Package ‘COMPASS’

May 29, 2024

Type Package

Title Combinatorial Polyfunctionality Analysis of Single Cells

Version 1.42.0

Date 2014-07-11

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Description COMPASS is a statistical framework that enables unbiased analysis of antigen-specific T-cell subsets. COMPASS uses a Bayesian hierarchical framework to model all observed cell-subsets and select the most likely to be antigen-specific while regularizing the small cell counts that often arise in multi-parameter space. The model provides a posterior probability of specificity for each cell subset and each sample, which can be used to profile a subject's immune response to external stimuli such as infection or vaccination.

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BugReports https://github.com/RGLab/COMPASS/issues

VignetteBuilder knitr

Depends R (>= 3.0.3)

LinkingTo Rcpp (>= 0.11.0)

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Imports methods, Rcpp, data.table, RColorBrewer, scales, grid, plyr, knitr, abind, clue, grDevices, utils, pdist, magrittr, reshape2, dplyr, tidyr, rlang, BiocStyle, rmarkdown, foreach, coda

Suggests flowWorkspace (>= 3.33.1), flowCore, ncdfFlow, shiny, testthat, devtools, flowWorkspaceData, ggplot2, progress

LazyLoad yes

LazyData yes

biocViews ImmunoOncology, FlowCytometry

Encoding UTF-8

RoxygenNote 7.1.1
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COMPASS-package

COMPASS (Combinatorial Polyfunctionality Analysis of Single-Cells)

Description

This package implements a model for the analysis of polyfunctionality in single-cell cytometry experiments. The model effectively identifies combinations of markers that are differentially expressed between samples of cells subjected to different stimulations.

See Also

- COMPASSContainer, for information on getting your cytometry data into a suitable format for use with COMPASS,
- COMPASS, for the main model fitting routine.

categories

Description

Returns the categories matrix in a COMPASSResult object.

Usage

categories(x, counts)

Arguments

x A COMPASSResult object.

counts Boolean; if TRUE we return the counts (degree of functionality) as well.
CellCounts

Compute Number of Cells Positive for Certain Cytokine Combinations

Description

Compute the number of cells expressing a particular combination of markers for each sample.

Usage

CellCounts(data, combinations)

Arguments

- **data**: Either a COMPASSContainer, or a list of matrices. Each matrix is of dimension \(N_i\) cells (rows) by \(K\) common markers (columns).
- **combinations**: A list of 'combinations', used to denote the subsets of interest. See the examples for usage.

See Also

Combinations

Examples

```r
set.seed(123)
## generate 10 simulated matrices of flow data
K <- 6  ## number of markers
data <- replicate(10, simplify=FALSE, {
  m <- matrix( rnorm(1E4 * K, 2000, 1000 ), ncol=K )
  m[m < 2500] <- 0
  colnames(m) <- c("IL2", "IL4", "IL6", "Mip1B", "IFNg", "TNFa")
  return(m)
})
names(data) <- sample(letters, 10)
head( data[[1]] )

## generate counts over all available combinations of markers in data
str(CellCounts(data)) ## 64 columns, as all 2^6 combinations expressed

## generate marginal counts
combos <- list(1, 2, 3, 4, 5, 6) ## marginal cell counts
cc <- CellCounts(data, combos)

## a base R way of doing the same thing
f <- function(data) {
  do.call(rbind, lapply(data, function(x) apply(x, 2, function(x) sum(x > 0))))
}
cc2 <- f(data)
```
## check that they’re identical
stopifnot(identical(unname(cc), unname(cc2)))

## We can also generate cell counts by expressing various combinations
## of markers (names) in the data.

## count cells expressing IL2 or IL4
CellCounts(data, "IL2|IL4")

## count cells expressing IL2, IL4 or IL6
CellCounts(data, "IL2|IL4|IL6")

## counts for each of IL2, IL4, IL6 (marginally)
CellCounts(data, c("IL2", "IL4", "IL6"))

## counts for cells that are IL2 positive and IL4 negative
CellCounts(data, "IL2 & !IL4")

## expressing the same intent with indices
CellCounts(data, list(c(1, -2)))

## all possible combinations
str(CellCounts(data, Combinations(6)))

## can also call on COMPASSContainers
data(COMPASS)
CellCounts(CC, "M1&M2")

---

### Combinations

**Generate Combinations**

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Given an integer n, generate all binary combinations of n elements, transformed to indices. This is primarily for use with the CellCounts function, but may be useful for users in some scenarios.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Usage</th>
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<tbody>
<tr>
<td>Combinations(n)</td>
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<table>
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<tr>
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<tr>
<td>n</td>
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<table>
<thead>
<tr>
<th>Examples</th>
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<tr>
<td>Combinations(3)</td>
</tr>
</tbody>
</table>
Description

This function fits the COMPASS model.

Usage

COMPASS(
  data,
  treatment,
  control,
  subset = NULL,
  category_filter = function(x) colSums(x > 5) > 2,
  filter_lowest_frequency = 0,
  filter_specific_markers = NULL,
  model = "discrete",
  iterations = 40000,
  replications = 8,
  keep_original_data = FALSE,
  verbose = TRUE,
  dropDegreeOne = FALSE,
  init_with_fisher = FALSE,
  run_model_or_return_data = "run_model",
  ...
)

Arguments

data An object of class COMPASSContainer.
treatment An R expression, evaluated within the metadata, that returns TRUE for those samples that should belong to the treatment group. For example, if the samples that received a positive stimulation were named "92TH023 Env" within a variable in meta called Stim, you could write Stim == "92TH023 Env". The expression should have the name of the stimulation vector on the left hand side.
control An R expression, evaluated within the metadata, that returns TRUE for those samples that should belong to the control group. See above for details.
subset An expression used to subset the data. We keep only the samples for which the expression evaluates to TRUE in the metadata.
category_filter A filter for the categories that are generated. This is a function that will be applied to the treatment counts matrix generated from the intensities. Only categories meeting the category_filter criteria will be kept.
COMPASS

filter_lowest_frequency
A number specifying how many of the least expressed markers should be removed.

filter_specific_markers
Similar to filter_lowest_frequency, but lets you explicitly exclude markers.

model
A string denoting which model to fit; currently, only the discrete model ("discrete") is available.

iterations
The number of iterations (per 'replication') to perform.

replications
The number of 'replications' to perform. In order to conserve memory, we only keep the model estimates from the last replication.

keep_original_data
Keep the original COMPASSContainer as part of the COMPASS output? If memory or disk space is an issue, you may set this to FALSE.

verbose
Boolean; if TRUE we output progress information.

dropDegreeOne
Boolean; if TRUE we drop degree one categories and merge them with the negative subset.

init_with_fisher
Boolean; initialize from fisher's exact test. Any subset and subject with lower 95% or greater than 95% ps <= pu. Otherwise initialize very subject and subset as a responder except those where ps <= pu.

run_model_or_return_data
character defaults to "run_model" otherwise set it to "return_data" in order to not fit the model just return the data set needed for modeling. Useful for extracting the boolean counts.

...
Other arguments; currently unused.

Value

A COMPASSResult is a list with the following components:

fit
A list of various fitted parameters resulting from the COMPASS model fitting procedure.

data
The data used as input to the COMPASS fitting procedure – in particular, the counts matrices generated for the selected categories, n_s and n_u, can be extracted from here.

orig
If keep_original_data was set to TRUE in the COMPASS fit, then this will be the COMPASSContainer passed in. This is primarily kept for easier running of the Shiny app.

The fit component is a list with the following components:

alpha_s
The hyperparameter shared across all subjects under the stimulated condition. It is updated through the COMPASS model fitting process.

A_alphas
The acceptance rate of alpha_s, as computed through the MCMC sampling process in COMPASS.
alpha_u The hyperparameter shared across all subjects under the unstimulated condition. It is updated through the COMPASS model fitting process.

A_alphau The acceptance rate of alpha_u, as computed through the MCMC sampling process in COMPASS.

gamma An array of dimensions I x K x T, where I denotes the number of individuals, K denotes the number of categories / subsets, and T denotes the number of iterations. Each cell in a matrix for a given iteration is either zero or one, reflecting whether individual i is responding to the stimulation for subset k.

mean_gamma A matrix of mean response rates. Each cell denotes the mean response of individual i and subset k.

A_gamma The acceptance rate for the gamma. Each element corresponds to the number of times an individual’s gamma vector was updated.

categories The category matrix, showing which categories entered the model.

model The type of model called.

posterior Posterior measures from the sample fit.

call The matched call used to generate the model fit.

The data component is a list with the following components:

n_s The counts matrix for stimulated samples.

n_u The counts matrix for unstimulated samples.

counts_s Total cell counts for stimulated samples.

counts_u Total cell counts for unstimulated samples.

categories The categories matrix used to define which categories will enter the model.

meta The metadata. Note that only individual-level metadata will be kept; sample-specific metadata is dropped.

sample_id The name of the vector in the metadata used to identify the samples.

individual_id The name of the vector in the metadata used to identify the individuals.

The orig component (included if keep_original_data is TRUE) is the COMPASSContainer object used in the model fit.

Category Filter

The category filter is used to exclude categories (combinations of markers expressed for a particular cell) that are expressed very rarely. It is applied to the treatment counts matrix, which is a N samples by K categories matrix. Those categories which are mostly unexpressed can be excluded here. For example, the default criteria,

category_filter=function(x) colSums(x > 5) > 2

indicates that we should only retain categories for which at least three samples had at least six cells expressing that particular combination of markers.

See Also

- COMPASSContainer, for constructing the data object required by COMPASS
COMPASSContainer

Examples

```r
data(COMPASS) ## loads the COMPASSContainer 'CC'
fit <- COMPASS(CC,
               category_filter=NULL,
               treatment=trt == "Treatment",
               control=trt == "Control",
               verbose=FALSE,
               iterations=100 ## set higher for a real analysis
)
```

COMPASSContainer Generate the Data Object used by COMPASS

Description

This function generates the data container suitable for use with COMPASS.

Usage

```r
COMPASSContainer(
    data,
    counts,
    meta,
    individual_id,
    sample_id,
    countFilterThreshold = 0
)
```

Arguments

data A list of matrices. Each matrix _M_i_ is made up of _N_i_ cells by _K_ markers; for example, it could be the intensity information from an intracellular cytokine experiment. Each element of the list should be named; this name denotes which sample the cell intensities were measured from.

counts A named integer vector of the cell counts (of the parent population) for each sample in `data`.

meta A `data.frame` of metadata, describing the individuals in the experiment. Each row in `meta` should correspond to a row in `data`. There should be one row for each sample; i.e., one row for each element of `data`.

individual_id The name of the vector in `meta` that denotes the individuals from which samples were drawn. In this sense an individual equates to a single subject, or person.

sample_id The name of the vector in `meta` that denotes the samples. The `sample_id` identifies a combination of a subject with visit (if any), cell subset measured (e.g. CD4), and stimulation. This vector should contain all of the names in the `data` input.

countFilterThreshold Numeric; if the number of parent cells is less than this threshold, we remove that file. Default is 0, which means filter is disabled.
Details

The names attributes for the data and counts objects passed should match.

Value

A COMPASSContainer returns a list made up of the same components as input the model, but checks and sanitizes the supplied data to ensure that it conforms to the expectations outlined above.

Examples

```r
set.seed(123)
n <- 10 ## number of samples
k <- 3 ## number of markers

## generate some sample data
sid_vec <- paste0("sid_{}, 1:n") ## sample ids; unique names used to denote samples
iid_vec <- rep_len( paste0("iid_{}, 1:(n/2) "), n ) ## individual ids

## generate n matrices of 'cell intensities'
data <- replicate(n, {
  nrow <- round(runif(1) * 1E2 + 1000)
  ncol <- k
  vals <- rexp( nrow * ncol, runif(1, 1E-5, 1E-3) )
  vals[ vals < 2000 ] <- 0
  output <- matrix(vals, nrow, ncol)
  output <- output[ apply(output, 1, sum) > 0, ]
  colnames(output) <- paste0("M", 1:k)
  return(output)
})
names(data) <- sid_vec

## make a sample metadata data.frame
meta <- data.frame(
  sid=sid_vec,
  iid=iid_vec,
  trt=rep( c("Control", "Treatment"), each=5 )
)

## generate an example total counts
## recall that cells not expressing any marker are not included
## in the 'data' matrices
counts <- sapply(data, nrow) + round( rnorm(n, 1E3, 1E2) )
counts <- setNames( as.integer(counts), names(counts) )

## insert everything into a COMPASSContainer
CC <- COMPASSContainer(data, counts, meta, "iid", "sid")
```
**COMPASSContainer-data**  Simulated COMPASSContainer

**Description**

This dataset contains simulated data for an intracellular cytokine staining experiment. In this data set, we have paired samples from five individuals, with each pair of samples being subjected to either a 'Control' condition or a 'Treatment' condition.

**Details**

Please see [COMPASSContainer](#) for more information on the components of this object.

The dataset is exported as `CC`, which is a short alias for COMPASSContainer.

---

**COMPASSContainerFromGatingSet**  
*Create a COMPASS Container from a GatingSet*

**Description**

This code expects a `GatingSet` or `GatingSetList`. It expects a regular expression for the node name (i.e. `/4+$` would match `/4+/` in a node name with the plus sign at the end of the string). Alternatively, you can supply a partial path. The user must supply the `individual_id`, which has the default value suitable for the data we commonly see. `sample_id` is the `rownames` of `pData` of `GatingSet`. Sometimes the child node names don't match the marker names exactly. This function will try to make some guesses about how to match these up. The `filter.fun` parameter is a function that does some regular expression string substitution to try and clean up the node names by removing various symbols that are often added to gates, {+/-}. The user can provide their own function to do string cleanup. Counts are extracted as well as metadata and single cell data, and these are fed into the COMPASSContainer constructor.

**Usage**

```r
COMPASSContainerFromGatingSet(
gs = NULL,
node = NULL,
filter.fun = NULL,
individual_id = "PTID",
mp = NULL,
matchmethod = c("Levenshtein", "regex"),
markers = NA,
swap = FALSE,
countFilterThreshold = 5000
)
```
Arguments

- **gs**: a `GatingSet` or `GatingSetList`.
- **node**: a regular expression to match a single node in the gating tree. If more than one node is matched, an error is thrown.
- **filter.fun**: a function that does string substitution to clean up node names, i.e., turns a 'CD4+' into a 'CD4' to try and match against the parameters slot of the `flowFrames` in `gs`.
- **individual_id**: a character identifying the subject id column in the `gs` metadata.
- **mp**: a list mapping node names to markers. This function tries to guess, but may fail. The user can override the guesswork.
- **matchmethod**: a character either 'regex' or 'Levenshtein' for matching nodes to markers.
- **markers**: a character vector of marker names to include.
- **swap**: a logical default `FALSE`. Set to `TRUE` if the marker and channel names are swapped.
- **countFilterThreshold**: numeric threshold. If the number of parent cells is less than this threshold, we remove that file. Default is 5000.

Details

There is likely not sufficient error checking.

See Also

COMPASSContainer

Examples

```r
## Not run:
## gs is a GatingSet from flowWorkspace
COMPASSContainerFromGatingSet(gs, "4+")
## End(Not run)
```

Description

This is used for setting an informative description used in the Shiny application.

Usage

COMPASSDescription(x)

COMPASSDescription(x) <- value
COMPASSfitToCountsTable

Arguments

x  A COMPASS fit.
value A set of paragraphs describing the experiment, as a character vector.

Details

Information about the COMPASS results will be auto-generated.

Description

Returns a table of counts and parent counts for each cell subset in a COMPASS fit.

Usage

COMPASSfitToCountsTable(
  x,
  idcol = NULL,
  parent = NULL,
  drop = NULL,
  stimName = NULL
)

Arguments

x   COMPASSResult
idcol   unquote variable name in the metadata for the subject id.
parent   character name of the parent population for this model fit. e.g. "CD4"
drop   numeric vector indicating the columns in the metadata to drop from the output. Usually sample-specific columns rather than subject specific columns.
stimName   the name of the stimulation
COMPASSMCMCDiagnosis  Diagnostic of a set of COMPASS Models.

Description
Diagnostic of a set of COMPASS Models.

Usage
COMPASSMCMCDiagnosis(x)

Arguments
x  a list of compass model fits of the same data with the same number of iterations, different seeds. Run some mcmc diagnostics on a series of COMPASS model fits. Assuming the input is a list of model fits for the same data with the same number of iterations and different seeds. Run Gelman’s Rhat diagnostics on the alpha_s and alpha_u hyperparameter chains, treating each model as an independent chain. Rhat should be near 1 but rarely are in practice. Very large values may be a concern. The method returns an average model, by averaging the mean_gamma matrices (equally weighted since each input has the same number of iterations). This mean model should be better then any of the individual models. It can be plotted via "plot(result$mean_model)".

COMPASSResult-accessors  COMPASSResult Accessors

Description
These functions can be used for accessing data within a COMPASSResult.

Usage
Gamma(x)

MeanGamma(x)

Arguments
x  A COMPASSResult object.
**COMPASSResult-data**  

Simulated COMPASS fit

**Description**

This dataset represents the result of fitting the COMPASS model on the accompanying dataset `CC`, as exported by `data(COMPASS)`. Please see the vignette (`vignette("COMPASS")`) for more details on how to interact with a COMPASS fit.

**Details**

The model is fit as follows, using the COMPASSContainer `CC`.

```r
CR <- COMPASS(CC, 
    treatment=trt == "Treatment", 
    control=trt == "Control", 
    iterations=1000 
)
```

The dataset is exported as `CR`, which is a short alias for `COMPASSResult`. Please see `COMPASS` for more information on the output from a COMPASS model fit.

---

**FunctionalityScore**  

Compute the Functionality Score for each subject fit in a COMPASS model

**Description**

Computes the functionality score for each observation from the gamma matrix of a COMPASS model fit. The scores are normalized according to the total number of possible subsets that could be observed \((2^M - 1)\).

**Usage**

```r
FunctionalityScore(x, n, markers = NULL)

## S3 method for class 'COMPASSResult'
FunctionalityScore(x, n, markers = NULL)

## Default S3 method:
FunctionalityScore(x, n, markers = NULL)
```
getCounts

Arguments

- `x`: An object of class `COMPASSResult`, as returned by `COMPASS`. Alternatively, a matrix of functionality scores, used under the assumption that the 'null' category has been dropped.
- `n`: The number of markers included in an experiment. It is inferred from the data when `x` is a `COMPASSResult`.
- `markers`: The set of markers for which to compute a Functionality score. By default uses all markers. Must match names returned by `markers()`.

Value

A numeric vector of functionality scores.

Note

The null category is implicitly dropped when computing the functionality score for a `COMPASS` result; this is not true for the regular matrix method.

Examples

```r
FunctionalityScore(CR)
```

---

**getCounts**

Get a data.table of counts of polyfunctional subsets

**Description**

Get a data.table of counts of polyfunctional subsets

**Usage**

```r
getCounts(object)
```

**Arguments**

- `object`: An object of class `COMPASSResult`

**Examples**

```r
getCounts(CR)
```
GetThresholdedIntensities

Extract Thresholded Intensities from a GatingSet

Description
This function extracts thresholded intensities for children of a node \texttt{node}, as specified through the \texttt{map} argument.

Usage
GetThresholdedIntensities(\texttt{gs, node, map})

Arguments
\begin{itemize}
  \item \texttt{gs} A \texttt{GatingSet} or \texttt{GatingSetList}.
  \item \texttt{node} The name, or path, of a single node in a \texttt{GatingSet} / \texttt{GatingSetList}.
  \item \texttt{map} A list, mapping node names to markers.
\end{itemize}

Details
\texttt{map} should be an \texttt{R} list, mapping node names (as specified in the gating hierarchy of the gating set) to channel names (as specified in either the desc or name columns of the parameters of the associated \texttt{flowFrames} in the \texttt{GatingSet}).

Value
A list with two components:
\begin{itemize}
  \item \texttt{data} A list of thresholded intensity measures.
  \item \texttt{counts} A named vector of total cell counts at the node \texttt{node}.
\end{itemize}

Examples
\begin{verbatim}
if (require("flowWorkspace")&require("flowCore")&require("tidyr")) {

  ## Generate an example GatingSet that could be used with COMPASS
  ## We then pull out the 'data' and 'counts' components that could
  ## be used within a COMPASSContainer

  n <- 10 ## number of samples
  k <- 4 ## number of markers

  sid_vec <- paste0("sid", 1:n) ## sample ids; unique names used to denote samples
  iid_vec <- rep_len( paste0("iid", 1:(n/10) ), n ) ## individual ids
  marker_names <- c("TNFa", "IL2", "IL4", "IL6")

  ## Generate n sets of 'flow' data -- a list of matrices, each row

  ...
\end{verbatim}
## is a cell, each column is fluorescence intensities on a particular
## channel / marker

data <- replicate(n, {
  nrow <- round(runif(1) * 1E4 + 1000)
  ncol <- k
  vals <- rexp( nrow * ncol, runif(1, 1E-5, 1E-3) )
  output <- matrix(vals, nrow, ncol)
  colnames(output) <- marker_names
  return(output)
})

names(data) <- sid_vec

## Put it into a GatingSet

fs <- flowSet( lapply(data, flowFrame) )
gs <- GatingSet(fs)

## Add some dummy metadata

meta <- pData(gs)
meta$PTID <- 1:10
pData(gs) <- meta

gate <- rectangleGate( list(TNFa=c(-Inf,Inf)))
gs_pop_add(gs, gate, parent="root", name="dummy")

## Add dummy gate

## Make some gates, and apply them

invisible(lapply(marker_names, function(marker) {
  .gate <- setNames( list( c( rexp(1, runif(1, 1E-5, 1E-3)), Inf ) ), marker )
  gate <- rectangleGate(.gate=.gate)
  gs_pop_add(gs, gate, parent="dummy", name=paste0(marker, "+"))
}))

recompute(gs)

## Map node names to channel names

map=list(
  "TNFa+"="TNFa",
  "IL2+"="IL2",
  "IL4+"="IL4",
  "IL6+"="IL6"
)

## Pull out the data as a COMPASS-friendly dataset

node <- "dummy"
map <- map

system.time(
  output <- GetThresholdedIntensities(gs, "dummy", map)
)

system.time(
  output <- COMPASSContainerFromGatingSet(gs, "dummy", individual_id="PTID")
)
markers

```r
str(output)
```

---

### markers

<table>
<thead>
<tr>
<th>Markers</th>
</tr>
</thead>
</table>

#### Description

Returns the markers associated with an experiment.

#### Usage

```r
markers(object)
```

#### Arguments

- `object`: An R object.

---

### melt_

| Make a 'Wide' data set 'Long' |

#### Description

Inspired by `reshape2::melt`, we melt data.frames and matrices. This function is built for speed.

#### Usage

```r
melt_(data, ...)
```

```r
## S3 method for class 'data.frame'
melt_(
data,
id.vars,
measure.vars,
variable.name = "variable",
...,
value.name = "value"
)
```

```r
## S3 method for class 'matrix'
melt_(data, ...)
```
Arguments

- **data**: The data.frame to melt.
- **id.vars**: Vector of id variables. Can be integer (variable position) or string (variable name). If blank, we use all variables not in `measure.vars`.
- **measure.vars**: Vector of measured variables. Can be integer (variable position) or string (variable name). If blank, we use all variables not in `id.vars`.
- **variable.name**: Name of variable used to store measured variable names.
- **value.name**: Name of variable used to store values.

Details

If items to be stacked are not of the same internal type, they will be promoted in the order logical > integer > numeric > character.

---

merge.COMPASSContainer

*Merge Two COMPASSContainers*

Description

This function merges two COMPASSContainers.

Usage

```r
## S3 method for class 'COMPASSContainer'
merge(x, y, ...)
```

Arguments

- **x**: A COMPASSContainer.
- **y**: A COMPASSContainer.
- **...**: other arguments passed to `COMPASSContainer` call.

Examples

```r
## Chop the example COMPASSContainer into two, then merge it back together
CC1 <- subset(CC, trt == "Control")
CC2 <- subset(CC, trt == "Treatment")
merged <- merge(CC1, CC2)
res <- identical(CC, merge(CC1, CC2)) ## should return TRUE in this case
stopifnot( isTRUE(res) )
```
**metadata**

**Metadata**

### Description

Functions for getting and setting the metadata associated with an object.

### Usage

```r
metadata(x)
```

### S3 method for class `COMPASSContainer`

```r
metadata(x)
```

### S3 method for class `COMPASSResult`

```r
metadata(x)
```

```r
metadata(x) <- value
```

### S3 replacement method for class `COMPASSContainer`

```r
metadata(x) <- value
```

### Arguments

- **x**
  
  An R object.

- **value**
  
  An R object appropriate for storing metadata in object `x`; typically a `data.frame`.

---

**pheatmap**

*A function to draw clustered heatmaps.*

### Description

A function to draw clustered heatmaps where one has better control over some graphical parameters such as cell size, etc.

### Usage

```r
pheatmap(
  mat,
  color = colorRampPalette(rev(brewer.pal(n = 7, name = "RdYlBu")))(100),
  kmeans_k = NA,
  breaks = NA,
  border_color = "grey60",
  cellwidth = NA,
  cellheight = NA,
```
scale = "none",
cluster_rows = TRUE,
cluster_cols = TRUE,
clustering_distance_rows = "euclidean",
clustering_distance_cols = "euclidean",
clustering_method = "complete",
treeheight_row = ifelse(cluster_rows, 50, 0),
treeheight_col = ifelse(cluster_cols, 50, 0),
legend = TRUE,
legend_breaks = NA,
legend_labels = NA,
annotation = NA,
annotation_colors = NA,
annotation_legend = TRUE,
drop_levels = TRUE,
show_rownames = TRUE,
show_colnames = TRUE,
main = NA,
fontsize = 10,
fontsize_row = fontsize,
fontsize_col = fontsize,
display_numbers = FALSE,
number_format = "%2f",
fontsize_number = 0.8 * fontsize,
filename = NA,
width = NA,
height = NA,
row_annotation = NA,
row_annotation_legend = TRUE,
row_annotation_colors = NA,
cytokine_annotation = NA,
headerplot = NA,
polar = FALSE,
order_by_max_functionality = TRUE,
...}

Arguments

mat numeric matrix of the values to be plotted.

color vector of colors used in heatmap.

kmeans_k the number of kmeans clusters to make, if we want to aggregate the rows before drawing heatmap. If NA then the rows are not aggregated.

breaks a sequence of numbers that covers the range of values in mat and is one element longer than color vector. Used for mapping values to colors. Useful, if needed to map certain values to certain colors, to certain values. If value is NA then the breaks are calculated automatically.

border_color color of cell borders on heatmap, use NA if no border should be drawn.
cellwidth  individual cell width in points. If left as NA, then the values depend on the size of plotting window.

cellheight individual cell height in points. If left as NA, then the values depend on the size of plotting window.

scale character indicating if the values should be centered and scaled in either the row direction or the column direction, or none. Corresponding values are "row", "column" and "none"

cluster_rows boolean values determining if rows should be clustered,

cluster_cols boolean values determining if columns should be clustered.

clustering_distance_rows distance measure used in clustering rows. Possible values are "correlation" for Pearson correlation and all the distances supported by dist, such as "euclidean", etc. If the value is none of the above it is assumed that a distance matrix is provided.

clustering_distance_cols distance measure used in clustering columns. Possible values the same as for clustering_distance_rows.

clustering_method clustering method used. Accepts the same values as hclust.

treeheight_row the height of a tree for rows, if these are clustered. Default value 50 points.

treeheight_col the height of a tree for columns, if these are clustered. Default value 50 points.

legend logical to determine if legend should be drawn or not.

legend_breaks vector of breakpoints for the legend.

legend_labels vector of labels for the legend_breaks.

annotation data frame that specifies the annotations shown on top of the columns. Each row defines the features for a specific column. The columns in the data and rows in the annotation are matched using corresponding row and column names. Note that color schemes takes into account if variable is continuous or discrete.

annotation_colors list for specifying annotation track colors manually. It is possible to define the colors for only some of the features. Check examples for details.

annotation_legend boolean value showing if the legend for annotation tracks should be drawn.

drop_levels boolean value showing if the legend for annotation tracks should be drawn.

show_rownames logical to determine if unused levels are also shown in the legend

show_colnames boolean specifying if column names are be shown.

main the title of the plot

fontsize base fontsize for the plot

fontsize_row fontsize for rownames (Default: fontsize)

fontsize_col fontsize for colnames (Default: fontsize)

display_numbers logical determining if the numeric values are also printed to the cells.
number_format  format strings (C printf style) of the numbers shown in cells. For example "%.2f" shows 2 decimal places and "%.1e" shows exponential notation (see more in gettext).

fontsize_number  fontsize of the numbers displayed in cells

filename  file path where to save the picture. Filetype is decided by the extension in the path. Currently following formats are supported: png, pdf, tiff, bmp, jpeg. Even if the plot does not fit into the plotting window, the file size is calculated so that the plot would fit there, unless specified otherwise.

width  manual option for determining the output file width in inches.

height  manual option for determining the output file height in inches.

row_annotation  data frame that specifies the annotations shown on the rows. Each row defines the features for a specific row. The rows in the data and rows in the annotation are matched using corresponding row names. The category labels are given by the data frame column names.

row_annotation_legend  same interpretation as the column parameters.

row_annotation_colors  same interpretation as the column parameters

cytokine_annotation  A data.frame of factors, with either levels 0 = unexpressed, 1 = expressed, or optionally with a third level -1 = 'left out'. of the categories for each column. They will be colored by their degree of functionality and ordered by degree of functionality and by amount of expression if column clustering is not done.

headerplot  is a list with two components, order and data. Order tells how to reorder the columns of the matrix.

polar  Boolean; if TRUE we draw a polar legend. Primarily for internal use. Data is some summary statistic over the columns which will be plotted in the header where the column cluster tree usually appears. Cytokine ordering is ignored when the headerplot argument is passed.

order_by_max_functionality  Boolean; re-order the cytokine labels by maximum functionality?

...  graphical parameters for the text used in plot. Parameters passed to grid.text, see gpar.

Details

The function also allows to aggregate the rows using kmeans clustering. This is advisable if number of rows is so big that R cannot handle their hierarchical clustering anymore, roughly more than 1000. Instead of showing all the rows separately one can cluster the rows in advance and show only the cluster centers. The number of clusters can be tuned with parameter kmeans_k.

Value

Invisibly a list of components

- tree_row the clustering of rows as hclust object
- tree_col the clustering of columns as \texttt{hclust} object
- kmeans the kmeans clustering of rows if parameter \texttt{kmeans_k} was specified

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\textbf{Examples}

```r
# Generate some data
test = matrix(rnorm(200), 20, 10)
test[1:10, seq(1, 10, 2)] = test[1:10, seq(1, 10, 2)] + 3
test[11:20, seq(2, 10, 2)] = test[11:20, seq(2, 10, 2)] + 2
test[15:20, seq(2, 10, 2)] = test[15:20, seq(2, 10, 2)] + 4
colnames(test) = paste("Test", 1:10, sep = "")
rownames(test) = paste("Gene", 1:20, sep = "")

# Draw heatmaps
pheatmap(test)
pheatmap(test, kmeans_k = 2)
pheatmap(test, scale = "row", clustering_distance_rows = "correlation")
pheatmap(test, color = colorRampPalette(c("navy", "white", "firebrick3"))(50))
pheatmap(test, cluster_row = FALSE)
pheatmap(test, legend = FALSE)
pheatmap(test, display_numbers = TRUE)
pheatmap(test, display_numbers = TRUE, number_format = "%e")
pheatmap(test, cluster_row = FALSE, legend_breaks = -1:4, legend_labels = c("0", "1e-4", "1e-3", "1e-2", "1e-1", "1"))
pheatmap(test, cellwidth = 15, cellheight = 12, main = "Example heatmap")
#pheatmap(test, cellwidth = 15, cellheight = 12, fontsize = 8, filename = "test.pdf")

# Generate column annotations
annotation = data.frame(Var1 = factor(1:10 %% 2 == 0,
labels = c("Class1", "Class2")), Var2 = 1:10)
annotation$Var1 = factor(annotation$Var1, levels = c("Class1", "Class2", "Class3"))
rownames(annotation) = paste("Test", 1:10, sep = "")

pheatmap(test, annotation = annotation)
pheatmap(test, annotation = annotation, annotation_legend = FALSE)
pheatmap(test, annotation = annotation, annotation_legend = FALSE, drop_levels = FALSE)

# Specify colors
Var1 = c("navy", "darkgreen")
names(Var1) = c("Class1", "Class2")
Var2 = c("lightgreen", "navy")
ann_colors = list(Var1 = Var1, Var2 = Var2)

#Specify row annotations
row_ann <- data.frame(foo=gl(2,nrow(test)/2),`Bar`=relevel(gl(2,nrow(test)/2),"2"))
```r
rownames(row_ann) <- rownames(test)
pheatmap(test, annotation = annotation, annotation_legend = FALSE,
         drop_levels = FALSE, row_annotation = row_ann)

# Using cytokine annotations
M <- matrix(rnorm(8 * 20), ncol = 8)
row_annotation <- data.frame(A = gl(4, nrow(M) / 4), B = gl(4, nrow(M) / 4))
eg <- expand.grid(factor(c(0, 1)), factor(c(0, 1)), factor(c(0, 1)))
colnames(eeg) <- c("IFNg", "TNFa", "IL2")
rownames(eeg) <- apply(eeg, 1, function(x) paste0(x, collapse = "")
rownames(M) <- 1:nrow(M)
colnames(M) <- rownames(eeg)
cytokine_annotation = eeg
pheatmap(M, annotation = annotation, row_annotation = row_annotation,
         annotation_legend = TRUE, row_annotation_legend = TRUE,
         cluster_rows = FALSE, cytokine_annotation = cytokine_annotation, cluster_cols = FALSE)

# Specifying clustering from distance matrix
drows = dist(test, method = "minkowski")
dcols = dist(t(test), method = "minkowski")
pheatmap(test, clustering_distance_rows = drows, clustering_distance_cols = dcols)
```

---

**plot.COMPASSResult**

**Plot a COMPASSResult**

**Description**

This function can be used to visualize the mean probability of response; that is, the probability
that there is a difference in response between samples subjected to the 'treatment' condition, and
samples subjected to the 'control' condition.

**Usage**

```r
## S3 method for class 'COMPASSResult'
plot(
  x,
  y,
  subset = NULL,
  threshold = 0.01,
  minimum_dof = 1,
  maximum_dof = Inf,
  must_express = NULL,
  row_annotation,
  palette = colorRampPalette(brewer.pal(10, "Purples"))(20),
  show_rownames = FALSE,
  show_colnames = FALSE,
  measure = NULL,
  order_by = FunctionalityScore,
  order_by_max_functionality = TRUE,
)```
Arguments

x  An object of class COMPASSResult.

y  This argument gets passed to row_annotation, if row_annotation is missing. It can be used to group rows (individuals) by different conditions as defined in the metadata.

subset  An R expression, evaluated within the metadata, used to determine which individuals should be kept.

threshold  A numeric threshold for filtering under-expressed categories. Any categories with mean score < threshold are removed.

minimum_dof  The minimum degree of functionality for the categories to be plotted.

maximum_dof  The maximum degree of functionality for the categories to be plotted.

must_express  A character vector of markers that should be included in each subset plotted. For example, must_express=c("TNFa & IFNg") says we include only subsets that are positive for both TNFa or IFNg, while must_express=c("TNFa", "IFNg") says we should keep subsets which are positive for either TNFa or IFNg.

row_annotation  A vector of names, pulled from the metadata, to be used for row annotation.

palette  The colour palette to be used.

show_rownames  Boolean; if TRUE we display row names (ie, the individual ids).

show_colnames  Boolean; if TRUE we display column names (ie, the column name associated with a cytokine; typically not needed)

measure  Optional. By default, we produce a heatmap of the mean gammas produced in a model fit. We can override this by supplying a matrix of suitable dimension as well; these can be generated with the Posterior* functions – see Posterior for examples.

order_by  Order rows within a group. This should be a function; e.g. FunctionalityScore, mean, median, and so on. Set this to NULL to preserve the original ordering of the data.

order_by_max_functionality  Order columns by functionality within each degree subset. to TRUE.

markers  specifies a subset of markers to plot. default is NULL, which means all markers.

...  Optional arguments passed to pheatmap.

Value

The plot as a grid object (grob). It can be redrawn with e.g. grid::grid.draw().
Examples

## visualize the mean probability of response
plot(CR)

## visualize the proportion of cells belonging to a category
plot(CR, measure=PosteriorPs(CR))

plot2

Plot a pair of COMPASSResults

Description

This function can be used to visualize the mean probability of response – that is, the probability that there is a difference in response between samples subjected to the 'treatment' condition, and samples subjected to the 'control' condition.

Usage

plot2(
  x,
  y,
  subset,
  threshold = 0.01,
  minimum_dof = 1,
  maximum_dof = Inf,
  must_express = NULL,
  row_annotation = NULL,
  palette = NA,
  show_rownames = FALSE,
  show_colnames = FALSE,
  ...
)

Arguments

x An object of class COMPASSResult.
y An object of class COMPASSResult.
subset An R expression, evaluated within the metadata, used to determine which individuals should be kept.
threshold A numeric threshold for filtering under-expressed categories. Any categories with mean score < threshold are removed.
minimum_dof The minimum degree of functionality for the categories to be plotted.
maximum_dof The maximum degree of functionality for the categories to be plotted.
must_express  A character vector of markers that should be included in each subset plotted. For example, must_express=c("TNFa & IFNg") says we include only subsets that are positive for both TNFa or IFNg, while must_express=c("TNFa", "IFNg") says we should keep subsets which are positive for either TNFa or IFNg.

row_annotation  A vector of names, pulled from the metadata, to be used for row annotation.

palette  The colour palette to be used.

show_rownames  Boolean; if TRUE we display row names (ie, the individual ids).

show_colnames  Boolean; if TRUE we display column names (ie, the column name associated with a cytokine; typically not needed)

...  Optional arguments passed to pheatmap.

Value

The plot as a grid object (grob). It can be redrawn with e.g. grid::grid.draw().

Description

This function can be used to visualize the mean probability of response; that is, the probability that there is a difference in response between samples subjected to the 'treatment' condition, and samples subjected to the 'control' condition. This version is used for plotting multiple COMPASS-Result objects. The COMPASS runs must all use the same markers. The code is heavily based on the plot.COMPASSResult and plot2 functions. Not all options from plot.COMPASSResult are supported yet.

Usage

plotCOMPASSResultStack(
  x,
  threshold = 0.01,
  minimum_dof = 1,
  maximum_dof = Inf,
  row_annotation,
  variable,
  palette = colorRampPalette(brewer.pal(9, "Purples"))(20),
  show_rownames = FALSE,
  ...
)
PolyfunctionalityScore

**Arguments**

- `x` A named list of objects of class `COMPASSResult`. The names are values of type `variable`.
- `threshold` A numeric threshold for filtering under-expressed categories. Any categories with mean score < `threshold` are removed.
- `minimum_dof` The minimum degree of functionality for the categories to be plotted.
- `maximum_dof` The maximum degree of functionality for the categories to be plotted.
- `row_annotation` A vector of names, pulled from the metadata, to be used for row annotation.
- `variable` What to call the variable that is different across the objects in `x`.
- `palette` The colour palette to be used.
- `show_rownames` Boolean; if `TRUE` we display row names (ie, the individual ids).
- ... Optional arguments passed to `pheatmap`.

**Value**

The plot as a grid object (`grob`). It can be redrawn with e.g. `grid::grid.draw()`.

**Examples**

```r
# Not run:
# allCompassResults is a list of 4 COMPASSResults
names(allCompassResults) <- c("Antigen 85A", "CFP-10", "CMV", "ESAT-6")
plotCOMPASSResultStack(allCompassResults,
  row_annotation = c("Antigen", "PATIENT ID", "Time"),
  variable = "Antigen", show_rownames = FALSE,
  main = "Heatmap of Mean Probability of Response to Antigens, CD8+",
  fontsize = 14, fontsize_row = 13, fontsize_col = 11)
```

## End(Not run)

---

**PolyfunctionalityScore**

*Compute the Polyfunctionality Score for each subject fit in a COMPASS model*

**Description**

Computes the Polyfunctionality score for each observation from the gamma matrix of a COMPASS model fit. The scores are normalized to one.
Usage

PolyfunctionalityScore(x, degree, n, markers = NULL)

## S3 method for class 'COMPASSResult'
PolyfunctionalityScore(x, degree, n, markers = NULL)

## Default S3 method:
PolyfunctionalityScore(x, degree, n, markers = NULL)

Arguments

x An object of class COMPASSResult, as returned by COMPASS. Alternatively, a
matrix of functionality scores.

degree A vector of weights. If missing, this is the 'degree of functionality', that is, the
number of markers expressed in a particular category.

n The total number of markers. This is inferred when x is a COMPASSResult, and
is unused in that case.

markers A character specifying the markers for which to compute the score. Must
match names in markers().

Value

A numeric vector of polyfunctionality scores.

Examples

PolyfunctionalityScore(CR)

Description

These functions can be used to retrieve different posterior measures from a COMPASS fit object.

Usage

Posterior(x)

PosteriorDiff(x)

PosteriorLogDiff(x)

PosteriorPs(x)

PosteriorPu(x)
print.COMPASSContainer

Arguments

  x  An object of class COMPASSResult.

Details

The posterior items retrieved are described as follows:

  PosteriorPs: The posterior estimate of the proportion of cells in the stimulated sample.
  PosteriorPu: The posterior estimate of the proportion of cells in the unstimulated sample.
  PosteriorDiff: The difference in posterior proportions, as described above.
  PosteriorLogDiff: The difference in the log posterior proportions, as described above.

Examples

  Posterior(CR)
  PosteriorPs(CR)
  PosteriorPu(CR)
  PosteriorDiff(CR)
  PosteriorLogDiff(CR)

print.COMPASSContainer

Print a COMPASSContainer Object

Description

This function prints a COMPASSContainer object, giving basic information about the object and the data it encapsulates.

Usage

  ## S3 method for class 'COMPASSContainer'
  print(x, ...)

Arguments

  x  An object of class COMPASSContainer.
  ... Optional arguments passed to cat.

Examples

  print(CC)
print.COMPASSResult | Print a COMPASSResult Object

Description
This function prints basic information about the model fit by a COMPASS call.

Usage
```r
## S3 method for class 'COMPASSResult'
print(x, ...)
```

Arguments
- `x`: An object of class COMPASSResult.
- `...`: Optional arguments; currently unused.

Examples
```
print(CR)
```

Response | Compute a response probability from COMPASS mcmc samples.

Description
Compute a response probability based on the selected markers, evaluating the probability that a subject exhibits a response of size `degree` or greater. i.e., the probability of at least `degree` markers exhibiting an antigen specific response.

Usage
```
Response(x, markers, degree, max.prob, at_least_n)
```

```r
## S3 method for class 'COMPASSResult'
Response(x, markers = NULL, degree = 1, max.prob = FALSE, at_least_n = NULL)
```

Arguments
- `x`: a COMPASSResult object.
- `markers`: a vector of marker names.
- `degree`: the numeric degree of functionality to test.
- `max.prob`: logical Use the max probability rather than the average across subsets. Defaults to FALSE.
- `at_least_n`: logical response of degree `x` or greater with at_least_n subsets responding.
Details

The response is computed from the sampled Gamma matrix, subsetting on the selected markers, and

Value

A vector of response probabilities for each subject.

Examples

Response(CR, markers = c("M1","M2","M3"), degree = 2)

| scores | Fetch the table of scores and metadata from a COMPASSResult Object |

Description

This function extracts the functionality and polyfunctionality scores from a COMPASS result merged with the sample metadata table, accounting for any dropped samples due to filtering.

Usage

scores(x, markers = NULL)

Arguments

x A COMPASSResult object.

markers A character vector of markers for which to compute the scores. Defaults to all markers. Must match the names returned by markers().

Examples

scores(CR)
select_compass_pops  Flag COMPASS boolean populations

Description

Returns a boolean vector indexing cell populations in cellpops that match the pattern for boolean combinations of markers.

Usage

select_compass_pops(cellpops, markers)

Arguments

cellpops vector of character names of cell populations.

markers vector of character names of markers in the order they appear in the population names.

Details

If markers A, B, C, D make up the population names in cellpops and they the names match the pattern e.g. "A+B-C+D+,Count" (typical of exports from some gating tools), then markers should be a vector of markers in the same order they appear in cellpops.

Value

A boolean vector indexing cellpops with TRUE for populations matching the pattern.

See Also

translate_marker_names

Examples

#Generate some population names
markers = LETTERS[1:4]
pos = c("+","-")

popnames = apply(expand.grid(pos,pos,pos,pos),1,
  function(x)paste(paste(paste(markers,x,sep=""),
    collapse=""),",Count",sep=""))

popnames = sample(c(popnames,paste(paste(markers,sample(c("+","-"),
    length(markers),replace=TRUE),sep=""),",Count",sep=""))),

popnames[select_compass_pops(popnames,LETTERS[1:4])]
Description

This function takes a COMPASSResult object, and generates a local Shiny application for visualizing the results.

Usage

shinyCOMPASS(
  x, 
  dir = NULL, 
  meta.vars, 
  facet1 = "None", 
  facet2 = "None", 
  facet3 = "None", 
  main = "Heatmap of Ag-Specificity Posterior Probabilities", 
  stimulation = NULL, 
  launch = TRUE, 
  ...
)

Arguments

x  An object of class COMPASSResult.
dir  A location to write out the .rds files that will be loaded and used by the Shiny application.
meta.vars  A character vector of column names that should be used for potential facetting in the Shiny app. By default, we take all metadata variables; you may want to limit this if you know certain variables are not of interest.
facet1, facet2, facet3  Default values for facets in the Shiny app. Each should be the name of a single vector in the metadata.
main  A title to give to the heatmap and subset histogram plots.
stimulation  The name of the stimulation applied. If this is NULL, the stimulations used are inferred from the data (ie, the COMPASS call used).
launch  Boolean; if TRUE we launch the Shiny application. Otherwise, the user can launch it manually by navigating to the directory dir and running shiny::runApp().
...

See Also

shinyCOMPASSDepso, for identifying packages that you need in order to run the Shiny application.
Examples

if (interactive()) {
    oldOpt <- getOption("example.ask")
    options(example.ask=FALSE)
    on.exit( options(example.ask=oldOpt) )
    shinyCOMPASS(CR)
    options(example.ask=TRUE)
}

shinyCOMPASSDeps List Shiny Dependencies

Description

This function can be used to identify the packages still needed in order to launch the Shiny app.

Usage

    shinyCOMPASSDeps(VERBOSE = TRUE)

Arguments

verbo se                  Boolean; if TRUE we print installation instructions to the screen.

Examples

    shinyCOMPASSDeps()

SimpleCOMPASS Fit the discrete COMPASS Model

Description

This function fits the COMPASS model from a user-provided set of stimulated / unstimulated matrices. See the NOTE for important details.

Usage

    SimpleCOMPASS(
        n_s,
        n_u,
        meta,
        individual_id,
        iterations = 10000,
        replications = 8,
        verbose = TRUE,
        seed = 100
    )

Arguments

n_s The cell counts for stimulated cells.
n_u The cell counts for unstimulated cells.
meta A data.frame of metadata, describing the individuals in the experiment. Each row in meta should correspond to a row in data. There should be one row for each subject; i.e., one row for each element of n_s and n_u.
individual_id The name of the vector in meta that denotes the individuals from which samples were drawn.
iters The number of iterations (per 'replication') to perform.
replications The number of 'replications' to perform. In order to conserve memory, we only keep the model estimates from the last replication.
verbose Boolean; if TRUE we output progress information.
seed A seed for the random number generator. Defaults to 100.

Value

A list with class COMPASSResult with two components, the fit containing parameter estimates and parameter acceptance rates, and data containing the generated data used as input for the model.

Note

n_s and n_u counts matrices should contain ALL $2^M$ possible combinations of markers, even if they are 0 for some combinations. The code expects the marker combinations to be named in the following way: "M1&M2&!M3" means the combination represents cells expressing marker "M1" and "M2" and not "M3". For 3 markers, there should be 8 such combinations, such that n_s and n_u have 8 columns.

Examples

set.seed(123)
n <- 10 ## number of subjects
k <- 3 ## number of markers

## generate some sample data
iid_vec <- paste0("iid_", 1:n) # Subject id
data <- replicate(2*n, {
  nrow <- round(runif(1) * 1E4 + 1000)
  ncol <- k
  vals <- rexp( nrow * ncol, runif(1, 1E-5, 1E-3) )
  vals[ vals < 2000 ] <- 0
  output <- matrix(vals, nrow, ncol)
  output <- output[ apply(output, 1, sum) > 0, ]
  colnames(output) <- paste0("M", 1:k)
  return(output)
})

meta <- cbind(iid=iid_vec, data.frame(trt=rep( c("Control", "Treatment"), each=n/2 )))
## generate counts for n_s, n_u
n_s <- CellCounts(data[1:n], Combinations(k))
n_u <- CellCounts(data[(n+1):(2*n)], Combinations(k))
rownames(n_s) = unique(meta$iid)
rownames(n_u) = rownames(n_s)
## A smaller number of iterations is used here for running speed;
## prefer using more iterations for a real fit
scr = SimpleCOMPASS(n_s, n_u, meta, "iid", iterations=1000)

---

### subset.COMPASSContainer

**Subset a COMPASSContainer**

**Description**

Use this function to subset a COMPASSContainer.

**Usage**

```r
## S3 method for class 'COMPASSContainer'
subset(x, subset, ...)
```

**Arguments**

- `x`: A COMPASSContainer.
- `subset`: A logical expression, evaluated within the metadata, indicating which entries to keep.
- `...`: Other arguments passed to 'COMPASSContainer' call.

**Examples**

```r
subset(CC, iid == "iid_1")
```

---

### summary.COMPASSContainer

**Summarize a COMPASSContainer Object**

**Description**

This function prints summary information about a COMPASSContainer object – the number of samples, basic information about the metadata, and so on.

**Usage**

```r
## S3 method for class 'COMPASSContainer'
summary(object, ...)
```
** Arguments  

object An object of class COMPASSContainer.  
... Optional arguments; currently ignored.  

** Examples  

summary(CC)
Arguments

data A COMPASSContainer.
subset An expression, evaluated within the metadata, defining the subset of data over which the counts are computed. If left unspecified, the counts are computed over all samples.
aggregate Boolean; if TRUE we sum over the individual, to get total counts across samples for each individual.

Examples

TotalCellCounts(CC, trt == "Treatment")
TotalCellCounts(CC, trt == "Control")
TotalCellCounts(CC)

describe_marker_names

Translate marker names to format use by COMPASS

Description

Translate boolean population names from format exported by common software tools to a format used by COMPASS.

Usage

describe_marker_names(cellpops)

Arguments

cellpops character vector of cell population names.

Value

character vector of cell population names used by COMPASS

See Also

select_compass_pops

Examples

#Generate marker names
markers = LETTERS[1:4]
pos = c("+","-"")
popnames = apply(expand.grid(pos,pos,pos,pos),1,
function(x) paste(paste(markers,x,sep=""),
collapse=""),"Count",sep=""))
popnames = sample(c(popnames,
paste(paste(markers, sample(c("+","-"), length(markers), replace=TRUE), sep=""), ",Count", sep="")))

popnames = popnames[select_compass_pops(popnames, LETTERS[1:4])]
#Translate
translate_marker_names(popnames)

transpose_list Transpose a List

Description
Transpose a matrix-like list.

Usage
transpose_list(x)

Arguments
x An R list.

Examples
l <- list( 1:3, 4:6, 7:9 )
stopifnot(identical(
  transpose_list( transpose_list(l) ), l
))

UniqueCombinations Generate Unique Combinations

Description
Generate all possible unique combinations of x. Primarily used as a helper function for CellCounts, but may be occasionally useful to the end user.

Usage
UniqueCombinations(x, as.matrix)

## S3 method for class 'COMPASSContainer'
UniqueCombinations(x, as.matrix = FALSE)

## Default S3 method:
UniqueCombinations(x, as.matrix = FALSE)
**Arguments**

- **x**
  - Either a COMPASSContainer, or a list of matrices.

- **as.matrix**
  - Boolean; if TRUE we return results as a matrix; otherwise, we return the results as a list.

**Examples**

```
UniqueCombinations(CC)
```
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