

Package ‘knowYourCG’

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Type Package

Title Functional analysis of DNA methylome datasets

Version 1.7.14

Description KnowYourCG (KYCG) is a supervised learning framework designed for the functional analysis of DNA methylation data. Unlike existing tools that focus on genes or genomic intervals, KnowYourCG directly targets CpG dinucleotides, featuring automated supervised screenings of diverse biological and technical influences, including sequence motifs, transcription factor binding, histone modifications, replication timing, cell-type-specific methylation, and trait-epigenome associations. KnowYourCG addresses the challenges of data sparsity in various methylation datasets, including low-pass Nanopore sequencing, single-cell DNA methylomes, 5-hydroxymethylation profiles, spatial DNA methylation maps, and array-based datasets for epigenome-wide association studies and epigenetic clocks (<[doi:10.1126/sciadv.adw3027](https://doi.org/10.1126/sciadv.adw3027)>).

Depends R (>= 4.4.0)

URL <https://github.com/zhou-lab/knowYourCG>

BugReports <https://github.com/zhou-lab/knowYourCG/issues>

License AGPL-3

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Config/testthat/edition 3**git_url** <https://git.bioconductor.org/packages/knowYourCG>**git_branch** devel**git_last_commit** 8df14d1**git_last_commit_date** 2026-01-22**Repository** Bioconductor 3.23**Date/Publication** 2026-02-01**Author** Wanding Zhou [aut, fnd] (ORCID:[<https://orcid.org/0000-0001-9126-1932>](https://orcid.org/0000-0001-9126-1932)),

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`aggregateTestEnrichments`*Aggregate test enrichment results*

Description

Aggregate test enrichment results

Usage

```
aggregateTestEnrichments(result_list, column = "estimate", return_df = FALSE)
```

Arguments

<code>result_list</code>	a list of results from <code>testEnrichment</code>
<code>column</code>	the column name to aggregate (Default: <code>estimate</code>)
<code>return_df</code>	whether to return a merged data frame

Value

a matrix for all results

Examples

```
## pick some big TFBS-overlapping CpG groups
kycgDataCache(data_titles=
c("KYCG.MM285.TFBSconsensus.20220116", "KYCG.MM285.chromHMM.20210210"))

sesameData::sesameDataCache(data_titles=
c("probeIDSignature", "MM285.address"))

cg_lists <- getDBs("MM285.TFBS")
queries <- cg_lists[(sapply(cg_lists, length) > 40000)]
result_list <- lapply(queries, testEnrichment, "MM285.chromHMM")
mtx <- aggregateTestEnrichments(result_list)
```

`annoProbes`*Annotate Probe IDs using KYCG databases*

Description

see `sesameData_annoProbes` if you'd like to annotate by genomic coordinates (in `GRanges`)

Usage

```
annoProbes(
  probeIDs,
  databases,
  db_names = NULL,
  platform = NULL,
  sep = ",",
  indicator = FALSE,
  silent = FALSE
)
```

Arguments

probeIDs	probe IDs in a character vector
databases	character or actual database (i.e. list of probe IDs)
db_names	specific database (default to all databases)
platform	EPIC, MM285 etc. will infer from probe IDs if not given
sep	delimiter used in paste
indicator	return the indicator matrix instead of a concatenated annotation (in the case of have multiple annotations)
silent	suppress message

Value

named annotation vector, or indicator matrix

Examples

```
kycgDataCache(data_titles="KYCG.MM285.designGroup.20210210")
probes <- names(sesameData::sesameData_getManifestGRanges("MM285"))
anno <- annoProbes(probeIDs=probes, "designGroup", silent = TRUE)
```

bedToCg

Convert BED CpG set to YAME .cg format

Description

Converts a BED file listing CpGs into a YAME-compressed .cg file using the YAME reference coordinate and command-line tools.

Usage

```
bedToCg(bed_file, ref_cr, out_file, sort_bed = TRUE, verbose = FALSE)
```

Arguments

bed_file	Character string. Path to BED file listing CpGs.
ref_cr	Character string. Path to YAME reference coordinate file (.cr).
out_file	Character string. Path to output .cg file.
sort_bed	Logical. Whether to sort the BED file with bedtools before intersecting. (Default: TRUE)
verbose	Logical. Whether to print the shell commands. (Default: FALSE)

Value

Invisibly returns the output file path.

Examples

```
# Download YAME reference coordinate file for mm10
ref_cr <- tempfile(fileext = ".cr")
download.file(
  "https://zenodo.org/records/18176270/files/mm10chr16_f7_cpg.cr",
  destfile = ref_cr, quiet = TRUE, mode = "wb"
)

# Create a BED file with CpG positions
bed_file <- tempfile(fileext = ".bed")
bed_data <- c(
  "chr16\t3003648\t3003650",
  "chr16\t3003766\t3003768",
  "chr16\t3004036\t3004038",
  "chr16\t3004052\t3004054",
  "chr16\t3004275\t3004277",
  "chr16\t3004470\t3004472",
  "chr16\t3004545\t3004547",
  "chr16\t3004633\t3004635",
  "chr16\t3004652\t3004654",
  "chr16\t3004739\t3004741",
  "chr16\t3004841\t3004843",
  "chr16\t3004959\t3004961",
  "chr16\t3005065\t3005067",
  "chr16\t3005444\t3005446",
  "chr16\t3005522\t3005524")

writeLines(bed_data, bed_file)

# Convert BED to YAME .cg format (requires bedtools and yame)
out_file <- tempfile(fileext = ".cg")

bedToCg(bed_file, ref_cr, out_file, verbose = TRUE)

# Clean up
unlink(c(bed_file, ref_cr, out_file))
```

buildGeneDBs	<i>build gene-probe association database</i>
--------------	--

Description

build gene-probe association database

Usage

```
buildGeneDBs(
  probeIDs = NULL,
  platform = NULL,
  genome = NULL,
  max_distance = 10000,
  silent = FALSE
)
```

Arguments

probeIDs	the query probe list. If NULL, use all the probes on the platform
platform	HM450, EPIC, MM285, Mammal40, will infer from query if not given
genome	hg38, mm10, ..., will infer if not given.
max_distance	probe-gene distance for association
silent	suppress messages

Value

gene databases

Examples

```
sesameData::sesameDataCache(data_titles=
  c("EPIC.address", "genomeInfo.hg38", "probeIDSignature"))
query <- c("cg04707299", "cg13380562", "cg00480749")
dbs <- buildGeneDBs(query, platform = "EPIC")
testEnrichment(query, dbs, platform = "EPIC")
```

dbStats	<i>dbStats aggregates methylation of a given betas matrix over specified database set features</i>
---------	--

Description

dbStats aggregates methylation of a given betas matrix over specified database set features

Usage

```
dbStats(  
  betas,  
  databases,  
  fun = mean,  
  na.rm = TRUE,  
  n_min = NULL,  
  f_min = 0.1,  
  long = FALSE  
)
```

Arguments

betas	matrix of beta values where probes are on the rows and samples are on the columns
databases	List of vectors corresponding to probe locations for which the features will be extracted
fun	aggregation function, default to mean
na.rm	whether to remove NA
n_min	min number of non-NA for aggregation function to apply, overrides f_min
f_min	min fraction of non-NA for aggregation function to apply
long	produce long-form result

Value

matrix with samples on the rows and database set on the columns

Examples

```
library(SummarizedExperiment)  
sesameData::sesameDataCache(data_titles=  
c("MM285.467.SE.tissue20Kprobes", "KYCG.MM285.probeType.20210630"))  
se <- sesameData::sesameDataGet("MM285.467.SE.tissue20Kprobes")  
head(dbStats(assay(se), "MM285.probeType")[,1:3])  
sesameData::sesameDataGet_resetEnv()
```

df_master	<i>Master data frame for all object to cache</i>
-----------	--

Description

This is an internal object which will be updated on every new release library(ExperimentHub) eh <- query(ExperimentHub(localHub=FALSE), "knowYourCG") eh <- query(ExperimentHub(localHub=FALSE), "sesameData") # older data data.frame(name=eh\$title, eh=names(eh))

Format

A data.frame with columns:

name Character. Resource name/title.

eh Character. ExperimentHub record ID.

Details

Cache location is default to /Users/zhouw3/Library/Caches/org.R-project.R/R/ExperimentHub/

Value

master sheet of knowYourCG objects

getDBs	<i>Get databases by full or partial names of the database group(s)</i>
--------	--

Description

Get databases by full or partial names of the database group(s)

Usage

```
getDBs(
  group_nms,
  db_names = NULL,
  platform = NULL,
  summary = FALSE,
  allow_multi = FALSE,
  type = "all",
  silent = FALSE
)
```


Arguments

group_nms	database group names
db_names	name of the database, fetch only the given databases
platform	EPIC, HM450, MM285, ... If given, will restrict to that platform.
summary	return a summary of database instead of db itself
allow_multi	allow multiple groups to be returned for each query
type	numerical, categorical, default: all
silent	no messages

Value

a list of databases, return NULL if no database is found. Each element includes ‘group’ and ‘db-name’ attributes.

Examples

```
kycgDataCache(data_titles=
c("KYCG.MM285.chromHMM.20210210", "KYCG.MM285.probeType.20210630"))
dbs <- getDBs("MM285.chromHMM")
dbs <- getDBs(c("MM285.chromHMM", "MM285.probeType"))
```

kycgDataCache	<i>Cache KnowYourCG data</i>
---------------	------------------------------

Description

Cache KnowYourCG data

Usage

```
kycgDataCache(data_titles = NULL)
```

Arguments

data_titles	data to cache, if not given will cache all
-------------	--

Value

TRUE

Examples

```
kycgDataCache("KYCG.HM27.Mask.20220123")
## to cache all data: kycgDataCache()
```

kycgDataGet	<i>Get KnowYourCG data</i>
Description	
Get KnowYourCG data	
Usage	
kycgDataGet(title, verbose = FALSE)	
Arguments	
title	title of the data
verbose	whether to output ExperimentHub message
Value	
data object	
Examples	
kycgDataCache("KYCG.MSA.CGI.20220904") EPIC.1.SigDF <- kycgDataGet('KYCG.MSA.CGI.20220904')	
KYCG_plotBar	<i>Bar plot to show most enriched CG groups from testEnrichment</i>

Description

The input data frame should have an "estimate" and a "FDR" columns.

Usage

KYCG_plotBar(df, y = "-log10(FDR)", n = 20, order_by = "FDR", label = FALSE)

Arguments

- df KYCG result data frame
- y the column to be plotted on y-axis
- n number of CG groups to plot
- order_by the column by which CG groups are ordered
- label whether to label significant bars

Details

Top CG groups are determined by estimate (descending order).

Value

grid plot object

Examples

```
KYCG_plotBar(data.frame(
  estimate=runif(10,0,10), FDR=runif(10,0,1), nD=10,
  overlap=as.integer(runif(10,0,30)), group="g", dbname=seq_len(10)))
```

KYCG_plotDot

Dot plot to show most enriched CG groups from testEnrichment

Description

The input data frame should have an "estimate" and a "FDR" columns.

Usage

```
KYCG_plotDot(
  df,
  y = "-log10(FDR)",
  n = 20,
  order_by = "FDR",
  title = "Enriched Knowledgebases",
  label_by = "dbname",
  size_by = "overlap",
  color_by = "estimate",
  short_label = FALSE
)
```

Arguments

df	KYCG result data frame
y	the column to be plotted on y-axis
n	number of CG groups to plot
order_by	the column by which CG groups are ordered
title	plot title
label_by	the column for label
size_by	the column by which CG group size plot
color_by	the column by which CG groups are colored
short_label	omit group in label

Details

Top CG groups are determined by estimate (descending order).

Value

grid plot object (by ggplot)

Examples

```
KYCG_plotDot(data.frame(
  estimate=runif(10,0,10), FDR=runif(10,0,1), nD=runif(10,10,20),
  overlap=as.integer(runif(10,0,30)), group="g", dbname=seq_len(10)))
```

KYCG_plotEnrichAll	<i>plot enrichment test result</i>
--------------------	------------------------------------

Description

plot enrichment test result

Usage

```
KYCG_plotEnrichAll(
  df,
  fdr_max = 25,
  n_label = 15,
  min_estimate = 0,
  short_label = TRUE
)
```

Arguments

df	test enrichment result data frame
fdr_max	maximum fdr for capping
n_label	number of database to label
min_estimate	minimum estimate
short_label	use short label

Value

grid object

Examples

```
query <- getDBs("MM285.designGroup")["PGCMeth"]
res <- testEnrichment(query, platform="MM285")
KYCG_plotEnrichAll(res)
```

KYCG_plotLollipop	<i>creates a lollipop plot of log(estimate) given data with fields estimate.</i>
-------------------	--

Description

creates a lollipop plot of log(estimate) given data with fields estimate.

Usage

```
KYCG_plotLollipop(df, label_column = "dbname", n = 20)
```

Arguments

df	DataFrame where each row is a database name with its estimate.
label_column	column in df to be used as the label (default: dbname)
n	Integer representing the number of top enrichments to report. Optional. (Default: 10)

Value

ggplot lollipop plot

Examples

```
KYCG_plotLollipop(data.frame(  
  estimate=runif(10,0,10), FDR=runif(10,0,1), nD=runif(10,10,20),  
  overlap=as.integer(runif(10,0,30)), group="g",  
  dbname=as.character(seq_len(10))))
```

KYCG_plotManhattan	<i>KYCG_plotManhattan makes a manhattan plot to summarize EWAS results</i>
--------------------	--

Description

KYCG_plotManhattan makes a manhattan plot to summarize EWAS results

Usage

```
KYCG_plotManhattan(
  vals,
  platform = NULL,
  genome = NULL,
  title = NULL,
  rasterize = FALSE,
  rasterize_thres = 3,
  label_min = 100,
  col = c("wheat1", "sienna3"),
  ylabel = "Value"
)
```

Arguments

vals	named vector of values (P,Q etc), vector name is Probe ID.
platform	String corresponding to the type of platform to use for retrieving GRanges coordinates of probes. Either MM285, EPIC, HM450, or HM27. If it is not provided, it will be inferred from the query set probeIDs (Default: NA).
genome	hg38, mm10, ..., will infer if not given. For additional mapping, download the GRanges object from http://zwdzwd.github.io/InfiniumAnnotation and provide the following argument ..., genome = sesameAnno_buildManifestGRanges("downloaded_file"),... to this function.
title	title for plot
rasterize	if true use ggrastr to rasterize non-significant data.
rasterize_thres	the threshold of rasterize
label_min	Threshold above which data points will be labelled with Probe ID
col	color
ylabel	y-axis label

Value

a ggplot object

Examples

```
library(sesameData)

## Create example with simulated -log10(p-values)
## Mix of non-significant (low values) and significant (high values)
probes <- names(sesameData_getManifestGRanges("HM450"))
set.seed(123)
vals <- setNames(
  c(runif(990, 0, 3),      # Non-significant probes
    runif(10, 5, 25)),    # Significant probes
  sample(probes, 1000))
```

```

)

KYCG_plotManhattan(vals,
  platform = "HM450",
  title = "Example Manhattan Plot",
  ylabel = "-log10(P-value)",
  label_min = 20)

```

KYCG_plotMeta

Plot meta gene or other meta genomic features

Description

Plot meta gene or other meta genomic features

Usage

```
KYCG_plotMeta(betas, platform = NULL)
```

Arguments

betas	a named numeric vector or a matrix (row: probes; column: samples)
platform	if not given and x is a SigDF, will be inferred the meta features

Value

a grid plot object

Examples

```

library(sesameData)
library(sesame)
sdf <- sesameDataGet("EPIC.1.SigDF")
KYCG_plotMeta(getBetas(sdf))

```

KYCG_plotMetaEnrichment

Plot meta gene or other meta genomic features

Description

Plot meta gene or other meta genomic features

Usage

```
KYCG_plotMetaEnrichment(result_list)
```

Arguments

result_list one or a list of testEnrichment

Value

a grid plot object

Examples

```
cg_lists <- getDBs("MM285.TFBS")
queries <- cg_lists[(sapply(cg_lists, length) > 40000)]
result_list <- lapply(queries, testEnrichment,
  "MM285.metagene", silent=TRUE, platform="MM285")

KYCG_plotMetaEnrichment(result_list)
```

KYCG_plotPointRange	<i>Plot point range for a list of enrichment testing results against the same set of databases</i>
---------------------	--

Description

Plot point range for a list of enrichment testing results against the same set of databases

Usage

```
KYCG_plotPointRange(result_list)
```

Arguments

result_list a list of testEnrichment resultsx

Value

grid plot object

Examples

```
## pick some big TFBS-overlapping CpG groups
cg_lists <- getDBs("MM285.TFBS")
queries <- cg_lists[(sapply(cg_lists, length) > 40000)]
result_list <- lapply(queries, testEnrichment,
  "MM285.chromHMM", platform="MM285")
KYCG_plotPointRange(result_list)
```

`KYCG_plotSetEnrichment`*Plot Set Enrichment*

Description

Plot Set Enrichment

Usage

```
KYCG_plotSetEnrichment(result, n_sample = 1000, n_presence = 200)
```

Arguments

<code>result</code>	result object as returned from an element of the list of <code>testEnrichmentSEA(..., prepPlot=TRUE)</code>
<code>n_sample</code>	number of CpGs to sample
<code>n_presence</code>	number of overlap to sample for the plot

Value

grid object for plot

Examples

```
query <- getDBs("KYCG.MM285.designGroup")[[ "VMR" ]]  
db <- getDBs("MM285.seqContextN", "distToTSS")  
res <- testEnrichmentSEA(query, db, prepPlot = TRUE)  
KYCG_plotSetEnrichment(res[[1]])
```

<code>KYCG_plotVolcano</code>	<i>creates a volcano plot of $-\log_2(p.value)$ and $\log(estimate)$ given data with fields estimate and p.value.</i>
-------------------------------	---

Description

creates a volcano plot of $-\log_2(p.value)$ and $\log(estimate)$ given data with fields estimate and p.value.

Usage

```
KYCG_plotVolcano(df, label_by = "dbname", alpha = 0.05)
```

Arguments

df	DataFrame where each field is a database name with two fields for the estimate and p.value.
label_by	column in df to be used as the label (default: dbname)
alpha	Float representing the cut-off alpha value for the plot. Optional. (Default: 0.05)

Value

ggplot volcano plot

Examples

```
KYCG_plotVolcano(data.frame(
  estimate=runif(10,0,10), FDR=runif(10,0,1), nD=runif(10,10,20),
  overlap=as.integer(runif(10,0,30)), group="g", dbname=seq_len(10)))
```

KYCG_plotWaterfall	<i>create a waterfall plot of log(estimate) given test enrichment</i>
--------------------	---

Description

create a waterfall plot of log(estimate) given test enrichment

Usage

```
KYCG_plotWaterfall(
  df,
  order_by = "Log2(OR)",
  size_by = "-log10(FDR)",
  label_by = "dbname",
  n_label = 10
)
```

Arguments

df	data frame where each row is a database with test enrichment result
order_by	the column by which CG groups are ordered
size_by	the column by which CG group size plot
label_by	column in df to be used as the label (default: dbname)
n_label	number of datapoints to label

Value

grid

Examples

```
library(SummarizedExperiment)
library(sesameData)
df <- rowData(sesameDataGet('MM285.tissueSignature'))
query <- df$Probe_ID[df$branch == "fetal_brain" & df$type == "Hypo"]
results <- testEnrichment(query, "TFBS", platform="MM285")
KYCG_plotWaterfall(results)
```

linkProbesToProximalGenes

find genes in genomic proximity to given Infinium probes

Description

This is a convenient function that uses `sesameData_getGenomeInfo()` to retrieve stored gene models.

Usage

```
linkProbesToProximalGenes(probeIDs, platform = NULL, genome = NULL)
```

Arguments

probeIDs	character vector of probe IDs
platform	HM450, EPIC, EPICv2, MM285, MSA, ..., will infer from probe ID if not given
genome	hg38, hg19, mm10, this is usually inferred from platform.

Details

For finer control, such as taking only genes by their promoters, please use `sesameData_getTxnGRanges` followed by `sesameData_annoProbes()`. See code of this convenient function for details.

Value

a data frame annotate gene list linked to each given probes

Examples

```
library(SummarizedExperiment)
probes = rowData(
  sesameData::sesameDataGet('MM285.tissueSignature'))$Probe_ID[1:10]
linkProbesToProximalGenes(probes, platform = "MM285")
```

listDBGroups	<i>List database group names</i>
--------------	----------------------------------

Description

List database group names

Usage

```
listDBGroups(filter = NULL, path = NULL)
```

Arguments

filter	keywords for filtering
path	file path to downloaded knowledgebase sets

Value

a list of db group names

Examples

```
head(listDBGroups("chromHMM"))  
## or listDBGroups(path = "~/Downloads")
```

loadDBs	<i>Load knowledgebase databases from TSV files</i>
---------	--

Description

This used to be an exported function. Now it's internal. Use RDS download directly.

Usage

```
loadDBs(in_paths)
```

Arguments

in_paths	Character vector of file paths, URLs to .tsv or .tsv.gz files, or a single directory path containing such files. If a directory is provided, all files in that directory will be loaded. URLs (starting with http:// or https://) are loaded directly.
----------	--

Details

This function loads knowledgebase sets from tab-delimited (.tsv or .tsv.gz) files downloaded from Zenodo or other sources. The TSV files should contain two columns: "Probe_ID" and "Knowledgebase". The function splits the data by knowledgebase name and returns a list of database vectors.

The input TSV file(s) must have a header row and contain at least two columns:

- Probe_ID - Probe identifiers (e.g., cg12345678)
- Knowledgebase - Knowledgebase/database name

Value

A list of database vectors. Each element contains Probe_IDs with attributes:

- group - The database group name (derived from filename)
- dbname - The knowledgebase name (from the Knowledgebase column)

testEnrichment

Test for enrichment of query in knowledgebase sets

Description

Test for enrichment of query in knowledgebase sets

Usage

```
testEnrichment(
  query,
  databases = NULL,
  universe = NULL,
  alternative = "greater",
  include_genes = FALSE,
  platform = NULL,
  silent = FALSE,
  mtc_by_group = TRUE,
  mtc_method = "fdr"
)
```

Arguments

query	For array input, a vector of probes of interest (e.g., significant differential methylated probes). For sequencing data input, the file name for YAME-compressed CG sets.
databases	List of vectors corresponding to the database sets of interest with associated meta data as an attribute to each element. If NULL, all available databases for the platform are used. (Default: NULL)

universe	Vector of probes in the universe set containing all probes to be considered in the test. If NULL, will be inferred from the provided platform. (Default: NULL)
alternative	Test alternative: "two.sided", "greater", or "less". (Default: "greater")
include_genes	Include gene link enrichment testing. (Default: FALSE)
platform	String corresponding to the type of platform to use: MM285, EPIC, HM450, or HM27. If NULL, will be inferred from query set probe IDs. (Default: NULL)
silent	Suppress output messages? (Default: FALSE)
mtc_by_group	Perform multiple testing correction within knowledgebase groups. (Default: TRUE)
mtc_method	Method for multiple test correction. (Default: "fdr")

Value

A data frame containing features corresponding to the test estimate, p-value, and type of test, ordered by significance.

Examples

```
library(SummarizedExperiment)
library(sesameData)
library(knowYourCG)
kycgDataCache(data_titles = "KYCG.MM285.chromHMM.20210210")
sesameDataCache("MM285.tissueSignature")
df <- rowData(sesameDataGet("MM285.tissueSignature"))
probes <- df$Probe_ID[df$branch == "B_cell"]
res <- testEnrichment(probes, "chromHMM", platform = "MM285")

# Define temporary directory and file URLs
temp_dir <- tempdir()
knowledgebase <- file.path(temp_dir, "ChromHMM.20220414.cm")
query <- file.path(temp_dir, "mm10_f3_10cells.cg")

# URLs for the knowledgebase and query files
knowledgebase_url <- paste0(
  "https://zenodo.org/records/18175656/files/",
  "ChromHMM.20220414.cm"
)
query_url <- paste0(
  "https://zenodo.org/records/18176004/files/",
  "mm10_f3_10cells.cg"
)

# Download the files
download.file(knowledgebase_url, destfile = knowledgebase)
download.file(query_url, destfile = query)

# Confirm file download
list.files(temp_dir)
res <- testEnrichment(query, knowledgebase)
```

testEnrichment2	<i>Test enrichment from YAME-compressed CG sets</i>
-----------------	---

Description

Tests for enrichment of genomic regions in YAME-compressed CG sets using Fisher's exact test. This function is optimized for sequencing data and uses compiled C code for efficient processing.

Usage

```
testEnrichment2(
  query_fn,
  knowledge_fn,
  universe_fn = NULL,
  alternative = "greater",
  min_overlap = 1,
  verbose = FALSE
)
```

Arguments

query_fn	Character string specifying the file path to the query CG set file (YAME-compressed format).
knowledge_fn	Character string or vector specifying the file path(s) to the knowledgebase file(s) (YAME-compressed format). Can be a single file or multiple files.
universe_fn	Optional character string specifying the file path to the universe CG set file. If NULL, universe will be inferred from the knowledgebase. (Default: NULL)
alternative	Character string specifying the alternative hypothesis: "greater" (enrichment), "less" (depletion), or "two.sided". (Default: "greater")
min_overlap	Minimum number of overlapping CGs required for a test to be included in results. (Default: 1)
verbose	Logical indicating whether to print progress messages. (Default: FALSE)

Value

A tibble containing enrichment test results with the following columns:

Mask Name/identifier of the knowledgebase mask

N_mask Number of CGs in the mask

N_query Number of CGs in the query

N_overlap Number of overlapping CGs

N_univ Total number of CGs in universe

estimate Log2 odds ratio

p.value P-value from Fisher's exact test
log10.p.value Log10-transformed p-value
test Type of test performed
Additional effect size metrics Jaccard, MCC, etc.

Examples

```
if (.Platform$OS.type != "windows") {
  kfn <- system.file("extdata", "chromhmm.cm", package = "knowYourCG")
  qfn <- system.file("extdata", "onecell.cg", package = "knowYourCG")
  res <- testEnrichment2(qfn, kfn)
  head(res)
}
```

testEnrichmentSEA	<i>uses the GSEA-like test to estimate the association of a categorical variable against a continuous variable.</i>
-------------------	---

Description

estimate represent enrichment score and negative estimate indicate a test for depletion

Usage

```
testEnrichmentSEA(
  query,
  databases,
  platform = NULL,
  silent = FALSE,
  precise = FALSE,
  prepPlot = FALSE
)
```

Arguments

query	query, if numerical, expect categorical database, if categorical expect numerical database
databases	database, numerical or categorical, but needs to be different from query
platform	EPIC, MM285, ..., infer if not given
silent	suppress message (default: FALSE)
precise	whether to compute precise p-value (up to numerical limit) of interest.
prepPlot	return the raw enrichment scores and presence vectors for plotting

Value

A DataFrame with the estimate/statistic, p-value, and name of test for the given results.

Examples

```
sesameData::sesameDataCache(data_titles=
c("KYCG.MM285.designGroup.20210210", "KYCG.MM285.seqContextN.20210630",
"probeIDSignature"))
query <- getDBs("KYCG.MM285.designGroup")[["TSS"]]
res <- testEnrichmentSEA(query, "MM285.seqContextN")
```

testEnrichmentSpearman

testEnrichmentSpearman uses the Spearman statistical test to estimate the association between two continuous variables.

Description

testEnrichmentSpearman uses the Spearman statistical test to estimate the association between two continuous variables.

Usage

```
testEnrichmentSpearman(num_query, num_db)
```

Arguments

num_query	named numeric vector of probes of interest where names are probe IDs (e.g significant probes)
num_db	List of vectors corresponding to the database set of interest with associated meta data as an attribute to each element.

Value

A DataFrame with the estimate/statistic, p-value, and name of test for the given results.

testProbeProximity	<i>testProbeProximity tests if a query set of probes share closer genomic proximity than if randomly distributed</i>
--------------------	--

Description

testProbeProximity tests if a query set of probes share closer genomic proximity than if randomly distributed

Usage

```
testProbeProximity(
  probeIDs,
  gr = NULL,
  platform = NULL,
  iterations = 100,
  bin_size = 1500
)
```

Arguments

probeIDs	Vector of probes of interest (e.g., significant probes)
gr	GRanges to draw samples and compute genomic distances
platform	String corresponding to the type of platform to use. Either MM285, EPIC, HM450, or HM27. If it is not provided, it will be inferred from the query set probeIDs (Default: NA).
iterations	Number of random samples to generate null distribution (Default: 100).
bin_size	the poisson interval size for computing neighboring hits

Value

list containing a dataframe for the poisson statistics and a data frame for the probes in close proximity

Examples

```
sesameData::sesameDataCache(data_titles=
c("MM285.tissueSignature", "MM285.address", "probeIDSignature"))
library(SummarizedExperiment)
df <- rowData(sesameData::sesameDataGet("MM285.tissueSignature"))
probes <- df$Probe_ID[df$branch == "B_cell"]
res <- testProbeProximity(probeIDs=probes, platform="MM285")
sesameData::sesameDataGet_resetEnv()
```

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