Package 'CaDrA'

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Description Performs both stepwise and backward heuristic search for candidate (epi)genetic drivers based on a binary multi-omics dataset. CaDrA's main objective is to identify features which, together, are significantly skewed or enriched pertaining to a given vector of continuous scores (e.g. sample-specific scores representing a phenotypic readout of interest, such as protein expression, pathway activity, etc.), based on the union occurence (i.e. logical OR) of the events. **Depends** R (>= 4.4.0) LazyData false License GPL-3 + file LICENSE URL https://github.com/montilab/CaDrA/ **Encoding UTF-8** RoxygenNote 7.3.2 Imports doParallel, ggplot2, gplots, graphics, grid, gtable, knnmi, MASS, methods, misc3d, plyr, ppcor, R.cache, reshape2, stats, SummarizedExperiment Suggests BiocManager, devtools, knitr, pheatmap, rmarkdown, testthat (>= 3.1.6)Config/testthat/edition 3 biocViews Microarray, RNASeq, GeneExpression, Software, FeatureExtraction VignetteBuilder knitr BugReports https://github.com/montilab/CaDrA/issues git_url https://git.bioconductor.org/packages/CaDrA git_branch devel git_last_commit 3f6af6c git_last_commit_date 2024-10-29

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Contents

BRCA_GISTIC_MUT_SIG	2
CaDrA	3
calc_rowscore	6
candidate_search	10
CCLE_MUT_SCNA	13
CTNBB1_reporter	13
generate_permutations	14
meta_plot	15
permutation_plot	16
perm_res	16
prefilter_data	17
sim_FS	18
sim_Scores	19
TAZYAP_BRCA_ACTIVITY	20
topn_best	20
topn_list	21
topn_plot	22
	24

 ${\tt BRCA_GISTIC_MUT_SIG} \qquad \textit{Genomic Data from TCGA BRCA MUT} + \textit{GISTIC}$

Description

A SummarizedExperiment object consists of 16,873 genomic features across 951 samples.

Usage

Index

data(BRCA_GISTIC_MUT_SIG)

Format

An object of class SummarizedExperiment from SummarizedExperiment package containing an assay of 16,873 rows (features) and 951 columns (samples) see SummarizedExperiment for more details.

CaDrA 3

Value

a SummarizedExperiment object

References

Kartha VK, Kern JG, Sebastiani P, Zhang L, Varelas X, Monti S (2019) CaDrA: A computational framework for performing candidate driver analyses using binary genomic features. (Frontiers in Genetics)

CaDrA

CaDrA Search

Description

Perform permutation-based testings on a sample of permuted input scores using candidate_search as the main iterative function for each run.

Usage

```
CaDrA(
 FS,
  input_score,
 method = c("ks_pval", "ks_score", "wilcox_pval", "wilcox_score", "revealer", "knnmi",
    "correlation", "custom"),
 method_alternative = c("less", "greater", "two.sided"),
 cmethod = c("spearman", "pearson"),
  custom_function = NULL,
  custom_parameters = NULL,
 weights = NULL,
  search_start = NULL,
  top_N = 1,
  search_method = c("both", "forward"),
 max\_size = 7,
  n_{perm} = 1000,
  perm_alternative = c("one.sided", "two.sided"),
  obs_best_score = NULL,
  smooth = TRUE,
  plot = FALSE,
 ncores = 1,
  cache = FALSE,
 cache_path = NULL,
  verbose = FALSE
)
```

4 CaDrA

Arguments

FS a matrix of binary features or a SummarizedExperiment class object from Sum-

marizedExperiment package where rows represent features of interest (e.g. genes, transcripts, exons, etc...) and columns represent the samples. The assay of FS contains binary (1/0) values indicating the presence/absence of omics features.

input_score a vector of continuous scores representing a phenotypic readout of interest such

as protein expression, pathway activity, etc.

NOTE: input_score object must have names or labels that match the column

names of FS object.

method a character string specifies a scoring method that is used in the search. There

are 6 options: ("ks_pval" or ks_score or "wilcox_pval" or wilcox_score or "revealer" (conditional mutual information from REVEALER) or "knnmi" (K-Nearest Neighbor Mutual Information Estimator from knnmi) or "correlation"

or "custom" (a user-defined scoring method)). Default is ks_pval.

method_alternative

a character string specifies an alternative hypothesis testing ("two.sided" or "greater" or "less"). Default is less for left-skewed significance testing.

NOTE: This argument only applies to ks_pval and wilcox_pval method

cmethod correlation method to use - spearman or pearson. Default is "spearman".

NOTE: This argument only applies to correlation method

custom_function

If method is "custom", specifies a user-defined function here. Default is NULL.

NOTE: custom_function must take FS and input_score as its input arguments and its final result must return a vector of row-wise scores where its labels or names match the row names of FS object.

custom_parameters

If method is "custom", specifies a list of additional arguments (excluding FS and input_score) to be passed to custom_function. For example: custom_parameters

= list(alternative = "less"). Default is NULL.

weights If method is ks_score or ks_pval, specifying a vector of weights will perform

a weighted-KS testing. Default is NULL.

NOTE: weights must have names or labels that match the labels of input_score.

search_start a vector of character strings (separated by commas) specifies feature names in

the FS object to start the search with. If search_start is provided, then top_N

parameter will be ignored and vice versa. Default is NULL.

top_N an integer specifies the number of features to start the search over. By default, it

starts with the feature that has the highest best score (top $_N = 1$).

NOTE: If top_N is provided, then search_start parameter will be ignored and

vice versa. If $top_N > 10$, it may result in a longer search time.

search_method a character string specifies an algorithm to filter out the best candidates ("forward"

or "both"). Default is both (i.e., backward and forward).

max_size an integer specifies a maximum size that a meta-feature can extend to do for a

given search. Default is 7.

n_perm an integer specifies the number of permutations to perform. Default is 1000.

CaDrA 5

perm_alternative

an alternative hypothesis type for calculating permutation-based p-value. Op-

tions: one.sided, two.sided. Default is one.sided.

obs_best_score a numeric value corresponding to the best observed score. This value is used

to compare against the n_perm calculated best scores. Default is NULL. If set to

NULL, we will compute the observed best score based on the given parameters.

smooth a logical value indicates whether or not to add a smoothing factor of 1 to the

calculation of permutation-based p-value. This option is used to avoid a returned

p-value of 0. Default is TRUE.

plot a logical value indicates whether or not to plot the empirical null distribution of

the permuted best scores. Default is FALSE.

ncores an integer specifies the number of cores to perform parallelization for permutation-

based testing. Default is 1.

cache a logical value determines whether or not to cache the permuted best scores. This

helps to save time for future loading instead of re-computing the permutation-

based testing every time. Default is FALSE.

cache_path a path to cache permuted best scores. Default is NULL. If NULL, the cache path

is set to system home directory (e.g. \$HOME/.Rcache) for future loading.

verbose a logical value indicates whether or not to print the diagnostic messages. Default

is FALSE.

Value

a list of 4 objects: key, perm_best_scores, obs_best_score, perm_pval

-key: a list of parameters that was used to cache the results of the permutation-based testing. This is useful as the permuted best scores can be recycled to save time for future loading.

-perm_best_scores: a vector of permuted best scores obtained by performing candidate_search over n_perm iterations of permuted input scores.

-obs_best_score: a user-provided best score or an observed best score obtained by performing candidate_search on a given dataset and input parameters. This value is later used to compare against the permuted best scores (perm_best_scores).

perm_pval: a permutation-based p-value obtained by calculating sum(perm_best_scores > obs_best_score)/n_perm

NOTE: If smooth = TRUE, a smoothing factor of 1 will be added to the calculation of perm_pval.

e.g. $(sum(perm_best_scores > obs_best_score) + 1) / (n_perm + c)$

This is just to not return a p-value of 0

Examples

```
# Load pre-computed feature set
data(sim_FS)
```

Load pre-computed input-score
data(sim_Scores)

Set seed for permutation

```
# Define additional parameters and start the search function
cadra_result <- CaDrA(
   FS = sim_FS, input_score = sim_Scores, method = "ks_pval",
   weights = NULL, method_alternative = "less", top_N = 1,
   search_start = NULL, search_method = "both", max_size = 7,
   n_perm = 10, perm_alternative = "one.sided", plot = FALSE,
   smooth = TRUE, obs_best_score = NULL,
   ncores = 1, cache = FALSE, cache_path = NULL
)</pre>
```

calc_rowscore

Calculate row-wise scores of a given binary feature set based on a given scoring method

Description

Calculate row-wise scores of a given binary feature set based on a given scoring method

Usage

```
calc_rowscore(
   FS,
   input_score,
   meta_feature = NULL,
   method = c("ks_pval", "ks_score", "wilcox_pval", "wilcox_score", "revealer", "knnmi",
        "correlation", "custom"),
   method_alternative = c("less", "greater", "two.sided"),
   cmethod = c("spearman", "pearson"),
   custom_function = NULL,
   custom_parameters = NULL,
   weights = NULL,
   do_check = TRUE,
   verbose = FALSE,
   ...
)
```

Arguments

FS

a matrix of binary features or a SummarizedExperiment class object from SummarizedExperiment package where rows represent features of interest (e.g. genes, transcripts, exons, etc...) and columns represent the samples. The assay of FS contains binary (1/0) values indicating the presence/absence of omics features.

input_score

a vector of continuous scores representing a phenotypic readout of interest such as protein expression, pathway activity, etc.

NOTE: input_score object must have names or labels that match the column names of FS object.

meta_feature a vector of one or more features representing known causes of activation or

features associated with a response of interest (e.g. input_score). Default is

NULL.

method a character string specifies a scoring method that is used in the search. There

are 6 options: ("ks_pval" or ks_score or "wilcox_pval" or wilcox_score or "revealer" (conditional mutual information from REVEALER) or "knnmi" (k-Nearest Neighbor Mutual Information Estimator from knnmi) or "correlation" (based on simple correlation - pearson or spearman) or "custom" (a user-defined

scoring method)). Default is ks_pval.

method_alternative

a character string specifies an alternative hypothesis testing ("two.sided" or "greater" or "less"). Default is less for left-skewed significance testing.

NOTE: This argument only applies to ks_pval and wilcox_pval method

cmethod correlation method to use - spearman or pearson. Default is "spearman" #'

NOTE: This argument only applies to correlation method only

custom_function

if method is "custom", specifies a user-defined function here. Default is NULL.

NOTE: custom_function must take FS and input_score as its input arguments, and its final result must return a vector of row-wise scores where its labels or names matched the row names of FS object.

custom_parameters

if method is "custom", specifies a list of additional arguments (excluding FS and input_score) to be passed to custom_function. For example: custom_parameters

= list(alternative = "less"). Default is NULL.

weights If method is ks_score or ks_pval, specifying a vector of weights will perform

a weighted-KS testing. Default is NULL.

NOTE: weights must have names or labels that match the names or labels of

input_score.

do_check a logical value indicates whether or not to validate if the given parameters (FS

and input_score) are valid inputs. Default is TRUE.

verbose a logical value indicates whether or not to print the diagnostic messages. Default

is FALSE.

... additional parameters to be passed to custom_function

Value

return a vector of row-wise positive scores where it is ordered from most significant to least significant (e.g. from highest to lowest values) and its labels or names must match the row names of FS object

```
colnames(mat) <- 1:10</pre>
row.names(mat) <- c("TP_1", "TP_2", "TP_3")
# Create a vector of observed input scores
set.seed(42)
input_score = rnorm(n = ncol(mat))
names(input_score) <- colnames(mat)</pre>
# Run the ks method
ks_rowscore_result <- calc_rowscore(
 FS = mat,
 input_score = input_score,
 meta_feature = NULL,
 method = "ks_pval",
 method_alternative = "less",
 weights = NULL
)
# Run the wilcoxon method
wilcox_rowscore_result <- calc_rowscore(</pre>
 FS = mat,
 input_score = input_score,
 meta_feature = NULL,
 method = "wilcox_pval";
 method_alternative = "less"
# Run the revealer method
revealer_rowscore_result <- calc_rowscore(</pre>
 FS = mat,
 input_score = input_score,
 meta_feature = NULL,
 method = "revealer"
# Run the revealer method
knnmi_rowscore_result <- calc_rowscore(</pre>
 FS = mat,
 input_score = input_score,
 meta_feature = NULL,
 method = "knnmi"
)
# Run the correlation method
corr_result <- calc_rowscore(</pre>
  FS = mat,
   input_score = input_score,
  meta_feature = NULL,
  method = "correlation",
  cmethod = "spearman"
)
```

```
# A customized function using ks-test function
customized_ks_rowscore <- function(FS, input_score, meta_feature=NULL, alternative="less"){</pre>
 # Check if meta_feature is provided
 if(!is.null(meta_feature)){
    # Getting the position of the known meta features
    locs <- match(meta_feature, row.names(FS))</pre>
    # Taking the union across the known meta features
    if(length(locs) > 1) {
      meta\_vector \leftarrow as.numeric(ifelse(colSums(FS[locs,]) == 0, 0, 1))
    }else{
      meta_vector <- as.numeric(FS[locs,])</pre>
    # Remove the meta features from the binary feature matrix
    # and taking logical OR btw the remaining features with the meta vector
    FS <- base::sweep(FS[-locs, , drop=FALSE], 2, meta_vector, `|`)*1
    # Check if there are any features that are all 1s generated from
    # taking the union between the matrix
    # We cannot compute statistics for such features and thus they need
    # to be filtered out
    if(any(rowSums(FS) == ncol(FS))){
      warning("Features with all 1s generated from taking the matrix union ",
              "will be removed before progressing...\n")
      FS <- FS[rowSums(FS) != ncol(FS), , drop=FALSE]
   }
 }
 # KS is a ranked-based method
 # So we need to sort input_score from highest to lowest values
 input_score <- sort(input_score, decreasing=TRUE)</pre>
 # Re-order the matrix based on the order of input_score
 FS <- FS[, names(input_score), drop=FALSE]
 # Compute the scores using the KS method
 ks <- apply(FS, 1, function(r){</pre>
   x = input_score[which(r==1)];
   y = input_score[which(r==0)];
   res <- ks.test(x, y, alternative=alternative)</pre>
    return(c(res$statistic, res$p.value))
 })
 # Obtain score statistics
 stat <- ks[1,]
 # Obtain p-values and change values of 0 to the machine lowest value
 # to avoid taking -\log(0)
 pval <- ks[2,]
 pval[which(pval == 0)] <- .Machine$double.xmin</pre>
```

10 candidate_search

```
# Compute the -log(pval)
# Make sure scores has names that match the row names of FS object
scores <- -log(pval)
names(scores) <- rownames(FS)

return(scores)

# Search for best features using a custom-defined function
custom_rowscore_result <- calc_rowscore(
FS = mat,
    input_score = input_score,
    meta_feature = NULL,
    method = "custom",
    custom_function = customized_ks_rowscore,
    custom_parameters = NULL
)</pre>
```

candidate_search

Candidate Search

Description

Performs heuristic search on a set of binary features to determine whether there are features whose union is more skewed (enriched at the extremes) than either features alone. This is the main functionality of the CaDrA package.

Usage

```
candidate_search(
 FS,
  input_score,
 method = c("ks_pval", "ks_score", "wilcox_pval", "wilcox_score", "revealer", "knnmi",
    "correlation", "custom"),
 method_alternative = c("less", "greater", "two.sided"),
  cmethod = c("spearman", "pearson"),
  custom_function = NULL,
  custom_parameters = NULL,
  weights = NULL,
  search_start = NULL,
  top_N = 1,
  search_method = c("both", "forward"),
 max\_size = 7,
  best_score_only = FALSE,
  do_plot = FALSE,
  verbose = FALSE
)
```

candidate_search 11

Arguments

FS a matrix of binary features or a SummarizedExperiment class object from Sum-

marizedExperiment package where rows represent features of interest (e.g. genes, transcripts, exons, etc...) and columns represent the samples. The assay of FS contains binary (1/0) values indicating the presence/absence of omics features.

input_score a vector of continuous scores representing a phenotypic readout of interest such

as protein expression, pathway activity, etc.

NOTE: input_score object must have names or labels that match the column

names of FS object.

method a character string specifies a scoring method that is used in the search. There

are 7 options: ("ks_pval" or ks_score or "wilcox_pval" or wilcox_score or "revealer" (conditional mutual information from REVEALER) or "knnmi" (K-Nearest Neighbor Mutual Information Estimator from knnmi) or "correlation" (based on simple correlation - pearson or spearman) or "custom" (a user-defined

scoring method)). Default is ks_pval.

method_alternative

a character string specifies an alternative hypothesis testing ("two.sided" or "greater" or "less"). Default is less for left-skewed significance testing.

NOTE: This argument only applies to ks_pval and wilcox_pval method

cmethod correlation method to use - spearman or pearson. Default is "spearman".

NOTE: This argument only applies to correlation method only

custom_function

if method is "custom", specifies a user-defined function here. Default is NULL.

NOTE: custom_function must take FS and input_score as its input arguments and its final result must return a vector of row-wise scores where its labels or

names match the row names of FS object.

custom_parameters

if method is "custom", specifies a list of additional arguments (excluding FS and input_score) to be passed to the custom_function. For example: cus-

tom_parameters = list(alternative = "less"). Default is NULL.

weights if method is ks_score or ks_pval, specifying a vector of weights will perform

a weighted-KS testing. Default is NULL.

NOTE: weights must have names or labels that match the labels of input_score.

search_start a vector of character strings (separated by commas) specifies feature names in

the FS object to start the search with. If search_start is provided, then top_N

parameter will be ignored and vice versa. Default is NULL.

top_N an integer specifies the number of features to start the search over. By default, it

starts with the feature that has the highest best score (top $_N = 1$).

NOTE: If top_N is provided, then search_start parameter will be ignored and

vice versa. If $top_N > 10$, it may result in a longer search time.

search_method a character string specifies an algorithm to filter out the best features ("forward"

or "both"). Default is both (i.e. backward and forward).

max_size an integer specifies a maximum size that a meta-feature can extend to do for a

given search. Default is 7.

12 candidate_search

best_score_only

a logical value indicates whether or not to return the best score corresponding to

each top N searches only. Default is FALSE.

do_plot a logical value indicates whether or not to plot the overlapping features of the

resulting meta-feature matrix.

NOTE: plot can only be produced if the resulting meta-feature matrix contains more than 1 feature (e.g. length(search_start) > 1 or top_N > 1). Default is

FALSE.

verbose a logical value indicates whether or not to print the diagnostic messages. Default

is FALSE.

Details

 $NOTE: The \ legacy \ function \ topn_eval \ is \ equivalent \ to \ the \ recommended \ candidate_search \ function$

Value

If best_score_only = TRUE, the heuristic search will return the best feature whose its union meta-feature matrix has the highest score among the top_N feature searches. If best_score_only = FALSE, a list of objects pertaining to top_N feature searches will be returned. For each top_N feature search, the candidate search will contain 7 objects: (1) its best meta-feature matrix (feature_set), (2) its observed input scores (input_score), (3) its corresponding best score pertaining to the union meta-feature matrix (score), (4) names of the best meta-features (best_features), (5) rank of the best meta-features in term of their best scores (best_indices), (6) marginal scores of the best meta-features (marginal_best_scores), (7) cumulative scores of the best meta-features (cumulative_best_scores).

```
# Load pre-computed feature set
data(sim_FS)

# Load pre-computed input scores
data(sim_Scores)

# Define additional parameters and run the function
candidate_search_result <- candidate_search(
   FS = sim_FS, input_score = sim_Scores,
   method = "ks_pval", method_alternative = "less", weights = NULL,
   search_start = NULL, top_N = 3, search_method = "both",
   max_size = 7, best_score_only = FALSE
)</pre>
```

CCLE_MUT_SCNA 13

CCLE_MUT_SCNA

Genomic Data from CCLE MUT + SCNA

Description

A SummarizedExperiment object consists of 17,724 genomic features across 82 samples.

Usage

```
data(CCLE_MUT_SCNA)
```

Format

An object of class SummarizedExperiment from SummarizedExperiment package containing a matrix of 17,724 rows (features) and 82 columns (samples). See SummarizedExperiment for more details.

Value

a SummarizedExperiment object

References

Kim, J., Botvinnik, O., Abudayyeh, O. et al. Characterizing genomic alterations in cancer by complementary functional associations. Nat Biotechnol 34, 539–546 (2016). https://doi.org/10.1038/nbt.3527

CTNBB1_reporter

Transcriptional Activity of Beta-Catenin in Cancers

Description

A vector of continuous scores represents the activation of B-catenin across multiple cancer cell lines

Usage

```
data(CTNBB1_reporter)
```

Format

Consists of a vector of continuous scores of B-catenin activity across 82 cancer cell lines. The mutation and copy number associated with this sample cohorts can be found in CCLE_MUT_SCNA dataset.

Value

a vector of continuous scores

References

Kim, J., Botvinnik, O., Abudayyeh, O. et al. Characterizing genomic alterations in cancer by complementary functional associations. Nat Biotechnol 34, 539–546 (2016). https://doi.org/10.1038/nbt.3527

generate_permutations Random permutation matrix generator

Description

Produces a random permutation score matrix given a vector of sample-specific scores representing a phenotypic readout of interest such as protein expression, pathway activity, etc.

Usage

```
generate_permutations(input_score, n_perm)
```

Arguments

input_score a vector of continuous scores of a molecular phenotype of interest such as pro-

tein expression, pathway activity, etc. NOTE: The input_score object must

have names or labels to track samples by.

n_perm a number of permutations to generate. This determines the number of rows in

the permutation matrix.

Value

a matrix of values where each row contains scores of a single permuted input_score.

```
# Load pre-simulated scores
data(sim_Scores)

# Set seed for permutation
set.seed(123)

# Define number of permutations
n_perm = 1000

# Generate permuted scores
perm_matrix <- generate_permutations(
   input_score = sim_Scores,
   n_perm = n_perm
)</pre>
```

meta_plot 15

meta_plot

Candidate Drivers Search Plot

Description

By utilizing the top N results obtained from candidate_search, we can find the best meta-feature among the top N searches using topn_best. meta_plot is then used to produce graphics including a tile plot for the top meta-features that associated with a molecular phenotype of interest (e.g. input_score), the KS enrichment plot of the meta-features, and lastly, a density diagram of the distribution of the observed input scores sorted from largest to smallest at the top.

Usage

```
meta_plot(topn_best_list, input_score_label = NULL, plot_title = NULL)
```

Arguments

topn_best_list a list of objects returned from candidate_search corresponding to the search of top N features given by top_N value. The topn_best_list contains the best meta-feature matrix, its corresponding best score, its observed input scores, rank of the best features based on their scores, marginal best scores, and cumulative best scores.

input_score_label

a label that references to the input_score variable that was used to compute the top N best features. Default is NULL.

plot_title a title t

a title to the plot. Default is NULL.

Value

3 plots stacked on top of each other: 1. a density diagram of observed input scores sorted from highest to lowest 2. a tile plot of the top features within the meta-feature set 3. a KS enrichment plot of the meta-feature set (this correspond to the logical OR of the features)

```
# Load pre-computed Top-N list generated for sim_FS dataset
data(topn_list)

# With the results obtained from top-N evaluation,
# We can find the combination of features that gives the best score in
# top N searches
topn_best_meta <- topn_best(topn_list = topn_list)

# Now we can plot this set of best meta-feature
meta_plot(topn_best_list = topn_best_meta)</pre>
```

perm_res

permutation_plot

Permutation Best Scores Plot

Description

Plot the Empirical Null Distribution of Permutation Best Scores returned from CaDrA function

Usage

```
permutation_plot(perm_res)
```

Arguments

perm_res

a list of objects returned from CaDrA function. The returning object contains a list of key parameters that are used to run the permutation-based testing, a vector of permuted best scores for a given n_perm, an observed best score, and a computed permutation p-value.

Value

a density plot

Examples

```
# Load pre-computed permutation results
data(perm_res)

# Plot the permutation results
permutation_plot(perm_res)
```

perm_res

Pre-computed permutation results for simulated data (sim_FS)

Description

The permutation result returned from CaDrA using pre-simulated dataset (FS = sim_FS), pre-simulated input scores (input_score = sim_Scores), top_N = 7, method = "ks_pval", alternative = "less", search_method = "both", max_size = 10, obs_best_score = NULL and n_perm = 1000 as inputs to the function.

Usage

```
data(perm_res)
```

prefilter_data 17

Format

A list of objects returned from CaDrA function. The resulting object contains a list of key parameters that was used to run the permutation-based testing, a vector of permuted best scores for a given n_perm, an observed best score, and a permuted p-value.

To visualize the Empirical Null Distribution of the permuted best scores over n_perm iterations, just pass the resulting list to permutation_plot.

See permutation_plot for more details.

Value

a list of objects returned from CaDrA function

References

Kartha VK, Kern JG, Sebastiani P, Zhang L, Varelas X, Monti S (2017) CaDrA: A computational framework for performing candidate driver analyses using binary genomic features. (Frontiers in Genetics)

Examples

```
# Load the pre-computed permutation results for sim_FS
data(perm_res)

# Plot the Empirical Null Distribution of the permuted best scores
# against its observed best score
permutation_plot(perm_res = perm_res)
```

prefilter_data

Pre-filter features

Description

Pre-filter a dataset prior running candidate_search to avoid testing features that are too prevalent or too sparse across samples in the dataset

Usage

```
prefilter_data(FS, max_cutoff = 0.6, min_cutoff = 0.03, verbose = FALSE)
```

Arguments

FS

a matrix of binary features or a SummarizedExperiment class object from SummarizedExperiment package where rows represent features of interest (e.g. genes, transcripts, exons, etc...) and columns represent the samples. The assay of FS contains binary (1/0) values indicating the presence/absence of 'omics' features.

18 sim_FS

max_cutoff	a numeric value between 0 and 1 describing the absolute prevalence of a feature across all samples in the FS object which the feature will be filtered out. Default is 0.6 (feature that occur in 60 percent or more of the samples will be removed)
min_cutoff	a numeric value between 0 and 1 describing the absolute prevalence of a feature across all samples in the FS object which the feature will be filtered out. Default is 0.03 (feature that occur in 3 percent or less of the samples will be removed)
verbose	a logical value indicates whether or not to print the diagnostic messages. Default is FALSE.

Value

A SummarizedExperiment object with only the filtered-in features given the filtered thresholds

Examples

```
# Load pre-computed feature set
data(sim_FS)

# Filter out features having < 3% and > 60% prevalence across all samples
# by (default)
sim_FS_filt1 <- prefilter_data(FS = sim_FS)

# Change the min cut-off to 1% prevalence, instead of the default of 3%
sim_FS_filt2 <- prefilter_data(FS = sim_FS, min_cutoff = 0.01)

# Change the max cut-off to 65% prevalence, instead of the default of 60%
sim_FS_filt3 <- prefilter_data(FS = sim_FS, max_cutoff = 0.65)</pre>
```

sim_FS

Simulated Genomic Data

Description

A simulated SummarizedExperiment object that comprises of 1000 genomic features (rows) and 100 sample profiles (columns). Each row is represented by a vector of binary values (1/0) indicating the presence/absence of the feature in the samples. This simulated data includes 10 left-skewed (i.e. True Positive or TP) and 990 uniformly-distributed (i.e. True Null or TN) features.

Usage

```
data(sim_FS)
```

Format

An object of class SummarizedExperiment from SummarizedExperiment package containing an assay of 1000 rows (features) and 100 columns (samples). Each row is represented by a vector of binary values (1/0) indicating the presence/absence of the feature in the samples.

See ?SummarizedExperiment for more details.

sim_Scores 19

Value

a SummarizedExperiment object

References

Kartha VK, Kern JG, Sebastiani P, Zhang L, Varelas X, Monti S (2019) CaDrA: A computational framework for performing candidate driver analyses using binary genomic features. (Frontiers in Genetics)

sim_Scores

Simulated Input Scores

Description

A simulated vector of continuous scores generated from rnorm(n=ncol(sim_FS), mean=0, sd=1) with set.seed(123) based on the number of samples in the simulated dataset (sim_FS)

Usage

data(sim_Scores)

Format

A vector of continuous scores randomly generated from rnorm(n=ncol(sim_FS), mean=0, sd=1) with set.seed(123) based on the number of samples in the simulated dataset (sim_FS)

Value

a vector of continuous scores

References

Kartha VK, Kern JG, Sebastiani P, Zhang L, Varelas X, Monti S (2019) CaDrA: A computational framework for performing candidate driver analyses using binary genomic features. (Frontiers in Genetics)

20 topn_best

TAZYAP_BRCA_ACTIVITY YAP/TAZ Activity in TCGA BRCA dataset

Description

A vector of continuous scores represents oncogenic YAP/TAZ activity in human breast carcinomas

Usage

```
data(TAZYAP_BRCA_ACTIVITY)
```

Format

consists of a vector of continuous scores of YAP/TAZ activity across 951 profiles. The mutation and copy number associated with this sample cohorts can be found in BRCA_GISTIC_MUT_SIG dataset.

Value

a vector of continuous scores

References

Kartha VK, Kern JG, Sebastiani P, Zhang L, Varelas X, Monti S (2019) CaDrA: A computational framework for performing candidate driver analyses using binary genomic features. (Frontiers in Genetics)

topn_best

Top 'N' Best Meta-Features

Description

Take the resulting list of meta-features returned from candidate_search over top N feature searches and fetch the meta-feature with the best score.

Usage

```
topn_best(topn_list)
```

Arguments

topn_list

A nested list of objects that are returned from candidate_search using the following parameters: FS = sim_FS, input_score = sim_Scores, top_N = 7, method = "ks_pval", alternative = "less", search_method = "both", max_size = 10, and best_score_only = FALSE.

topn_list 21

Value

A list of objects containing the best meta-feature matrix, its corresponding best score, its observed input scores, rank of best meta-features based on their scores, its marginal and cumulative best scores.

Examples

```
# Load pre-computed Top-N list generated for sim_FS dataset
data(topn_list)

# Get the best meta-features list
topn_best_meta <- topn_best(topn_list = topn_list)</pre>
```

topn_list

Top-N Results for Simulated Data (sim_FS)

Description

A list of objects returned from candidate_search using simulated dataset FS = sim_FS, input_score = sim_Scores, top_N = 7, method = "ks_pval", alternative = "less", search_method = "both", max_size = 10, and best_score_only = FALSE.

Usage

```
data(topn_list)
```

Format

A list of objects returned from candidate_search including a set of best meta-feature matrix, its corresponding best score, its observed input scores, rank of the best features based on their scores, marginal best scores, and cumulative best scores. pertaining to each top N feature searches.

See candidate_search for more information.

Details

NOTE: max_size is set to 10 as we would like to account for the presence of 10 left-skewed (i.e. true positive or TP) features in sim_FS dataset.

Value

a list of objects returned from candidate_search function

References

Kartha VK, Kern JG, Sebastiani P, Zhang L, Varelas X, Monti S (2019) CaDrA: A computational framework for performing candidate driver analyses using binary genomic features. (Frontiers in Genetics)

22 topn_plot

Examples

```
# Load pre-computed Top-N list generated for sim_FS and sim_Scores dataset
data(topn_list)

# Fetch the first meta-feature
topn_list[[1]]$feature_set

# Fetch the second meta-feature
topn_list[[2]]$feature_set

# Retrieve the meta-feature with the best score among top_N = 7 runs
topn_best_meta <- topn_best(topn_list = topn_list)

# Visualize the best meta-feature using meta_plot function
meta_plot(topn_best_list = topn_best_meta)

# Visualize overlap of meta-features across top_N = 7
# using topn_plot function
topn_plot(topn_list = topn_list)</pre>
```

topn_plot

Top 'N' Plot

Description

Generate a heatmap representation of overlapping meta-features across top N feature searches using candidate_search function

Usage

```
topn_plot(topn_list)
```

Arguments

```
topn_list
```

a list of objects obtained from candidate_search using the following parameters: FS = sim_FS, input_score = sim_Scores, top_N = 7, method = "ks_pval", alternative = "less", search_method = "both", max_size = 10, and best_score_only = FALSE.

Value

a heatmap of overlapping meta-features across top N feature searches

topn_plot 23

```
# Load pre-computed Top-N list generated for sim_FS dataset
data(topn_list)

# Get the overlapping top N plot
topn_plot(topn_list = topn_list)
```

Index

```
* datasets
    BRCA_GISTIC_MUT_SIG, 2
    CCLE_MUT_SCNA, 13
    CTNBB1_reporter, 13
    perm_res, 16
    sim_FS, 18
    sim_Scores, 19
    TAZYAP_BRCA_ACTIVITY, 20
    topn_list, 21
BRCA_GISTIC_MUT_SIG, 2
CaDrA, 3
calc_rowscore, 6
candidate_search, 10, 21
CCLE_MUT_SCNA, 13
CTNBB1_reporter, 13
generate_permutations, 14
meta_plot, 15
perm_res, 16
permutation_plot, 16
prefilter_data, 17
sim_FS, 18
sim_Scores, 19
TAZYAP_BRCA_ACTIVITY, 20
topn_best, 20
topn_list, 21
topn_plot, 22
```