Package 'RcometsAnalytics'

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Type Package

Title Comets Analytics for consortium based metabolomic analyses
Version 1.9.9.13
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Description This R package support all cohort-specific analyses of the COMETS consortium. Data are not saved in the system but output must be downloaded and submitted for meta-analyses. This package can be used in several ways: supports the CBIIT implementation of comets-analytics, local shiny based add-in called shinycomets, or console based analysis. See example vignette. The version of the package is noted as: level 1 and 2 reflect the comets-analytics version, 3rd level reflects the number of major revision and 4th level for bug fixes.
Depends R (>= $3.5.0$)
Imports readxl, rio, dplyr, plyr, plotly, tidyr, heatmaply, stringr, data.table, caret, subselect, broom, psych, MASS, ppcor
Suggests Hmisc, knitr, testthat, rmarkdown
VignetteBuilder knitr
License GPL-3
RoxygenNote 7.1.2
NeedsCompilation no
<pre>URL http://comets-analytics.org/</pre>
BugReports https://github.com/CBIIT/nci-webtools-comets-analytics/issues
R topics documented: correlation.options Effects
Errors_Warnings
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lm.options
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corr	Plation options options list for model="correlation"	

Description

A list of 1:

• method Correlation method to use. It must be one of "spearman", "pearson", "kendall". The default value is "spearman".

Examples

```
model.options <- list(method="pearson")</pre>
```

Effects

Effects table

Description

The Effects data frame contains the estimates for each exposure, and will contain multiple rows for categorical exposure variables. Depending on the model run and options specified, all the below names may not appear in the data frame.

- corr The correlation between term and the outcome
- estimate The regression coefficient of term
- estimate.lower The lower confidence limit for term
- estimate.upper The upper confidence limit for term
- \bullet $\mbox{exp.estimate}$ The exponentiated regression coefficient of \mbox{term}
- exp.estimate.lower The exponentiated lower confidence limit for term
- exp.estimate.upper The exponentiated upper confidence limit for term
- exp.std.error The standard error of exp.estimate from the delta method

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- exposurespec Exposure variable
- outcomespec Outcome variable
- pvalue The p-value of the test
- run Run number that can be used to link with the ModelSummary table
- statistic The test statistic for term
- std.error The standard error of estimate
- stratavar Stratum variable(s)
- strata Stratum level(s)
- term Variable in the model

Details

Missing values will appear if a model did not converge, produced an error, or not run because of too many missing values in the outcome.

Errors_Warnings

Errors and Warnings table

Description

Columns in the Errors_Warnings table.

- type WARNING or ERROR
- object The object that produced the warning or error. This is typically a variable or a particular stratum.
- message Message describing the warning or error

Details

The kinds of warnings and errors stored in this matrix are ones that apply to all models or all outcomes for an exposure variable. An error message for a particular exposure-outcome pair will be stored in the message column of the ModelSummary table.

 ${\tt filter COMETS input}$

Function that subsets input data based on "where variable"

Description

Function that subsets input data based on "where variable"

```
filterCOMETSinput(readData, where = NULL)
```

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Arguments

readData list from readComets

where users can specify which subjects to perform the analysis by specifying this pa-

rameter. 'where' expects a vector with a variable name, a comparison operator ("<", ">", "=","<=",">="), and a value. For example, "where = c("Gender=Female")".

Value

filtered list

getModelData Prepares data for the models to be run as specified in the input. Can

be run in interactive or batch mode. Each model is checked for validity

(correlation between predictors, zero variance, etc.).

Description

Prepares data for the models to be run as specified in the input. Can be run in interactive or batch mode. Each model is checked for validity (correlation between predictors, zero variance, etc.).

Usage

```
getModelData(
  readData,
  modelspec = "Batch",
  modlabel = "",
  outcomes = "All metabolites",
  exposures = "",
  adjvars = NULL,
  strvars = NULL,
  wgtvar = NULL,
  offvar = NULL,
  where = NULL
```

Arguments

readData	List from readCOMETSinput
modelspec	How model is specified (Interactive or Batch). The default is Batch
modlabel	If batch, chosen model specified by batch mode (the MODEL column in the Models sheet). If interactive, then the model label.
outcomes	If Interactive, a vector of outcome variables (see details), the default is All metabolites)
exposures	If Interactive, a vector of exposure variables (see details)
adjvars	If Interactive, a vector adjustment covariates (see details)
strvars	If Interactive, stratification covariates (see details)
wgtvar	If Interactive, a variable of weights (see details)
offvar	If Interactive, an offset variable (see details)

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where

users can specify which subjects to perform the analysis on by specifying this parameter. 'where' expects a vector of strings with a variable name, a comparison operator (e.g. "<", ">=", ">=", "!=", "="), and a value. For example, where = c("age>50", "bmi > 22") uses all subjects with age > 50 AND bmi > 22. Note that when running in Batch mode, rules in the WHERE column of the Models sheet must be separated by a comma.

Details

All metabolite variables specified should be listed in the metabolite_name column of the Metabolites sheet of the Excel file. All non-metabolite variables should be listed in the VARREFERENCE column of the VarMap sheet. The wgtvar and offvar are only used when the model function is glm or lm, see the model option in options.

Value

a list comprising:

1: subset data: gdta

2: exposure variables: ccovs

3: outcome variables: rcovs

4: adjustment variables: acovs

5: stratification variable: scovs

6: model specification: modspec

7: model label: modlab

8: whether all metabolites vs all metabolites is run: allvsall

9: weight variables: wgtcov 10: offset variables: offcov

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,modlabel="1 Gender adjusted")</pre>
```

glm.options

options list for glm

Description

A list of 6:

- family One of: "binomial", "gaussian", "Gamma", "inverse.gaussian", "poisson", "quasi", "quasibinomial", "quasipoisson". The default is "gaussian".
- link NULL or a string for the link function to use (see family). The default is to use the canonical link for family.
- weights A variable name to specify weights. The default is NULL.
- offset A variable name to specify an offset. The default is NULL.
- control See glm. The default is glm. control.
- singular.ok See glm. The default is TRUE.

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Examples

```
model.options <- list(family="binomial", weights="weightVarInData")</pre>
```

lm.options

options list for lm

Description

A list of 4:

- weights A variable name to specify weights. The default is NULL.
- offset A variable name to specify an offset. The default is NULL.
- tol See 1m. The default is 1e-7.
- singular.ok See lm. The default is TRUE.

Examples

```
model.options <- list(weights="weightVarInData")</pre>
```

ModelSummary

ModelSummary table

Description

The ModelSummary data frame contains one row of model summary results for each exposure/outcome combination. Depending on the model run and options specified, all the below names may not appear in the data frame.

- · adjspec Original adjustment variables specified
- adjvars Adjustment variables included in the model
- adjvars.removed Adjustment variables removed from the model
- adj_uid Adjustment variable universal ids
- adj.r.squared Adjusted R-squared
- · aic Akaike information criterion
- bic Bayesian information criterion
- cohort String passed into runModel
- converged TRUE or FALSE for model convergence
- · deviance Deviance of the fitted model
- df.null NULL model degrees of freedom
- df.residual Residual degrees of freedom
- exposure Exposure variable
- exposure_uid Exposure universal id
- exposurespec Exposure variable
- loglik Log-likelihood of the fitted model

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- message Error message produced from the modeling function
- model Model label from getModelData
- model_function Model function used in runModel
- · nobs Number of observations used
- null.deviance Deviance of the NULL model
- outcome Outcome variable
- outcomespec Outcome variable
- outcome_uid Outcome universal id
- run Run number that can be used to link with the Effects table
- runmode "Batch" or "Interactive"
- r. squared R-squared, the fraction of variance explained by the model
- sigma Square root of the estimated variance of the random error
- stratavar Stratum variable(s)
- strata Stratum level(s)
- term Variable in the model
- wald.pvalue P-value from the Wald test of the given variable. Note that this test may be a multi-df test if the variable is categorical.

Details

Missing values will appear if a model did not converge, produced an error, or not run because of too many missing values in the outcome.

options

options list

Description

A list of 12:

- check.cor.cutoff Cutoff value to remove highly correlated columns in the design matrix. The default value is 0.97.
- check.cor.method Correlation method to remove highly correlated columns in the design matrix. It must be one of "spearman", "pearson", "kendall". The default value is "spearman".
- check.design TRUE or FALSE to check the design matrix for linearly dependent columns, highly correlated columns, or for being ill-conditioned whenever it is updated. The default is TRUE.
- check.illCond TRUE or FALSE to check for an ill-conditioned design matrix. The default is TRUE.
- check.nsubjects Minimum number of subjects. The default is 25.
- max.nstrata The maximum number of strata for a stratified analysis. The default is 10.
- model String for the model function. Currently, it must be one of "correlation", "lm", or "glm". The default is "correlation".

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• model.options List of options specific for the model. See correlation.options, glm.options, and lm.options for options specific to model="correlation", "lm", "glm" respectively. The default is NULL.

- output.ci_alpha Confidence interval level for estimated from glm models. This option must be a number >= 0 and < 1, where 0 is for not creating confidence intervals. The default value is 0.95.
- output.Effects A string to define the terms output in the returned Effects and ModelSummary data frames. Currently, it must be "exposure" or "all". If set to "all", then summary statistics for the exposure and adjustment variables will be output. Otherwise, only summary statistics for the exposure will be output. This option is ignored with model = "correlation". The default is "exposure".
- output.exp_parms TRUE, FALSE or NULL to exponentiate glm parameter estimates. Standard errors are obtained from the delta method. The default is NULL, so that estimates from glm models with family="binomial" will be exponentiated, and not otherwise.
- output.metab.cols Character vector of column names in the METABOLITES sheet to be output in the ModelSummary and Effects data frames. Metabolite ids are matched first using the outcomespec column and then using the exposurespec column. The default is "metabolite name".
- output. ModelSummary A string to defines the columns output in the returned ModelSummary data frame. Currently, it must be "anova" or "all". This option is ignored with model = "correlation". The default is "anova".

Details

Before any analysis is performed, an initial design matrix is constructed using the above options as follows:

- 1. Adjustment variables with less than two distinct non-missing values, or with less than check nsubjects non-missing values are removed.
- 2. The design matrix is created from the remaining adjustment variables and any linearly dependent columns are removed.
- 3. If check.cor.cutoff > 0, then highly correlated columns are removed by computing the correlation matrix cor_matrix = cor(design_matrix, method=check.cor.method).
- 4. If check.illCond is TRUE, then the function subselect::trim.matrix is applied to the correlation matrix to determine if additional columns will be removed.

If check.design is TRUE, then steps 2-4 are repeated each time an exposure variable is added to the design matrix or when rows of the design matrix are removed due to missing values in an outcome variable.

Examples

```
# Logistic regression with all default options
model.op <- list(family="binomial")
op <- list(model="glm", model.options=model.op)
# Compute Pearson correlations requiring at least 100 subjects
model.op <- list(method="pearson")
op <- list(model="correlation", check.nsubjects=100, model.options=model.op)</pre>
```

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OutputCSVResults Create output CSV file

Description

Create output CSV file

Usage

```
OutputCSVResults(filename, dataf, cohort = "")
```

Arguments

filename name of CSV file and can include path dataf correlation output (from function runCorr())

cohort cohort name

Value

the filename of the CSV file with results named with cohort

Examples

```
## Not run:
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,exposures="age",modlabel="1 Gender adjusted",
outcomes=c("lactose","lactate"))
corrmatrix <-runCorr(modeldata,exmetabdata,"DPP")
# Get correlation results
OutputCSVResults(filename="corr",dataf=corrmatrix,cohort="DPP")
# Get harmonization results
OutputCSVResults(filename="harmonization",dataf=exmetabdata$metab,cohort="DPP")
## End(Not run)</pre>
```

OutputListToExcel

Create an excel xlsx file from a list of data frames

Description

Create an excel xlsx file from a list of data frames

Usage

```
OutputListToExcel(filename, obj)
```

Arguments

filename Name of file and can include path. It must have a ".xlsx" extension.

obj List of data frames or matrices

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OutputXLSResults

Create output XLSX file

Description

Create output XLSX file

Usage

```
OutputXLSResults(filename, datal, cohort = "")
```

Arguments

filename name of file and can include path

data list to output (each item on list outputs to a worksheet)

cohort cohort name

Value

the filename of the XLSX file with results named with cohort

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,exposures="age",modlabel="1 Gender adjusted",
outcomes=c("lactose","lactate"))
# Get descriptive data
descdata <-runDescrip(exmetabdata)
OutputXLSResults(filename="corr",datal=descdata,cohort="DPP")</pre>
```

plotMinvalues

Plot the distribution of the number of missing values for each metabo-

Description

Plot the distribution of the number of missing values for each metabolite

```
plotMinvalues(
  cometsdata,
  title = "Distribution of the Number/Missing Values",
  xlabel = "Number of minimum/missing values",
  ylabel = "Frequency",
  xylabelsize = 12,
  titlesize = 16
```

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Arguments

cometsdata output of readCOMETSinput function

title main title for the plot (default is "Distribution of the Number/Missing Values")

xlabel x-axis label (default is "Number of minimum/missing values")

ylabel y-axis label (default is "Frequency")

xylabelsize size of x and y labels (default=8)

titlesize size of title (default, 20)

Value

a distribution plot

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
plotMinvalues(exmetabdata)</pre>
```

plotVar

Plot the variance distribution of transformed metabolite abundances

Description

Plot the variance distribution of transformed metabolite abundances

Usage

```
plotVar(
  cometsdata,
  title = "Distribution of Variance",
  titlesize = 16,
  xlabel = "Variance of transformed metabolite abundances",
  ylabel = "Frequency",
  xylabelsize = 12
)
```

Arguments

cometsdata output of readCOMETSinput function

title main title for the plot (default is "Distribution of Variance")

titlesize size of title (default, 20)

xlabel x-axis label (default is "Variance of transformed metabolite abundances")

ylabel y-axis label (default is "Frequency")

xylabelsize size of x and y labels (default=8)

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Value

a distribution plot

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
plotVar(exmetabdata)</pre>
```

RcometsAnalytics

RcometsAnalytics R package

Description

This R package supports all cohort-specific analyses of the COMETS consortium https://www.comets-analytics.org/. Data are not saved in the system but output must be downloaded and submitted for meta-analyses. import only functions needed

Details

Functions for analysis:

runCorr (correlation analysis)
runModel (correlation, glm or lm)
runAllModels (run models in batch mode from models sheet)

Functions for graphics:

plotVar (metabolite variance distribution plot)
plotMinvalues (distribution of missing values)
showHeatmap (heat map of metabolite correlations)
showHClust (interactive heat map with hierarchical clustering)

Functions for saving results to files:

OutputCSVResults (write to .csv file)
OutputXLSResults (write to excel file)

OutputListToExcel (write list of data frames to excel file with multiple sheets)

 ${\tt readCOMETSinput}$

Read in Excel file that contains metabolite data, covariate data, models, and model options.

Description

Read in Excel file that contains metabolite data, covariate data, models, and model options.

```
readCOMETSinput(file)
```

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Arguments

file

path of Excel file to be read in. This file must contain sheets with names **Metabolites**, **SubjectMetabolites**, **SubjectData**, **VarMap**, and optionally **Models**, **ModelOptions** (see details).

Details

Additional information regarding each sheet in the input Excel file is given below.

Metabolites

A table with the columns METABID, METABOLITE_NAME, and possibly other columns of information about the metabolites. The METABID column is used for harmonizing the metabolite names across different cohorts when meta-analyses are performed.

SubjectMetabolites

A table with the subject ids in the first column and metabolites as the other columns.

SubjectData

A table with the subject ids in the first column and covariates as the other columns.

VarMap

A table with at least the required columns VARREFERENCE, COHORTVARIABLE, VARTYPE, and VARDEFINITION. The COHORTVARIABLE column must contain names that match the column names in the **SubjectData** table. These names will be renamed to their corresponding name in the VARREFERENCE column. The VARTYPE column should have values continuous or categorical for each row.

Models

A table where each row represents a model to be run, and with columns MODEL, OUTCOMES, EXPOSURE, ADJUSTMENT, STRATIFICATION, WHERE, and optionally MODELSPEC. All variable names in this table must match variable names in the VARREFERENCE column of the VarMap sheet. The MODEL column is a label for the model. The OUTCOMES and EXPOSURE columns define the outcome and exposure variables for the model. Use All metabolites to specify that all metabolite variables are to be included as outcomes or exposures, otherwise use a space separated list of variable names. The ADJUSTMENT column contains a space separated list of covariate adjustment variables; use an empty cell for no covariate adjustment. The STRATIFICATION column is used for stratified analyses, with a space separated list of stratification variables. If more than one stratification variable is specified, then the strata are defined by all unique combinations of the stratification variables that appear in the data. The WHERE column is used to define a subset of subjects to include in the analysis, and has the form variable operator value, where operator can be one of the following >, <, >=, <= !=,=. An example WHERE condition is age > 50, which will include all subjects older than 50 in the analysis. Multiple WHERE conditions must be separated by a &. For example, age > 50 & bmi >= 22 will include the subjects older than 50 AND with bmi >= 22. Values in the MODELSPEC column must match with the MODELSPEC column in the ModelOptions sheet. This sheet is not required when running in interactive mode, but is required when running in batch mode.

ModelOptions

A table where each row specifies an option and has columns MODELSPEC, FUNCTION, OPTION, and VALUE. For an example sheet and additional information about this sheet, see the Excel file /extdata/cometsInput.xlsx. This sheet is optional, but is required when the **Models** sheet contains the column MODELSPEC.

Value

a list comprising of data and information needed for getModelData.

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)</pre>
```

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```
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)</pre>
```

runAllModels

This function allows users to run all models that are provided in the "Models" sheet of the input Excel file.

Description

This function allows users to run all models that are provided in the "Models" sheet of the input Excel file.

Usage

```
runAllModels(readData, cohortLabel = "", writeTofile = T)
```

Arguments

readData list from readCOMETSinput

cohortLabel cohort label (e.g. DPP, NCI, Shanghai)

writeTofile T/F (whether or not to write results for each model into separate xlsx files). Files

are written to current directory. Default is TRUE.

Value

A list of return objects from runModel or runCorr. The ith element in this list is the output from the ith model run.

Examples

```
## Not run:
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
allmodeloutput <- runAllModels(exmetabdata)
## End(Not run)</pre>
```

runCorr

Calculate correlations for input model.

Description

Calculate correlations for input model.

```
runCorr(modeldata, metabdata, cohort = "")
```

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Arguments

modeldata list from function getModelData

metabdata metabolite data list from readCOMETSinput cohort cohort label (e.g DPP, NCI, Shanghai)

Details

This function is a special case of runModel with the option op\$model = "correlation", however for backwards compatibility, it returns a data frame as in the original version of the **COMETS** R package.

Value

data frame with each row representing the correlation for each combination of outcomes and exposures represented as specified in the model (*spec), label (*lab), and universal id (*_uid) with additional columns for n, pvalue, method of model specification (Interactive or Batch), universal id for outcomes (outcome_uid) and exposures (exposure_uid) name of the cohort, adjustment (adjvars) and stratification (stratavar,strata) variables. Attribute of dataframe includes ptime for processing time of model run.

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,exposures="age",modlabel="1 Gender adjusted",
outcomes=c("lactose","lactate"), modelspec="Interactive")
corrmatrix <- runCorr(modeldata,exmetabdata, "DPP")</pre>
```

runDescrip

This function provides a description of the input data (for categorical data, the number of samples of each type; for continous data, the median and other statistics for each variable)

Description

This function provides a description of the input data (for categorical data, the number of samples of each type; for continous data, the median and other statistics for each variable)

Usage

```
runDescrip(readData)
```

Arguments

readData list from readComets

Value

a list with 2 data frames, continuous and categorical summaries. Type of variable is defined in varmap

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Examples

```
## Not run:
dir <- system.file("extdata", package="COMETS", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
allmodeloutput <- runAllModels(exmetabdata)
# Get descriptive data
descdata <-runDescrip(exmetabdata)
OutputXLSResults(filename="corr",datal=descdata,cohort="DPP")
## End(Not run)</pre>
```

runModel

Main function for running the models

Description

Main function for running the models

Usage

```
runModel(modeldata, metabdata, cohortLabel = "", op = NULL)
```

Arguments

modeldata list from function getModelData

metabdata metabolite data list from readCOMETSinput cohortLabel cohort label (e.g DPP, NCI, Shanghai)

op list of options when running in Interactive mode (see options).

Details

This function will check for stratification variables, and if present run within each stratum. The design matrix is checked for validity (see options). When running in Batch mode, the options are obtained from the Options sheet in the excel file.

Value

A list of objects with names ModelSummary, Effects, and Errors_Warnings.

Important: check the adjvars.removed column in the ModelSummary data frame to see if any adjustment variables were dropped from the model, and use caution interpreting results when variables are removed. The Errors_Warnings object may also contain additional variables removed. Attribute of this list includes ptime for processing time of model run.

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,exposures="age",modlabel="1 Gender adjusted",
outcomes=c("lactose","lactate"))
obj <- runModel(modeldata,exmetabdata, cohortLabel="DPP")</pre>
```

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showCorr

Function that returns top N lines of the runCorr output

Description

Function that returns top N lines of the runCorr output

Usage

```
showCorr(corr, nlines = 50)
```

Arguments

corr returned object from runCorr
nlines number of lines to return (default 50)

Value

first 50 lines of output

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,modlabel="1 Gender adjusted")
corrmatrix <- runCorr(modeldata,exmetabdata,"DPP")
showCorr(corrmatrix)</pre>
```

showHClust

Show interactive heatmap using heatmaply_cor with hierarchical clustering

Description

This function outputs a heatmap with hierarchical clustering. It thus requires you to have at least 2 outcome and 2 exposure variables in your models.

```
showHClust(
  ccorrList,
  strata = NULL,
  clust = TRUE,
  colscale = "RdYlBu",
  showticklabels = TRUE
)
```

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Arguments

ccorrList	correlation object (output of runCorr)
strata	Only valid if ccorrList is from a stratified analysis. If NULL, then results from the first stratum will be used in the plot.
clust	TRUE or FALSE to show hierarchical clustering. The default is TRUE.
colscale	$colors cale, can \ be \ custom \ or \ named \ ("Hots", "Greens", "Blues", "Greys", "Purples") \ see \ RColorBrewer_colors$
showticklabels	TRUE or FALSE to show axis labels. The default is TRUE.

Value

a heatmap with outcomes as rows and exposures in columns.

References

For colorscale reference: RColorBrewer_colors

Examples

```
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata, modelspec="Interactive",
exposures=c("age","bmi_grp"))
corrmatrix <-runCorr(modeldata,exmetabdata,"DPP")
showHClust(corrmatrix)</pre>
```

showHeatmap

Show interactive heatmap using plot_ly

Description

Show interactive heatmap using plot_ly

```
showHeatmap(
  ccorrList,
  strata = NULL,
  rowsortby = "corr",
  plothgt = 700,
  plotwid = 800,
  colscale = "RdYlBu"
```

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Arguments

ccorrList correlation object (output of runCorr)

strata Only valid if ccorrList is from a stratified analysis. If NULL, then results from the first stratum will be used in the plot.

rowsortby How row labels are sorted

plothgt Plot height default 700

plotwid Plot width default 800

colscale colorscale, can be custom or named ("Hots", "Greens", "Blues", "Greys", "Purples")

see https://plot.ly/ipython-notebooks/color-scales/

Value

a heatmap with outcomes as rows and exposures in columns.

References

For colorscale reference: https://plot.ly/ipython-notebooks/color-scales/

Examples

```
## Not run:
dir <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata <- getModelData(exmetabdata,modlabel="1 Gender adjusted")
corrmatrix <-runCorr(modeldata,exmetabdata,"DPP")
showHeatmap(corrmatrix)
## End(Not run)</pre>
```

showModel

Function that displays the first N rows of each data frame in the the runModel output.

Description

Function that displays the first N rows of each data frame in the the runModel output.

Usage

```
showModel(obj, nlines = 10)
```

Arguments

obj returned object from runModel

nlines number of lines to display (default 10)

20 showModel

Examples

```
dir      <- system.file("extdata", package="RcometsAnalytics", mustWork=TRUE)
csvfile      <- file.path(dir, "cometsInputAge.xlsx")
exmetabdata <- readCOMETSinput(csvfile)
modeldata      <- getModelData(exmetabdata,modlabel="1 Gender adjusted")
result      <- runModel(modeldata,exmetabdata, cohortLabel="DPP")
showModel(result)</pre>
```

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