

Package ‘BiocOncoTK’

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Title Bioconductor components for general cancer genomics

Description Provide a central interface to various tools for genome-scale analysis of cancer studies.

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Depends R (>= 3.6.0), methods, utils

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<code>add_sym</code>	<i>add symbols in rowData to a SummarizedExperiment that has Entrez IDs for rownames</i>
----------------------	--

Description

add symbols in rowData to a SummarizedExperiment that has Entrez IDs for rownames

Usage

```
add_sym(x)
```

Arguments

x SummarizedExperiment instance

Note

Will fail if 'symbol' is a column of rowData(x)

Examples

```
if (interactive()) {  
  bq = pancan_BQ()  
  rnse = try(buildPancanSE(bq, assay="RNASeqv2"))  
  if (inherits(rnse, "try-error")) stop("probably need CGC_BILLING set in environment or with pancan_BQ")  
  add_sym(rnse)  
}
```

annotTabs	<i>table names in Annotated pancancer data release</i>
-----------	--

Description

table names in Annotated pancancer data release

Usage

annotTabs

Format

character vector

Source

pancancer-atlas in BigQuery

Examples

```
BiocOncoTK::annotTabs
```

bindMSI	<i>bind MSI data to a SummarizedExperiment</i>
---------	--

Description

bind MSI data to a SummarizedExperiment

Usage

```
bindMSI(se, useDing = TRUE, onlyHL = TRUE)
```

Arguments

se	SummarizedExperiment instance
useDing	logical(1) if TRUE, use MSIsensor outputs from Ding et al. Cell 2018, otherwise use firehose labelings msi-h,msi-l
onlyHL	logical(1) if TRUE, retain only msi-h, msi-l records; ignored if useDing is TRUE

Value

SummarizedExperiment instance with expanded colData, samples limited to those with microsatellite instability values. The additional variable is called 'msiTest' and is numerical if useDing is TRUE and is character (msi-h,l,s) otherwise.

Note

This function adds the column `msiTest` to `colData(se)`. The contents of the column are given by `fireMSI`. Samples in `se` that do not correspond to a row of `fireMSI` are dropped. If there is already a column named `msiTest` in `colData(se)`, it is replaced and samples are filtered as described, and a message is given. If none of the samples in `se` have rows in `fireMSI`, an error is thrown. *OF NOTE:* The MSI sensor data from Ding's cell paper (see `help(dingMSI)` for URL) provides the participant barcode. The participant barcode is a substring of the sample barcode. Be sure to filter the input `SummarizedExperiment` to include only tumor samples, using the `substr(colnames(se),14,15)` (values "10"..."14" correspond to normal, non-tumor samples.) Additionally, `bindMSI` will only work if the `colnames` of the (filtered) `SummarizedExperiment` have been truncated to the participant barcode, that is, the first 12 characters of the sample barcode.

Examples

```
bindMSI
```

<code>bipg_tests</code>	<i>configure a bipartite graph relating tumor type to gene, using graphNEL</i>
-------------------------	--

Description

configure a bipartite graph relating tumor type to gene, using graphNEL

Usage

```
bipg_tests(
  statab,
  genes_adverse = NA,
  genes_favorable = NA,
  gpar_cex = 0.65,
  gpar_lwd = 0
)
```

Arguments

<code>statab</code>	a data.frame with columns 'tumor', 'gene', and 'tstat'
<code>genes_adverse</code>	a vector of genes whose increased expression is regarded as adverse
<code>genes_favorable</code>	a vector of genes whose increased expression is regarded as favorable
<code>gpar_cex</code>	tune size of graph labels
<code>gpar_lwd</code>	tune appearance of node boundaries

Value

a graphNEL instance (graph package)

Examples

```
bigg_tests(k23sig)
```

buildPancanSE	<i>helper for SummarizedExperiment construction from pancan</i>
---------------	---

Description

helper for SummarizedExperiment construction from pancan

Usage

```
buildPancanSE(
  bq,
  acronym = "BLCA",
  assay = "meth450k",
  sampType = "TP",
  subjectIDName = "ParticipantBarcode",
  seTransform = force,
  bindMethRowranges = TRUE,
  featIDMap = featIDMapper()
)
```

Arguments

bq	instance of BigQueryConnection for pancancer-atlas.Annotated Dataset
acronym	character(1) 'cohort' label, e.g., 'BLCA'
assay	character(1) element from names(BiocOncoTK::annotTabs), e.g., 'meth450k'. If 'assay == "mc3_MAF"' an error is thrown as the mutation data are inconsistently annotated; the message produced directs the user to 'mc3toGR'.
sampType	character(1) element from BiocOncoTK::pancan_sampTypeMap\$"SampleTypeLetterCode", e.g., 'TP' for Primary solid Tumor samples, or 'TB' for peripheral blood sample from primary blood derived cancer
subjectIDName	character(1) field name for subject identifier
seTransform	a function that accepts a SummarizedExperiment and returns a SummarizedExperiment; useful for feature name remapping, defaults to force (does nothing)
bindMethRowranges	logical(1) if true and assay is meth27k
featIDMap	a named character() vector defining, for each assay type, what field should be used to label features in rownames. or meth450k, annotation from FDb.InfiniumMethylation.hg19 and EnsDb.Hsapiens.v75 is obtained for available features and bound into the rowRanges component of returned object

Value

SummarizedExperiment, with metadata on acronym, assay, and sampleType propagated; if the assay is a methylation assay and bindMethRowranges is TRUE, a RangedSummarizedExperiment is returned.

Note

Note that pancancer-atlas is distinguished from TCGA by the presence of more sample types. The default type is 'TP' for primary solid tumor. Codes and their interpretations are available in Bioconductor::pancan_sampTypeMap.

Examples

```
if (interactive() && Biobase::testBioCConnection()) {
  billco = Sys.getenv("CGC_BILLING")
  if (nchar(billco)>0) {
    bq = pancan_BQ()
    methSE_BLCA = try(buildPancanSE(bq))
    methSE_BLCA
  }
}
```

CCLE_DRUG_BROAD	<i>CCLE_DRUG_BROAD: serialization of legacy CCLE 'Drug data' from Broad Institute</i>
-----------------	---

Description

CCLE_DRUG_BROAD: serialization of legacy CCLE 'Drug data' from Broad Institute

Usage

```
CCLE_DRUG_BROAD
```

Format

S4Vectors DataFrame instance

Source

data.broadinstitute.org/ccle_legacy_data/pharmacological_profiling/CCLE_NP24.2009_Drug_data_2015.02.24.csv

Examples

```
data(CCLE_DRUG_BROAD)
requireNamespace("S4Vectors")
S4Vectors::metadata(CCLE_DRUG_BROAD) # imported using read.csv, stringsAsFactors=FALSE, coerced to DataFrame
head(CCLE_DRUG_BROAD)
```

`cell_70138`*cell_70138: a table with cell-line information from LINCS*

Description

cell_70138: a table with cell-line information from LINCS

Usage

```
cell_70138
```

Format

```
data.frame
```

Source

```
GEO GSE70138 GSE70138_Broad_LINCS_cell_info_2017-04-28.txt.gz
```

Examples

```
data(cell_70138)
```

`clueDemos`*generate lists to generate clue API queries*

Description

generate lists to generate clue API queries

Usage

```
clueDemos()
```

Value

a list of lists of strings with 'where' and substructure as appropriate

Note

These are converted to JSON (

Examples

```
clueDemos()
```

clueServiceNames	<i>Provide names of some clue.io services for which examples are available in this package.</i>
------------------	---

Description

Provide names of some clue.io services for which examples are available in this package.

Usage

```
clueServiceNames()
```

Value

a character vector of service names

Note

See <https://clue.io/api>.

Examples

```
clueServiceNames()
```

darmGBMcls	<i>Data in count_lstpm format from Darmanis 2017 (PMC 5810554) single cell RNA-seq in GBM</i>
------------	---

Description

Data in count_lstpm format from Darmanis 2017 (PMC 5810554) single cell RNA-seq in GBM

Usage

```
darmGBMcls
```

Format

SummarizedExperiment with HDF Object store back end

Note

<https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5810554/> is the main source article.

Source

http://imlspenticton.uzh.ch/robinson_lab/conquer/data-mae/GSE84465.rds

Examples

```
BiocOncoTK::darmGBMcls
```

dingMSI	<i>microsatellite instability data in TCGA, collected from Ding et al. Cell 173(2) 2018.</i>
---------	--

Description

microsatellite instability data in TCGA, collected from Ding et al. Cell 173(2) 2018.

Usage

```
dingMSI
```

Format

```
DataFrame
```

Source

<https://www.cell.com/cms/10.1016/j.cell.2018.03.033/attachment/0ac495ba-3578-41cf-8fb1-94487f554bb5mmc5.xlsx> retrieved 9/17/2018.

Examples

```
str(BiocOncoTK::dingMSI)
```

featIDMapper	<i>define assay-specific feature names in a character vector</i>
--------------	--

Description

define assay-specific feature names in a character vector

Usage

```
featIDMapper()
```

Note

We may want to use Symbol instead of Entrez when retrieving expression data. The value of this function is supplied as a default for `buildPancanSE`'s `featIDMap` parameter, and alternatives can be selected by passing similarly named vectors in `featIDMap`.

Examples

```
featIDMapper()
```

fireMSI	<i>microsatellite instability data in TCGA, collected from curatedTCGAData</i>
---------	--

Description

microsatellite instability data in TCGA, collected from curatedTCGAData

Usage

```
fireMSI
```

Format

DataFrame

Source

firehose via curatedTCGAData; see metadata(BiocOncoTK::fireMSI)

Examples

```
str(S4Vectors::metadata(BiocOncoTK::fireMSI))
```

get_plates	<i>use curatedTCGAData request to acquire plate codes for samples</i>
------------	---

Description

use curatedTCGAData request to acquire plate codes for samples

Usage

```
get_plates(
  tumcode = "BLCA",
  assay = "RNASeq2GeneNorm",
  samptypes = c("01", "02", "03", "04", "06", "09", "40")
)
```

Arguments

tumcode	a TCGA tumor code, usually 3 or for characters
assay	a curatedTCGAData assay code, run curatedTCGAData() to see a message with available options
samptypes	a character vector with codes as defined at https://gdc.cancer.gov/resources-tcga-users/tcga-code-tables/sample-type-codes

Value

a data.frame with a row for each TCGA contribution for the selected tumor type and assay type

Examples

```
if (interactive()) {
  plts_blca_rnaseq = get_plates()
  dim(plts_blca_rnaseq)
  head(plts_blca_rnaseq)
}
```

ggFeatDens

create ggplot for density of starts of a GRanges in an interval

Description

create ggplot for density of starts of a GRanges in an interval

Usage

```
ggFeatDens(
  gr,
  mcolvbl,
  chrname = "chr15",
  start = 20450000,
  end = 20730000,
  binwidth.in = 5000,
  basicfilt = function(data) dplyr::filter(data, Consequence ==
    "non_coding_transcript_exon_variant"),
  ylab.in = "feature\ndensity",
  slstyle = "UCSC"
)
```

Arguments

gr	GRanges instance of interest
mcolvbl	character(1) mcols(gr) has this variable that will be used to specify different groups for computing/colouring the density traces
chrname	character(1) chromosome/seqname
start	numeric(1) start of interval
end	numeric(1) end of interval
binwidth.in	numeric(1) for geom_freqpoly binwidth setting
basicfilt	a dplyr::filter operation, defaulting to select non-coding variants in mc3 MAF
ylab.in	character(1) label for y axis
slstyle	character(1) for GenomeInfoDb::seqlevelsStyle

Value

ggplot instance

Examples

```
ggFeatDens
```

ggFeatureSegs	<i>generate a ggplot of segments of gene-like regions</i>
---------------	---

Description

generate a ggplot of segments of gene-like regions

Usage

```
ggFeatureSegs(  
  chrname = "chr15",  
  start = 20450000,  
  end = 20730000,  
  db = EnsDb.Hsapiens.v75::EnsDb.Hsapiens.v75,  
  slstyle = "UCSC",  
  ylab.in = "ensembl\nnoncoding"  
)
```

Arguments

chrname	character(1) chromosome tag
start	numeric(1) start of interval
end	numeric(1) end of interval
db	EnsDb instance for example
slstyle	character(1) tag for seqlevelsStyle
ylab.in	character(1) for use as y axis tag

Value

ggplot instance

Note

Most annotation is turned off with `element_blank()`

Examples

```
ggFeatureSegs
```

ggMutDens	<i>make a ggplot with density traces of mutations per base pair, for 'most mutated' tumor types in a given interval</i>
-----------	---

Description

make a ggplot with density traces of mutations per base pair, for 'most mutated' tumor types in a given interval

Usage

```
ggMutDens(  
  bq,  
  basicfilt = function(data) dplyr::filter(data, Consequence ==  
    "non_coding_transcript_exon_variant"),  
  chrname = "15",  
  start = 20450000,  
  end = 20730000,  
  project_volume = 5,  
  maxnrec = 50000,  
  binwidth = 5000,  
  xlab.in = " "  
)
```

Arguments

bq	bigquery BigQueryConnection instance
basicfilt	a dplyr::filter operation, defaulting to select non-coding variants in mc3 MAF
chrname	character(1) chromosome token in NCBI seqlevels style
start	numeric(1) base coordinate to start
end	numeric(1) base coordinate to end
project_volume	numeric(1) tumor types will have different numbers of contributions; this parameter tells how many tumor types to represent, counting down from the most frequently represented
maxnrec	numeric(1) for as.data.frame
binwidth	numeric(1) passed to geom_freqpoly
xlab.in	character(1) passed to ggplot2::xlab

Value

instance of ggplot

Examples

```

if (interactive()) {
  if (!requireNamespace("ggplot2")) stop("install ggplot2 to run this function")
  bq = try(pancan_BQ())
  if (!inherits(bq, "try-error")) {
    ggMutDens(bq)
  }
}

```

icd10_c	<i>helper for interpreting ICD-10 codes</i>
---------	---

Description

helper for interpreting ICD-10 codes

Usage

```
icd10_c
```

Format

data.frame

Source

ICD-10

Examples

```
BiocOncoTK::icd10_c
```

k23sig	<i>a table of 'significant' MSIsensor-score/expression relationships in TCGA</i>
--------	--

Description

a table of 'significant' MSIsensor-score/expression relationships in TCGA

Usage

```
k23sig
```

Format

data.frame

Note

provided to demonstrate bipartite graph construction

Examples

```
head(k23sig)
```

kang_DNArepair	<i>list of 151 genes annotated as DNA repair pathway members</i>
----------------	--

Description

list of 151 genes annotated as DNA repair pathway members

Usage

```
kang_DNArepair
```

Format

named list

Note

The zipped PDF was read using `pdftools::pdf_text` and then manually organized. All gene symbols present in `curatedTCGAData.RNASeq2GeneNorm` rownames. The list elements are ATM, BER, FA.HR, MMR, NER, NHEJ, OTHER, TLS, RECQ, and XLR. These denote, respectively, ataxia-telangiectasia-mutated, base excision repair, Fanconi anemia/homologous recombination, mismatch repair, nucleotide excision repair, non-homologous end joining, other, translesion synthesis, recQ helicase pathway, and cross-link repair.

Source

<https://academic.oup.com/jnci/article/104/9/670/872781#supplementary-data>

loadPatel	<i>use BiocFileCache discipline to acquire patelGBMSC SummarizedExperiment</i>
-----------	--

Description

use BiocFileCache discipline to acquire patelGBMSC SummarizedExperiment

Usage

```
loadPatel(
  remotePath = "https://s3.us-east-2.amazonaws.com/biocfound-scrna/patelGBMSC.rds",
  cache = BiocFileCache::BiocFileCache()
)
```

Arguments

remotePath	character(1) identifying remote RDS
cache	instance of BiocFileCache, defaults to BiocFileCache::BiocFileCache()

Value

a SummarizedExperiment instance

Note

The RDS for the SummarizedExperiment is in an AWS S3 bucket. This function will check local cache for the data and will download to cache if not found. That download is a one-time operation for any given value of cache.

Examples

```
loadPatel
```

load_ccleNRAS	<i>utilities for mock data (not involving internet access for vignette)</i>
---------------	---

Description

utilities for mock data (not involving internet access for vignette)

Usage

```
load_ccleNRAS()
load_NRAS_AHR()
load_nrasdf()
```

Value

a list of DRProfSet instances

a data.frame with fields 'Cell_line_primary_name', 'RMA_normalized_expression', 'HGNC_gene_symbol'

a data.frame

Note

These functions are provided only for avoiding reliance on internet connectivity for document production.

Examples

```
load_ccleNRAS()  
dim(load_nrasdf())
```

log10p11

log10(x+p) transformation for use with scales/ggplot2

Description

log10(x+p) transformation for use with scales/ggplot2

Usage

```
log10p11(p = 1)
```

Arguments

p value of shift before taking log10

Value

an instance of custom trans() for scales package

map_tcga_ncit	<i>a manually constructed table mapping TCGA acronyms to NCIT thesaurus tags</i>
---------------	--

Description

a manually constructed table mapping TCGA acronyms to NCIT thesaurus tags

Usage

```
map_tcga_ncit
```

Format

```
data.frame
```

Note

Constructed using `ontoProc::getOncotreeOnto()` result. See the vignette on Mapping TCGA tumor codes to NCIT for elaborating the mapping to aggregate tumors into NCIT organ systems.

mc3toGR	<i>create a GRanges from the MC3 mutation data</i>
---------	--

Description

create a GRanges from the MC3 mutation data

Usage

```
mc3toGR(
  bq,
  basicfilt = function(data) dplyr::filter(data, Consequence ==
    "non_coding_transcript_exon_variant"),
  maxnrec = 1e+05
)
```

Arguments

bq	bigquery BigQueryConnection instance
basicfilt	a dplyr::filter instance or NULL to convert entire MAF
maxnrec	numeric(1) used with dplyr::as.data.frame en route to GRanges

Value

a GRanges instance

Examples

```
if (interactive()) {
  con = try(pancan_BQ()) # need CGC_BILLING set
  if (!inherits(con, "try-error")) {
    aut = as.character(1:22) # some records in BQ have missing Chromosome
    chk = mc3toGR(con, basicfilt=function(data) dplyr::filter(data,
      project_short_name=="TCGA-BRCA",
      SYMBOL=="TP53", Chromosome %in% aut))
    print(chk[,1:5]) # lots of mcol fields
    table(chk$Variant_Classification)
  }
}
```

molpo_3utr

representation of 3'UTR MSI events in TCGA from Cortes-Ciriano et al. 2017

Description

representation of 3'UTR MSI events in TCGA from Cortes-Ciriano et al. 2017

Usage

molpo_3utr

Format

SummarizedExperiment

Note

Supplementary data 6 from publication noted in Source. See metadata() component of this SummarizedExperiment for more details.

Source

<https://www.nature.com/articles/ncomms15180#Sec22>

Examples

molpo_3utr

molpo_5utr	<i>representation of 5'UTR MSI events in TCGA from Cortes-Ciriano et al. 2017</i>
------------	---

Description

representation of 5'UTR MSI events in TCGA from Cortes-Ciriano et al. 2017

Usage

molpo_5utr

Format

SummarizedExperiment

Note

Supplementary data 7 from publication noted in Source. See metadata() component of this SummarizedExperiment for more details.

Source

<https://www.nature.com/articles/ncomms15180#Sec22>

Examples

molpo_5utr

molpo_CDS	<i>representation of MSI events in coding regions TCGA from Cortes-Ciriano et al. 2017</i>
-----------	--

Description

representation of MSI events in coding regions TCGA from Cortes-Ciriano et al. 2017

Usage

molpo_CDS

Format

SummarizedExperiment

Note

Supplementary data 5 from publication noted in Source. See metadata() component of this SummarizedExperiment for more details.

Source

<https://www.nature.com/articles/ncomms15180#Sec22>

Examples

```
molpo_CDS
```

```
molpo_WGS
```

representation of events detected in 708 WGS experiments TCGA from Cortes-Ciriano et al. 2017

Description

representation of events detected in 708 WGS experiments TCGA from Cortes-Ciriano et al. 2017

Usage

```
molpo_WGS
```

Format

```
SummarizedExperiment
```

Note

Supplementary data 10 from publication noted in Source. See metadata() component of this SummarizedExperiment for more details.

Source

<https://www.nature.com/articles/ncomms15180#Sec22>

Examples

```
molpo_WGS
```

MSIsensor.10k	<i>MSIsensor microsatellite instability scores for TCGA, collected from Ding et al. Cell 173(2) 2018.</i>
---------------	---

Description

MSIsensor microsatellite instability scores for TCGA, collected from Ding et al. Cell 173(2) 2018.

Usage

```
MSIsensor.10k
```

Format

```
DataFrame
```

Source

<https://www.cell.com/cms/10.1016/j.cell.2018.03.033/attachment/0ac495ba-3578-41cf-8fb1-94487f554bb5/mmc5.xlsx> retrieved 9/17/2018.

Examples

```
str(BiocOncoTK::dingMSI)
```

multiviz	<i>visualize aspects of MSIsensor/expression relationships</i>
----------	--

Description

visualize aspects of MSIsensor/expression relationships

Usage

```
multiviz(  
  tum = "MESO",  
  gene = "TYMS",  
  intrans = log10p11(p = 1),  
  inmeth = "auto",  
  topmsi = Inf,  
  indata,  
  nvar = 6  
)
```

Arguments

tum	a TCGA tumor code
gene	a gene symbol used in the indata data.frame
intrans	an instance of the trans() transformation method of scales package
inmeth	a valid setting for method parameter for geom_smooth
topmsi	maximum numeric value for x-axis when plotting against MSI value
indata	a data.frame instance with values for acronym, gene, msival
nvar	numeric() number of variables to show in biplot

oncoPrintISB	<i>interactive interface to ComplexHeatmap oncoPrint with inputs from ISB Cancer Genomics Cloud BigQuery back end</i>
--------------	---

Description

interactive interface to ComplexHeatmap oncoPrint with inputs from ISB Cancer Genomics Cloud BigQuery back end

Usage

```
oncoPrintISB(bq)
```

Arguments

bq	an instance of BigQueryConnection-class authenticated for ISB Cancer Genomics Cloud access
----	--

Value

only used for side effect of running shiny app

Note

This function will start a shiny app and will generate queries to Google BigQuery tables representing TCGA.

Examples

```
if (interactive()) {
  bcode = Sys.getenv("CGC_BILLING")
  if (nchar(bcode)>0) {
    con <- DBI::dbConnect(bigquery::bigquery(), project = "isb-cgc",
      dataset = "tcga_201607_beta", billing = bcode)
    oncoPrintISB(con)
  }
}
```

pancan.clin.varnames *pancan.clin.varnames: a data.frame with a list of variable names for clinical patient data*

Description

pancan.clin.varnames: a data.frame with a list of variable names for clinical patient data

Usage

```
pancan.clin.varnames
```

Format

data.frame

Source

pancancer-atlas in BigQuery

Examples

```
BiocOncoTK::pancan.clin.varnames[1:5,]
```

pancan_app *provide a shiny app to 'glimpse' structure and content of pancan atlas*

Description

provide a shiny app to 'glimpse' structure and content of pancan atlas

Usage

```
pancan_app(dataset = "Annotated", nrecs = 5)
```

Arguments

dataset	character(1) name of a BigQuery dataset in the pancan-atlas project
nrecs	numeric(1) number of records to request (limited through the n= parameter to as.data.table)

Value

currently only as a side effect of starting app

Examples

```
if (interactive()) pancan_app()
```

pancan_BQ *provide bigquery connection to pancancer Annotated datasets*

Description

provide bigquery connection to pancancer Annotated datasets

Usage

```
pancan_BQ(dataset = "Annotated", billing = Sys.getenv("CGC_BILLING"), ...)
```

Arguments

dataset	character(1) dataset name
billing	character(1) Google cloud platform billing code; authentication will be attempted when using the resulting connection
...	passed to dbConnect , for example, quiet=TRUE

Value

BigQueryConnection instance

Examples

```
pancan_BQ
```

pancan_clinicalTabVarnames
give an interface to tablenames

Description

give an interface to tablenames

Usage

```
pancan_clinicalTabVarnames()
```

Value

interactive datatable from DT

Examples

```
if (interactive()) pancan_clinicalTabVarnames()
```

pancan_longname	<i>utility to help find long table names</i>
-----------------	--

Description

utility to help find long table names

Usage

```
pancan_longname(guess, ...)
```

Arguments

guess	a regexp to match the table of interest
...	passed to agrep

Value

character vector of matches

Note

Note that `ignore.case=TRUE` is set in the function.

Examples

```
pancan_longname("rnaseq")
```

pancan_sampTypeMap	<i>helper for interpreting pancan-atlas sample type codes</i>
--------------------	---

Description

helper for interpreting pancan-atlas sample type codes

Usage

```
pancan_sampTypeMap
```

Format

data.frame

Note

The sample type codes are not straightforward to interpret. Primary solid tumor is denoted "TP", and metastatic samples are denoted "TM". This data frame pairs code and natural language terms.

Source

ISB BigQuery pancan-atlas project

Examples

```
BiocOncoTK::pancan_sampTypeMap
```

pancan_tabulate	<i>tabulate a variable in a table</i>
-----------------	---------------------------------------

Description

tabulate a variable in a table

Usage

```
pancan_tabulate(dataset = "Annotated", tblname, vblname)
```

Arguments

dataset	character(1) dataset name
tblname	character(1) table name in dataset
vblname	character(1) field name in table

Value

instance of tbl_dbi, constituting summarise result

Examples

```
if (interactive()) pancan_tabulate(tblname=
  "clinical_PANCAN_patient_with_followup", vblname="icd_10")
```

patient_to_tumor_code	<i>data.frame mapping from TCGA patient_barcode to TCGA tumor code</i>
-----------------------	--

Description

data.frame mapping from TCGA patient_barcode to TCGA tumor code

Usage

```
patient_to_tumor_code
```

Format

data.frame

Note

Used IDs recorded in MSISensor.10k; one is unmatched at TCGA portal metadata() component of this SummarizedExperiment for more details.

Source

<https://portal.gdc.cancer.gov/exploration?uploadCaseTab=matched>

Examples

```
head(patient_to_tumor_code)
```

pertClasses	<i>enumerate perturbation classes</i>
-------------	---------------------------------------

Description

enumerate perturbation classes

Usage

```
pertClasses(key = Sys.getenv("CLUE_KEY"))
```

Arguments

key character(1) API key provided by clue.io

Value

a character vector

Examples

```
if (nchar(Sys.getenv("CLUE_KEY"))>0) {  
  pc = pertClasses()  
  head(vapply(pc, "[[", character(1), 1))  
}
```

pert_70138

pert_70138: a table with perturbagen information from LINCS

Description

pert_70138: a table with perturbagen information from LINCS

Usage

```
pert_70138
```

Format

```
data.frame
```

Source

GEO GSE70138 GSE70138_Broad_LINCS_pert_info.txt.gz

Examples

```
data(pert_70138)
```

query_clue

run the api.clue.io API to acquire information on LINCS experiments

Description

run the api.clue.io API to acquire information on LINCS experiments

Usage

```
query_clue(
  service = "profiles",
  filter = list(where = (list(pert_iname = "sirolimus", cell_id = "MCF7", assay =
    "L1000"))),
  key = Sys.getenv("CLUE_KEY")
)
```

Arguments

service	a character(1) service name
filter	a list to be converted to JSON for submission as a GET request
key	character(1) API key provided by clue.io

Value

API return value processed by fromJSON

Examples

```
if (nchar(Sys.getenv("CLUE_KEY"))>0) {
  demos = clueDemos()
  nd = length(demos)
  chk = lapply(seq_len(nd), function(x) query_clue( service=names(demos)[x],
    filter=demos[[x]] ) )
  names(chk) = names(demos)
  sapply(chk,length)
}
```

reexports

Objects exported from other packages

Description

These objects are imported from other packages. Follow the links below to see their documentation.

dplyr [filter](#), [select](#), [tbl](#)

magrittr [%>%](#)

replaceRownames

map rownames of an SE to another vocabulary

Description

map rownames of an SE to another vocabulary

Usage

```
replaceRownames(se, sourceVocab = "ENTREZID", targetVocab = "SYMBOL")
```

Arguments

se	SummarizedExperiment instance
sourceVocab	character(1) must be a keytype of org.Hs.eg.db, defaults to 'ENTREZID'
targetVocab	character(1) must be a column of org.Hs.eg.db

small_msi	<i>filtered MSI data for demonstrating exploratory app</i>
-----------	--

Description

filtered MSI data for demonstrating exploratory app

Usage

```
small_msi
```

Format

DataFrame

Source

MSI values from dingMSI, expression from curatedTCGAData for three genes, two tumors

Examples

```
head(BiocOncoTK::small_msi)
```

TcgaMutCounts	<i>obtain data frame with counts of mutation per gene symbol for selected tumor type</i>
---------------	--

Description

obtain data frame with counts of mutation per gene symbol for selected tumor type

Usage

```
TcgaMutCounts(tumor, limit = NULL, db = "isb-cgc:tcga_201607_beta", project)
```

Arguments

tumor	character(1) defaults to 'BRCA'
limit	numeric(1) defaults to NULL, appended as limit to number of records returned if non-null
db	character(1) BigQuery database name
project	character(1) project code

Value

table as returned by bigrquery::bq_project_query

Note

This function returns overall mutation count, and many individuals have multiple mutations recorded per gene.

Examples

```
if (interactive()) {
  requireNamespace("bigrquery")
  tt = TcgaMutCounts("BRCA", project="cgc-05-0009") # substitute your project name
  head(tt)
} # need authentication
```

TcgaNIndWithAnyMut	<i>Give count of individuals with a mutation recorded for selected tumor</i>
--------------------	--

Description

Give count of individuals with a mutation recorded for selected tumor

Usage

```
TcgaNIndWithAnyMut(
  tumor = "BRCA",
  limit = NULL,
  db = "isb-cgc:tcga_201607_beta",
  project
)
```

Arguments

tumor	character(1) defaults to 'BRCA'
limit	numeric(1) defaults to NULL, appended as limit to number of records returned if non-null
db	character(1) BigQuery database name
project	character(1) project code

Value

numeric(1)

Examples

```
if (interactive()) TcgaNIndWithAnyMut(project="cgc-05-0009")
```

tumNorSet	<i>create list with SEs for tumor and normal for a tumor/assay pairing</i>
-----------	--

Description

create list with SEs for tumor and normal for a tumor/assay pairing

Usage

```
tumNorSet(  
  bq,  
  code = "PRAD",  
  assayDataTableName = pancan_longname("rnaseq"),  
  assayValueFieldName = "normalized_count",  
  assayFeatureName = "Entrez"  
)
```

Arguments

bq	a BigQuery connection
code	character(1) a TCGA tumor code, defaults to "PRAD" for prostate tumor
assayDataTableName	character(1) name of table in BigQuery
assayValueFieldName	character(1) field from which assay quantifications are retrieved
assayFeatureName	character(1) field from which assay feature names are retrieved

Examples

```
if (interactive()) {  
  bqcon = try(pancan_BQ())  
  if (!inherits(bqcon, "try-error")) {  
    tn = tumNorSet(bqcon)  
    tn  
  }  
}
```

viz_msi_raw	<i>small app to survey MSI sensor against expression</i>
-------------	--

Description

small app to survey MSI sensor against expression

Usage

```
viz_msi_raw(df, inmeth = MASS::rlm, nvar = 3)
```

Arguments

df	a data.frame instance
inmeth	a method for geom_smooth
nvar	number of variables to show in biplot

Note

Use ask=FALSE if running example.

Examples

```
if (interactive()) viz_msi_raw(BiocOncoTK::small_msi, nvar=3)
```

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