
  legacy = FALSE,
  data.type = "Masked Somatic Mutation",
  workflow.type = "Aliquot Ensemble Somatic Variant Merging and Masking"
)
GDCdownload(query)
mut <- GDCprepare(query)
TCGAvsualize_oncprint(mut = mut, genes = mut$Hugo_Symbol[1:10], rm.empty.columns = TRUE)
TCGAvsualize_oncprint(mut = mut, genes = mut$Hugo_Symbol[1:10],
  filename = "onco.pdf",
  color=c("background"="#CCCCCC", "DEL"="purple", "INS"="yellow", "SNP"="brown"))
clin <- GDCquery_clinic("TCGA-ACC", "clinical")
clin <- clin[,c("bcr_patient_barcode", "disease", "gender", "tumor_stage", "race", "vital_status")]
TCGAvsualize_oncprint(mut = mut, genes = mut$Hugo_Symbol[1:20],
  filename = "onco.pdf",
  annotation = clin,
  color=c("background"="#CCCCCC", "DEL"="purple", "INS"="yellow", "SNP"="brown"),
  rows.font.size=10,
  heatmap.legend.side = "right",
  dist.col = 0,
  label.font.size = 10)

## End(Not run)
```

Description

TCGAvisualize_PCA performs a principal components analysis (PCA) on the given data matrix and returns the results as an object of class prcomp, and shows results in PCA level.

Usage

```
TCGAvisualize_PCA(dataFilt, dataDEGsFiltLevel, ntopgenes, group1, group2)
```

Arguments

dataFilt	A filtered dataframe or numeric matrix where each row represents a gene, each column represents a sample from function TCGAanalyze_Filtering
dataDEGsFiltLevel	table with DEGs, log Fold Change (FC), false discovery rate (FDR), the gene expression level, etc, from function TCGAanalyze_LevelTab.
ntopgenes	number of DEGs genes to plot in PCA
group1	a string containing the barcode list of the samples in in control group
group2	a string containing the barcode list of the samples in in disease group the name of the group

Value

principal components analysis (PCA) plot of PC1 and PC2

Examples

```
# normalization of genes
dataNorm <- TCGAanalyze_Normalization(tabDF = dataBRCA, geneInfo = geneInfo,
method = "geneLength")
# quantile filter of genes
dataFilt <- TCGAanalyze_Filtering(tabDF = dataBRCA, method = "quantile", qnt.cut = 0.25)
# Principal Component Analysis plot for ntop selected DEGs
# selection of normal samples "NT"
group1 <- TCGAquery_SampleTypes(colnames(dataFilt), typesample = c("NT"))
# selection of normal samples "TP"
group2 <- TCGAquery_SampleTypes(colnames(dataFilt), typesample = c("TP"))
pca <- TCGAvisualize_PCA(dataFilt, dataDEGsFiltLevel, ntopgenes = 200, group1, group2)
```

TCGAvisualize_starburst

Create starburst plot

Description

Create Starburst plot for comparison of DNA methylation and gene expression. The log10 (FDR-corrected P value) is plotted for beta value for DNA methylation (x axis) and gene expression (y axis) for each gene.

The black dashed line shows the FDR-adjusted P value of 0.01.

You can set names to TRUE to get the names of the significant genes.

Candidate biologically significant genes will be circled in the plot.

Candidate biologically significant are the genes that respect the expression (logFC.cut), DNA methylation (diffmean.cut) and significance thresholds (exp.p.cut, met.p.cut)

Usage

```
TCGAvisualize_starburst(
  met,
  exp,
  group1 = NULL,
  group2 = NULL,
  exp.p.cut = 0.01,
  met.p.cut = 0.01,
  diffmean.cut = 0,
  logFC.cut = 0,
  met.platform = c("Illumina Human Methylation 450", "Illumina Human Methylation 27",
    "Illumina Methylation Epic"),
  genome,
  names = FALSE,
  names.fill = TRUE,
  filename = "starburst.png",
  return.plot = FALSE,
  ylab = expression(atop("Gene Expression", paste(-Log[10],
    " (FDR corrected P values)"))),
  xlab = expression(atop("DNA Methylation", paste(-Log[10],
    " (FDR corrected P values)"))),
  title = "Starburst Plot",
  legend = "DNA Methylation/Expression Relation",
  color = NULL,
  label = c("Not Significant", "Up regulated & Hypo methylated",
    "Down regulated & Hypo methylated", "hypo methylated", "hyper methylated",
    "Up regulated", "Down regulated", "Up regulated & Hyper methylated",
    "Down regulated & Hyper methylated"),
  xlim = NULL,
  ylim = NULL,
  height = 10,
  width = 20,
  dpi = 600
)
```

Arguments

met	A SummarizedExperiment with methylation data obtained from the TCGAPrepare or Data frame from DMR_results file. Expected colData columns: diffmean, p.value.adj and p.value Execute volcanoPlot function in order to obtain these values for the object.
exp	Object obtained by DEArnaSEQ function
group1	The name of the group 1 Obs: Column p.value.adj.group1.group2 should exist
group2	The name of the group 2. Obs: Column p.value.adj.group1.group2 should exist
exp.p.cut	expression p value cut-off
met.p.cut	methylation p value cut-off
diffmean.cut	If set, the probes with diffmean higher than methylation cut-off will be highlighted in the plot. And the data frame return will be subseted.
logFC.cut	If set, the probes with expression fold change higher than methylation cut-off will be highlighted in the plot. And the data frame return will be subseted.
met.platform	DNA methylation platform "Illumina Human Methylation 450", "Illumina Human Methylation 27", "Illumina Methylation Epic"
genome	Genome of reference ("hg38" or "hg19") used to identify nearest probes TSS
names	Add the names of the significant genes? Default: FALSE
names.fill	Names should be filled in a color box? Default: TRUE
filename	The filename of the file (it can be pdf, svg, png, etc)
return.plot	If true only plot object will be returned (pdf will not be created)
ylab	y axis text
xlab	x axis text
title	main title
legend	legend title
color	vector of colors to be used in graph
label	vector of labels to be used in graph
xlim	x limits to cut image
ylim	y limits to cut image
height	Figure height
width	Figure width
dpi	Figure dpi

Details

Input: data with gene expression/methylation expression Output: starburst plot

Value

Save a starburst plot

Examples

```
## Not run:
library(SummarizedExperiment)
met <- TCGAbiolinks::getMetPlatInfo(
  genome = "hg38",
  platform = "Illumina Human Methylation 27"
)
values(met) <- NULL
met$probeID <- names(met)
nrows <- length(met); ncols <- 20
counts <- matrix(runif(nrows * ncols, 1, 1e4), nrows)
colData <- S4Vectors::DataFrame(
  Treatment = rep(c("ChIP", "Input"), 5),
  row.names = LETTERS[1:20],
  group = rep(c("group1", "group2"), c(10, 10))
)
met <- SummarizedExperiment::SummarizedExperiment(
  assays = S4Vectors::SimpleList(counts=counts),
  rowRanges = met,
  colData = colData
)
rowRanges(met)$diffmean.g1.g2 <- c(runif(nrows, -0.1, 0.1))
rowRanges(met)$diffmean.g2.g1 <- -1*(rowRanges(met)$diffmean.g1.g2)
rowRanges(met)$p.value.g1.g2 <- c(runif(nrows, 0, 1))
rowRanges(met)$p.value.adj.g1.g2 <- c(runif(nrows, 0, 1))
exp <- TCGAbiolinks::get.GRCh.bioMart("hg38")
exp$logFC <- runif(nrow(exp), -5, 5)
exp$FDR <- runif(nrow(exp), 0.01, 1)

result <- TCGAvisualize_starburst(
  met,
  exp,
  exp.p.cut = 0.05,
  met.p.cut = 0.05,
  logFC.cut = 2,
  group1 = "g1",
  group2 = "g2",
  genome = "hg38",
  met.platform = "27k",
  diffmean.cut = 0.0,
  names = TRUE
)

## End(Not run)
```

Description

TCGAvisualize_SurvivalCoxNET can help an user to identify a group of survival genes that are significant from univariate Kaplan Meier Analysis and also for Cox Regression. It shows in the end a network build with community of genes with similar range of pvalues from Cox regression (same color) and that interaction among those genes is already validated in literatures using the STRING database (version 9.1). TCGAvisualize_SurvivalCoxNET perform survival analysis with univariate Cox regression and package (dnet) using following functions wrapping from these packages:

1. survival::coxph
2. igraph::subgraph.edges
3. igraph::layout.fruchterman.reingold
4. igraph::spinglass.community
5. igraph::communities
6. dnet::dRDataLoader
7. dnet::dNetInduce
8. dnet::dNetPipeline
9. dnet::visNet
10. dnet::dCommSignif

Usage

```
TCGAvisualize_SurvivalCoxNET(
  clinical_patient,
  dataGE,
  Genelist,
  org.Hs.string,
  scoreConfidence = 700,
  titlePlot = "TCGAvisualize_SurvivalCoxNET Example"
)
```

Arguments

clinical_patient	is a data.frame using function 'clinic' with information related to barcode / samples such as bcr_patient_barcode, days_to_death , days_to_last_followup , vital_status, etc
dataGE	is a matrix of Gene expression (genes in rows, samples in cols) from TCGAprepare
Genelist	is a list of gene symbols where perform survival KM.
org.Hs.string	an igraph object that contains a functional protein association network in human. The network is extracted from the STRING database (version 10).
scoreConfidence	restrict to those edges with high confidence (eg. score>=700)
titlePlot	is the title to show in the final plot.

Details

TCGAVisualize_SurvivalCoxNET allow user to perform the complete workflow using coxph and dnet package related to survival analysis with an identification of gene-active networks from high-throughput omics data using gene expression and clinical data.

1. Cox regression survival analysis to obtain hazard ratio (HR) and p-values
2. fit a Cox proportional hazards model and ANOVA (Chisq test)
3. Network communities
4. An igraph object that contains a functional protein association network in human. The network is extracted from the STRING database (version 9.1). Only those associations with medium confidence (score $\geq$ 400) are retained.
5. restrict to those edges with high confidence (score $\geq$ 700)
6. extract network that only contains genes in pvals
7. Identification of gene-active network
8. visualisation of the gene-active network itself
9. the layout of the network visualisation (fixed in different visuals)
10. color nodes according to communities (identified via a spin-glass model and simulated annealing)
11. node sizes according to degrees
12. highlight different communities
13. visualize the subnetwork

Value

net IGRAPH with related Cox survival genes in community (same pval and color) and with interactions from STRING database.

TCGAVisualize_volcano *Creates a volcano plot for DNA methylation or gene expression*

Description

Creates a volcano plot from the gene expression and DNA methylation analysis.

Usage

```
TCGAVisualize_volcano(
  x,
  y,
  filename = "volcano.pdf",
  ylab = expression(paste(-Log[10], " (FDR corrected P-values)")),
  xlab = NULL,
  title = "Volcano plot",
```

```

legend = NULL,
label = NULL,
xlim = NULL,
ylim = NULL,
color = c("black", "red", "green"),
names = NULL,
names.fill = TRUE,
show.names = "significant",
x.cut = 0,
y.cut = 0.01,
height = 5,
width = 10,
highlight = NULL,
highlight.color = "orange",
names.size = 4,
dpi = 300
)

```

Arguments

x	x-axis data (i.e. Diff mean beta-values or Log2FC).
y	FDR adjusted p-value (q-value). This data will be transformed to -log10 values.
filename	File name: volcano.pdf, volcano.svg, volcano.png. If NULL returns the ggplot object.
ylab	y axis text. Default: -Log10 FDR corrected P-values
xlab	x axis text. Default: No text. Examples of input: expression(paste(Log[2], "FoldChange"))
title	main title. If not specified it will be "Volcano plot (group1 vs group2)
legend	Legend title
label	vector of labels to be used in the figure. Example: c("Not Significant", "Hypermethylated in group1", "Hypomethylated in group1"))#'
xlim	x limits to cut image (i.e. c(-4,4))
ylim	y limits to cut image (i.e. c(-1,10))
color	vector of colors to be used in graph
names	Names to be plotted if significant. Should be the same size of x and y
names.fill	Names should be filled in a color box? Default: TRUE
show.names	What names will be showed? Possibilities: "both", "significant", "highlighted"
x.cut	x-axis threshold. Default: 0.0 If you give only one number (e.g. 0.2) the cut-offs will be -0.2 and 0.2. Or you can give different cut-offs as a vector (e.g. c(-0.3,0.4))
y.cut	q-values threshold (i.e. 0.01, 10^-10)
height	Figure height
width	Figure width

highlight	List of genes/probes to be highlighted. It should be in the names argument.
highlight.color	Color of the points highlighted
names.size	Size of the names text
dpi	Figure dpi

Details

Creates a volcano plot from the gene expression and DNA methylation analysis. Please see the vignette for more information

Value

Saves the volcano plot in the current folder

Examples

```
log2_foldchange <- runif(200, -2, 2)
fdr <- runif(200, 0.01, 1)
TCGAVisualize_volcano(
  x = log2_foldchange,
  y = fdr,
  x.cut = 1.5,
  y.cut = 0.01,
  title = "Title example",
  xlab = expression(paste(Log[2], "FoldChange")))
)
beta_diff <- runif(200, -1, 1)
fdr <- runif(200, 0.01, 1)
TCGAVisualize_volcano(
  x = beta_diff,
  y = fdr,
  x.cut = 1.5,
  y.cut = 0.01,
  title = "Title example",
  xlab = expression(paste("DNA Methylation difference (", beta, "-values)"))
)
## Not run:
TCGAVisualize_volcano(
  x,
  y,
  filename = NULL,
  y.cut = 10000000,
  x.cut=0.8,
  names = rep("AAAA",length(x)),
  legend = "Status",
  names.fill = FALSE
)

TCGAVisualize_volcano(
  x,
```

```

    y,
    filename = NULL,
    y.cut = 10000000,
    x.cut = 0.8,
    names = as.character(1:length(x)),
    legend = "Status",
    names.fill = TRUE, highlight = c("1","2"),
    show = "both"
  )
  TCGAVisualize_volcano(
    x,
    y,
    filename = NULL,
    y.cut = 10000000,
    x.cut = c(-0.3,0.8),
    names = as.character(1:length(x)),
    legend = "Status",
    names.fill = TRUE,
    highlight = c("1","2"),
    show = "both"
  )

  ## End(Not run)
  while (!(is.null(dev.list()["RStudioGD"]))) {dev.off()}

```

TCGA_MolecularSubtype *Retrieve molecular subtypes for given TCGA barcodes*

Description

TCGA_MolecularSubtype Retrieve molecular subtypes from TCGA consortium for a given set of barcodes

Usage

```
TCGA_MolecularSubtype(barcodes)
```

Arguments

barcodes is a vector of TCGA barcodes

Value

List with \$subtypes attribute as a dataframe with barcodes, samples, subtypes, and colors. The \$filtered attribute is returned as filtered samples with no subtype info

Examples

```
TCGA_MolecularSubtype("TCGA-60-2721-01A-01R-0851-07")
```

Tumor.purity	<i>TCGA samples with their Tumor Purity measures</i>
--------------	--

Description

A dataset containing the Sample Ids from TCGA tumor purity measured according to 4 estimates attributes of 9364 tumor patients

Usage

```
Tumor.purity
```

Format

A data frame with 9364 rows and 7 variables:

Sample.ID Sample ID from TCGA barcodes, character string

Cancer.type Cancer type, character string

ESTIMATE uses gene expression profiles of 141 immune genes and 141 stromal genes, 0-1 value

ABSOLUTE uses somatic copy-number data (estimations were available for only 11 cancer types), 0-1 value

LUMP (leukocytes unmethylation for purity), which averages 44 non-methylated immune-specific CpG sites, 0-1 value

IHC as estimated by image analysis of haematoxylin and eosin stain slides produced by the Nationwide Childrens Hospital Biospecimen Core Resource, 0-1 value

CPE derived consensus measurement as the median purity level after normalizing levels from all methods to give them equal means and s.ds, 0-1 value ...

Source

<https://images.nature.com/original/nature-assets/ncomms/2015/151204/ncomms9971/extref/ncomms9971-s2.xlsx>

UseRaw_afterFilter	<i>Use raw count from the DataPrep object which genes are removed by normalization and filtering steps.</i>
--------------------	---

Description

function to keep raw counts after filtering and/or normalizing.

Usage

```
UseRaw_afterFilter(DataPrep, DataFilt)
```

Arguments

DataPrep	DataPrep object returned by TCGAanalyze_Preprocessing()
DataFilt	Filtered data frame containing samples in columns and genes in rows after normalization and/or filtering steps

Value

Filtered return object similar to DataPrep with genes removed after normalization and filtering process.

Examples

```
## Not run:  
dataPrep_raw <- UseRaw_afterFilter(dataPrep, dataFilt)  
  
## End(Not run)
```

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