

MeTime R package

September 12, 2022

add_col_stats

Function to check normality and add data to col data

Description

A method applied on the s4 object of class "metime_analyser" to check normality of the metabolites and add it to corresponding columns

Usage

```
add_col_stats(object, which_data, type, metab_names)
```

Arguments

| | |
|-------------|---|
| object | An object of class metime_analyser |
| which_data | dataset on which the method is to be applied |
| type | type of test, "shapiro" and "kruskal" are available |
| metab_names | column that has the metabolite names in col_data. |
| all | logical to add all kinds of available stats. |

Value

S4 object with shapiro wilk test related data in the col_data

Examples

```
object <- add_col_normality(object=data, which_data=c("lipid_data", "nmr_data"), type="shapiro", metab_names=c("m
```

```
add_distribution_vars_to_rows
```

Function to add measurements taken at screening time for samples to be added to all timepoints in row data

Description

A method applied on the s4 object of class "metime_analyser" to add all those datapoints that were measured only during screening to all the respective samples at all timepoints in row_data lists

Usage

```
add_distribution_vars_to_rows(
  object,
  screening_vars,
  distribution_vars,
  which_data
)
```

Arguments

| | |
|------------|---|
| object | An object of class metime_analyser |
| which_data | dataset to which the information is to be added(only 1 can be used at a time) |
| vars | A character naming the vars of interest |

Value

object of class metime_analyser with phenotype data added to row data

Examples

```
# adding APOEGrp, PTGENDER, and diag group to all data points and prepping the object for viz_distribution_plotter()
object <- add_distribution_vars_to_rows(object=data, screening_vars=c("APOEGrp", "DXGrp_longi", "PTGENDER"),
distribution_vars=c("Age", "BMI", "ADNI_MEM", "ADNI_LAN", "ADNI_EF", "APOEGrp", "DXGrp_longi", "PTGENDER"), which_data="screening")
```

```
add_metabs_as_covariates
```

Function to add metabolites as covariates for network construction

Description

Method applied on metime_analyser object to add other metabolite data to a certain dataset

Usage

```
add_metabs_as_covariates(object, which_data, which_metabs)
```

Arguments

| | |
|--------------|---|
| object | A S4 object of class metime_analyser |
| which_data | Dataset to which the metab data is to be added(please note that this a single character) |
| which_metabs | list of names of metabs and name of the list represents the dataset from which the metabs are to be acquired. eg: which_metabs=list(nmr_data=c("metab1", "metab2"), lipid_data=c("")) |

Value

S4 object with metabs added for GGM to another dataset

add_phenotypes_as_covariates

Function to add covariates to the dataset of interest for GGMs

Description

adds Covariates to data matrices in metime_analyser S4 object

Usage

```
add_phenotypes_as_covariates(
  object,
  which_data,
  covariates,
  class.ind,
  phenotype
)
```

Arguments

| | |
|------------|---|
| object | object of class metime_analyser |
| which_data | Dataset to which the covariates is to be added |
| covariates | character vector names of covariates. |
| class.ind | Logical to convert factor variables into class.ind style or not |
| phenotype | Logical. If True will extract from phenotype dataset else uses row data |

Value

S4 object with covariates added to the dataset

| | |
|--------------------|---|
| add_screening_vars | <i>Function to add measurements taken at screening time for samples to be added to all timepoints</i> |
|--------------------|---|

Description

A method applied on the s4 object of class "metime_analyser" to add all those datapoints that were measured only during screening to all the respective samples at all timepoints

Usage

```
add_screening_vars(object, vars)
```

Arguments

| | |
|--------|---|
| object | An object of class metime_analyser |
| vars | A character naming the vars of interest |

Value

phenotype data which can be replaced into the original object or use it separately with a different object

Examples

```
# adding APOEGrp, PTGENDER to all data points
new_with_apoegrp_sex <- add_screening_vars(object=metime_analyser_object, vars=c("APOEGrp", "PTGENDER"))
```

| | |
|------------------------------|--|
| calc_conservation_metabolite | <i>Function to calculate metabolite conservation index</i> |
|------------------------------|--|

Description

Method applied on the object metime_analyser to calculate the metabotype conservation index

Usage

```
calc_conservation_metabolite(object, which_data, timepoints, verbose)
```

Arguments

| | |
|------------|---|
| object | An object of class metime_analyser |
| which_data | Name of the dataset to be used |
| timepoints | character vector with timepoints of interest |
| verbose | Information provided on steps being processed |

Value

List of conservation index results

Examples

```
#calculating metabolite_conservation_index  
out <- calc_metabolite_conservation(object=metime_analyser_object, which_data="Name of the dataset")
```

calc_conservation_metabotype

Function to calculate metabotype conservation index

Description

Method applied on the object metime_analyser to calculate the metabotype conservation index

Usage

```
calc_conservation_metabotype(object, which_data, timepoints, verbose)
```

Arguments

| | |
|------------|---|
| object | An object of class metime_analyser |
| which_data | Name of the dataset to be used |
| timepoints | character vector with timepoints of interest |
| verbose | Information provided on steps being processed |

Value

List of conservation index results

Examples

```
#calculating metabotype_conservation_index  
out <- calc_metabotype_conservation(object=metime_analyser_object, which_data="Name of the dataset")
```

calc_correlation_pairwise

Function to calculate correlation

Description

calculate pairwise correlations This function creates a dataframe for plotting from a dataset.

Usage

```
calc_correlation_pairwise(object, which_data, method)
```

Arguments

| | |
|------------|---|
| object | S4 Object of class metime_analyser |
| which_data | specify datasets to calculate on. One or more possible |
| method | default setting: method="pearson", Alternative "spearman" also possible |

Value

data.frame with pairwise results

Examples

```
# Example to calculate correlations
dist <- calc_correlation(object=metime_analyser_object, which_data="name of the dataset",
  method="pearson")
```

calc_dimensionality_reduction

Function to calculate dimensionality reduction methods such as tsne, umap and pca.

Description

A method to apply on s4 object of class metime_analyse in order to obtain information after dimensionality reduction on a dataset/s

Usage

```
calc_dimensionality_reduction(object, which_data, type, ...)
```

Arguments

| | |
|------------|--|
| object | An object of class metime_analyser |
| which_data | a character vector - Names of the dataset from which the samples will be extracted |
| type | type of the dimensionality reduction method to be applied. Accepted inputs are "UMAP", "tSNE", "PCA" |
| ... | additional arguments that can be passed on to prcomp(), M3C::tsne() and umap::umap() |

Value

a list with two dataframes containing the dimensionality reduction information 1) samples - data of the individuals(".\$samples") 2) metabs - data of the metabolites(".\$metabs")

Examples

```
#calculate PCA
pca <- calc_dimensionality_reduction(object=metime_analyser_object, which_data="name/s of the dataset/s", type="PCA")
#calculate UMAP
pca <- calc_dimensionality_reduction(object=metime_analyser_object, which_data="name/s of the dataset/s", type="UMAP")
#calculate tSNE
pca <- calc_dimensionality_reduction(object=metime_analyser_object, which_data="name/s of the dataset/s", type="tSNE")
```

calc_distance_pairwise

Function to calculate dissimilarity using distance measures

Description

calculate pairwise distances This function creates a dataframe for plotting from a dataset.

Usage

```
calc_distance_pairwise(object, which_data, method)
```

Arguments

| | |
|------------|---|
| object | S4 Object of class metime_analyser |
| which_data | specify datasets to calculate on. One or more possible |
| method | default setting: method="euclidean", Alternative "maximum", "minimum", "manhattan", "canberra", "minkowski" are also possible |

Value

data.frame with pairwise results

Examples

```
# Example to calculate pairwise distances
dist <- calc_pairwise_distance(object=metime_analyser_object, which_data="name of the dataset",
                              method="euclidean")
```

calc_featureselection_boruta

Function to calculate dependent variables

Description

An S4 method to be applied on the metime_analyser object so as to calculate dependent variables

Usage

```
calc_featureselection_boruta(
  object,
  which_x,
  which_y,
  verbose,
  output_loc,
  file_name
)
```

Arguments

| | |
|------------|---|
| object | An object of class metime_analyser |
| which_x | Name of the dataset to be used for training |
| which_y | Name of the dataset to be used for testing |
| verbose | Information provided on steps being processed |
| output_loc | path to the parent directory where in the out file will be stored |
| file_name | name of the out file |

Value

List of conservation index results

 calc_ggm_genenet_crosssectional

An automated fuction to calculate GGM from genenet crosssectional version

Description

automated funtion that can be applied on metime_analyser object to obtain geneNet network along with threshold used

Usage

```
calc_ggm_genenet_crosssectional(
  object,
  which_data,
  threshold,
  timepoint,
  all,
  ...
)
```

Arguments

| | |
|------------|---|
| object | S4 object of class metime_analyser |
| which_data | a character or a character vector naming the datasets of interest |
| threshold | type of threshold to be used for extracting significant edges. allowed inputs are "li", "FDR", "bonferroni" |
| all | Logical to extract all edges without any pval correction |
| ... | additional arguments for GeneNet |
| timepoints | timepoints of interest that are to be used to build networks(as per timepoints in rows) |

Value

Network data with edgelist, partial correlation values and associated p-values and corrected p-values

 calc_ggm_genenet_longitudnal

An automated fuction to calculate GGM from genenet longitudinal version

Description

automated funtion that can be applied on metime_analyser object to obtain geneNet network along with threshold used

Usage

```
calc_ggm_genenet_longitudnal(
  object,
  which_data,
  threshold,
  timepoints,
  all,
  ...
)
```

Arguments

| | |
|------------|---|
| object | S4 object of class metime_analyser |
| which_data | a character or a character vector naming the datasets of interest |
| threshold | type of threshold to be used for extracting significant edges |
| timepoints | timepoints of interest that are to be used to build networks(as per timepoints in rows) |
| all | Logical to get all edges without any cutoff. |
| ... | additional arguments for genenet network |

Value

Network data with edgelist, partial correlation values and associated p-values and corrected p-values

calc_ggm_multibipartite_lasso

An automated fuction to calculate GGM from multibipartite lasso approach

Description

automated funtion that can be applied on s4 object of class metime_analyser to calculate a network using multibipartite lasso

Usage

```
calc_ggm_multibipartite_lasso(object, which_data, alpha, nfolds, timepoints)
```

Arguments

| | |
|------------|---|
| object | S4 object of class metime_analyser |
| which_data | a character or a character vector naming the datasets of interest |
| alpha | tuning parameter for lasso + ridge regression in glmnet |
| nfolds | nfolds for cv.glmnet |
| timepoints | timepoints of interest that are to be used to build networks(as per timepoints in rows) |

Value

Network data with edges and their respective betas

| | |
|--------------|---|
| calc_parafac | <i>Function to perform PARAFAC analysis</i> |
|--------------|---|

Description

Method to be applied on S4 object of class metime_analyser to perform PARAFAC analysis

Usage

```
calc_parafac(object, which_data, timepoints, nfac = 3, ...)
```

Arguments

| | |
|------------|--|
| object | S4 object of class metime_analyser |
| which_data | character vector for dataset to be used |
| timepoints | character vector to define timepoints of interest |
| nfac | parameter nfac for parafac(). Numeric value to define the number of factors. Default is set to 3 |
| ... | Additional arguments to be used for the function parafac() |

Value

An object of class PARAFAC. See multiway library for more information

| | |
|-------------------|--|
| calc_temporal_ggm | <i>An automated function to calculate temporal network with lagged model</i> |
|-------------------|--|

Description

calculates temporal networks for each dataset with a lagged model as used in graphical VAR

Usage

```
calc_temporal_ggm(object, which_data, lag, timepoints, alpha, nfolds)
```

Arguments

| | |
|------------|---|
| object | S4 object of class metab_analyser |
| which_data | dataset or datasets to be used |
| lag | which lagged model to use. 1 means one-lagged model, similiary 2,3,..etc |
| timepoints | timepoints of interest that are to be used to build networks(in the order of measurement) |
| alpha | parameter for regression coefficient |
| nfolds | nfolds parameter for glmnet style of regression |

Value

temporal network data with edgelist and regression values

calc_ttest

Function to calculate students t-test

Description

Method for S4 object of class metime_analyser for performing t-test

Usage

```
calc_ttest(object, which_data, timepoints, split_var)
```

Arguments

| | |
|------------|---|
| object | S4 object of class metime_analyser |
| which_data | dataset or datasets to be used for the analysis |
| timepoints | two timepoints of interest to perform the test on |
| split_var | split variable for testing such as diagnostic group etc |

Value

t-test result as a list or a list of t-test results

check_col_normality *Function to check for col_normality data whether it is added or not.*

Description

function to check whether col_normality data is added to the object or not

Usage

```
check_col_normality(object, which_data)
```

Arguments

| | |
|------------|---------------------------------------|
| object | S4 object of class of metime_analyser |
| which_data | dataset/s to check |

Value

NULL if it passes all the sanity checks

check_ids_and_classes *Function to check the ids in the data and data format*

Description

sanity check to check for ids and order of the data

Usage

```
check_ids_and_classes(object)
```

Arguments

| | |
|--------|---------------------------------------|
| object | S4 object of class of metime_analyser |
|--------|---------------------------------------|

Value

NULL if it passes all the sanity checks

check_rownames_and_colnames

Function to check the format of rownames and colnames and if they are same or not

Description

sanity check to check for rownames of the data

Usage

check_rownames_and_colnames(object)

Arguments

object S4 object of class of metime_analyser

Value

NULL if it passes all the sanity checks

comp_network_with_established

Function to compare the network generated from data to an existing network

Description

function to perform fishers exact test to decide which network is the best.

Usage

comp_network_with_established(calc_networks, est_network)

Arguments

calc_networks list of networks calculated from data
 est_network established network to compare with the established network results Make sure that this network has only two columns with names as node1 and node2

Value

fisher test results with pval and test statistic

`get_append_analyser_object`

This function appends an object of class metime_analyser with a new dataset.

Description

function to apply on metime_analyse object to append a new dataset into the existing object

Usage

```
get_append_analyser_object(object, data, col_data, row_data, name)
```

Arguments

| | |
|----------|---|
| object | S4 object of class metime_analyser |
| data | data.frame containing data |
| col_data | data.frame containing col_data: id column of col data has to match colnames of data |
| row_data | data.frame containing row_data: id column of row data has to match rownames of data |
| name | Name of the new dataset |

Value

An object of class metime_analyser

Examples

```
# append data frames into the metime_analyser object
appended_object <- get_append_metab_object(object=metime_analyser_object, data=data, row_data=data, col_data=col_data)
```

`get_betas_for_multibipartite_lasso`

Function to perform multibipartite style regression on a list of matrices

Description

Performs multibipartite lasso in cv.glmnet style on a list of matrices that have metabolite information from different platforms

Usage

```
get_betas_for_multibipartite_lasso(list_of_mats, alpha, nfolds)
```

Arguments

| | |
|--------------|---|
| list_of_mats | a list with matrices and samples ordered similarly |
| alpha | alpha for cv.glmnet regression. Defines style of penalty. |
| nfolds | nfolds for cv.glmnet |

Value

returns a list with information of the combinations in context

get_class_info_from_edges

Function to get information on how many class edges are present

Description

Function to check how the different edges in a GGM are associated to their respective classes(it could be super-pathway or sub-pathway)

Usage

```
get_class_info_from_edges(calc_networks, metadata)
```

Arguments

| | |
|---------------|-------------------------------|
| calc_networks | list of calculated networks |
| metadata | metadata of the edges present |

Value

table with information on different type of edges present

| | |
|---------------------|--|
| get_files_and_names | <i>Function to pack all the data into a single object of class "metime_analyser"</i> |
|---------------------|--|

Description

This function loads all the files from the parent directory. It assumes a certain naming pattern as follows: "datatype_Noncollrow_data.rds" Any other naming pattern is not allowed. The function first writes all files into a list and each type of data is packed into its respective class i.e. col_data, row_data or data

Usage

```
get_files_and_names(path, annotations_index)
```


Arguments

path Path to the parent directory

annotations_index a list to be filled as follows = list(phenotype="Name or index of the files", medication="Name or index of the files")

Value

An object of class metime_analyser

Examples

```
# Input in the parent directory from which the data files are to be extracted along with annotations_index to specify
get_files_and_names(path=/path/to/parent/directory, annotations_index=list(phenotype="Name of phenotype file", m
```

| | |
|-----------------|--|
| get_ggm_genenet | <i>Function to calculate a dynamic GeneNet GGM from a longitudinal data matrix</i> |
|-----------------|--|

Description

calculates GGM on longitudinal data matrix and returns a dataframe with edges, partial correlation and associated p-values

Usage

```
get_ggm_genenet(data, threshold = c("bonferroni", "FDR", "li"), all, ...)
```

Arguments

data data matrix in a longitudinal format

threshold type of multiple hypothesis correction. Available are Bonferoni("bonferroni"), Benjamini-Hochberg("FDR") and independent tests method("li", also see Li et al)

all Logical to get all edges without any cutoff.

... additional arguments for ggm.estimate.pcor()

Value

a dataframe with edges, partial correlation and associated p-values

```
get_make_analyser_object
```

Function to pack all the data into a single object of class "metime_analyser"

Description

This function creates an object of class metime_analyser from a dataset.

Usage

```
get_make_analyser_object(
  data,
  col_data,
  row_data,
  annotations_index,
  name = NULL
)
```

Arguments

| | |
|-------------------|---|
| data | data.frame containing data |
| col_data | data.frame containing col_data: id column of col data has to match colnames of data |
| row_data | data.frame containing row_data: id column of row data has to match rownames of data |
| annotations_index | a list to be filled as follows = list(phenotype="Name or index of the file/list", medication="Name or index of the files/list") |
| name | character. Name you want to assign to the new dataset that is being added on |

Value

An object of class metime_analyser

Examples

```
# new_metime_analyser_object <- get_make_metab_object(data=data_frame, col_data=col_data_frame, row_data=row_data,
  annotations_index=list(phenotype="name of phenotype", medication="name of medication"))
```

`get_make_plotter_object`*Function to make a plottable object for viz functions*

Description

function to generate metime_plotter object from plot data and metadata

Usage

```
get_make_plotter_object(data, metadata, calc_type, calc_info, plot_type, style)
```

Arguments

| | |
|------------------------|--|
| <code>data</code> | dataframe of plotable data obtained from any calc object |
| <code>metadata</code> | dataframe with the metadata for the plot table mentioned above. To obtain these see <code>get_metadata_for_rows()</code> and <code>get_metadata_for_columns()</code> |
| <code>calc_type</code> | A character to specify type of calculation - will be used for <code>comp_</code> functions For networks the accepted notations are "genenet_ggm", "multibipartite_ggm", and "temporal_network" |
| <code>calc_info</code> | A string to define the information about calculation |
| <code>plot_type</code> | type of the plot you want to build. eg: "box", "dot" etc. Its a character vector |
| <code>style</code> | Style of plot, accepted inputs are "ggplot", "circo" and "visNetwork". Is a singular option. |

`get_metadata_for_columns`*Get metadata for columns(in most cases for metabolites)*

Description

function to generate a metadata list for building the MeTime plotter object

Usage

```
get_metadata_for_columns(object, which_data, columns, names, index_of_names)
```

Arguments

| | |
|-----------------------------|---|
| <code>object</code> | S4 object of class MeTime Analyser |
| <code>which_data</code> | Names of dataset/s to be used |
| <code>columns</code> | A list of character vectors for the columns of interest. Length of the list should be same as length of <code>which_data</code> |
| <code>names</code> | A Character vector with the new names for the columns mentioned above |
| <code>index_of_names</code> | character vector to define the name of the column in which names of the variables are stored |

Value

data.frame with metadata information

get_metadata_for_rows *Get metadata for rows(in most cases for samples)*

Description

function to generate a metadata list for building the MeTime plotter object

Usage

```
get_metadata_for_rows(object, which_data, columns)
```

Arguments

| | |
|------------|--|
| object | S4 object of class MeTime Analyser |
| which_data | Names of dataset/s to be used |
| columns | A list of character vectors for the columns of interest. Length of the list should be same as length of which_data |

Value

data.frame with metadata information for rows

get_palette *Get a palette of "n" distinct colorblind friendly colors*

Description

Function to get a palette of distinct colorblind friendly colors, the distinctiveness is determined by the difference in their hue values.

Usage

```
get_palette(n)
```

Arguments

| | |
|---|--|
| n | number of colors wanted in the palette |
|---|--|

Value

a color palette vector with colors in the form of hex codes

Examples

```
# colors=get_palette(n=10)
```

`get_samples_and_timepoints`

Function to know the number of timepoints and the total number of samples available at that point

Description

A method applied onto s4 object of class "metime_analyser" so as to obtain the number of unique samples available at each timepoint.

Usage

```
get_samples_and_timepoints(object, which_data)
```

Arguments

| | |
|-------------------------|---|
| <code>object</code> | An object of class <code>metime_analyser</code> |
| <code>which_data</code> | Name of the dataset in context |

Value

A data table with timepoints and number of samples at each timepoint

Examples

```
# newdata <- get_samples_and_timepoints(object=metime_analyser_object, which_data="Name of dataset of interest")
```

`get_text_for_plot`

Function to Obtain textual information for visualization in interactive plots

Description

a standard function to be applied on data matrices or dataframes with the colnames of interest such that the information from columns is visualized in the interactive plot

Usage

```
get_text_for_plot(data, colnames)
```

Arguments

| | |
|-----------------------|---|
| <code>data</code> | a dataframe with plotting data along with other variables for visualization |
| <code>colnames</code> | a character vector with the names of the variables that you want to see on the plot |

Value

a vector with strings that can be parsed into plot_ly text.

Examples

```
# text = get_text(data=data.frame, colnames=c("names", "of", "columns", "of", "interest"))
```

metime_analyser-class *Constructor to generate an object of class metime_analyser. contains slots - list_of_data: For the list of all data matrices. - list_of_col_data: list of all the col data files in the same order. - list_of_row_data: list of all the row data files in the same order. - annotations: list with phenotype and medication. Each of which is character that represents the name of the aforementioned dataset types.*

Description

Constructor to generate an object of class metime_analyser. contains slots - list_of_data: For the list of all data matrices. - list_of_col_data: list of all the col data files in the same order. - list_of_row_data: list of all the row data files in the same order. - annotations: list with phenotype and medication. Each of which is character that represents the name of the aforementioned dataset types.

Constructor to generate an object of class metime_analyser. contains slots - list_of_data: For the list of all data matrices. - list_of_col_data: list of all the col data files in the same order. - list_of_row_data: list of all the row data files in the same order. - annotations: list with phenotype and medication. Each of which is character that represents the name of the aforementioned dataset types.

Constructor to generate an object of class metime_analyser. contains slots - list_of_data: For the list of all data matrices. - list_of_col_data: list of all the col data files in the same order. - list_of_row_data: list of all the row data files in the same order. - annotations: list with phenotype and medication. Each of which is character that represents the name of the aforementioned dataset types.

Constructor to generate an object of class metime_analyser. contains slots - list_of_data: For the list of all data matrices. - list_of_col_data: list of all the col data files in the same order. - list_of_row_data: list of all the row data files in the same order. - annotations: list with phenotype and medication. Each of which is character that represents the name of the aforementioned dataset types.

Constructor to generate an object of class metime_analyser. contains slots - list_of_data: For the list of all data matrices. - list_of_col_data: list of all the col data files in the same order. - list_of_row_data: list of all the row data files in the same order. - annotations: list with phenotype and medication. Each of which is character that represents the name of the aforementioned dataset types.

`metime_plotter-class` *creating metime_plotter class that converts calculations and meta-data as a plotable object to parse into viz_plotter Contains slots - plot_data: list of Dataframe(s) with plotting data and metadata for visualization. Dataframes is an option only for visNetwork() plots. Need a list of two dataframes: Nodes dataframe and edge dataframe named as .\$.node and .\$.edge - plot: ggplot(), circos() or visNetwork() object - calc_type: A vector to specify type of calculation - will be used for comp_functions - calc_info: string to define the information about calculation - plot_type: A character vector to define the type of plots that are needed. - style: Character that defines the style of plot i.e. a ggplot(), circos() or visNetwork() plot. Is always a singular input. Cannot have two styles in one object.*

Description

creating metime_plotter class that converts calculations and metadata as a plotable object to parse into viz_plotter Contains slots - plot_data: list of Dataframe(s) with plotting data and metadata for visualization. Dataframes is an option only for visNetwork() plots. Need a list of two dataframes: Nodes dataframe and edge dataframe named as .\$.node and .\$.edge - plot: ggplot(), circos() or visNetwork() object - calc_type: A vector to specify type of calculation - will be used for comp_functions - calc_info: string to define the information about calculation - plot_type: A character vector to define the type of plots that are needed. - style: Character that defines the style of plot i.e. a ggplot(), circos() or visNetwork() plot. Is always a singular input. Cannot have two styles in one object.

creating metime_plotter class that converts calculations and metadata as a plotable object to parse into viz_plotter Contains slots - plot_data: Dataframe with plotting data and metadata for visualization - plot: ggplot(), circos() or visNetwork() object with predefined aesthetics - calc_type: A vector to specify type of calculation - will be used for comp_functions - calc_info: string to define the information about calculation - plot_type: A character vector to define the type of plots that are needed.

`mod_convert_s4_to_s3` *Function to Convert S4 object of class metime_analyser to an S3 object with same architecture*

Description

converter function to be applied onto metime_analyse object to convert into a standard list of S3 type.

Usage

```
mod_convert_s4_to_s3(object)
```

Arguments

object An object of class metime_analyser

Value

An S3 object of the same data as metime_analyser in other words all slots are now converted into nested lists

Examples

```
# convert S4 object to a list
s3_list <- mod_convert_s4_to_s3(object=metime_analyser_object)
```

```
mod_extract_common_samples
```

Function to get only common samples from the dataframes in list_of_data

Description

A method applied on object of class metime_analyse to extract common samples across datasets. Also has an option to split the data according timepoints(mod_split_acc_time()).

Usage

```
mod_extract_common_samples(object, time_splitter = FALSE)
```

Arguments

object An object of class metime_analyser

time_splitter A boolean input: True leads to splitting of the data wrt time, False returns all the dataframes as they are with common rows

Value

list_of_data with common samples across all time points

Examples

```
# extracting common samples across all datasets
new_list_of_data <- mod_common_sample_extractor(object=metime_analyser_object)
```

| | |
|---------------|--|
| mod_filter_tp | <i>Functions for selecting time points</i> |
|---------------|--|

Description

a method applied onto class metime_analyser in order to extract timepoints of interest from a dataset

Usage

```
mod_filter_tp(object, timepoints, full, which_data)
```

Arguments

| | |
|------------|--|
| object | An object of class metime_analyser |
| timepoints | time points to be selected |
| full | if TRUE subjects are only selected if measured in all selected time points |
| which_data | Name of the dataset to be used |

Value

An object of class metime_analyser with processed data

Examples

```
#example to use this function
object <- mod_filter_tp(object, timepoints=c(0,12,24), full=TRUE, which_data="Name of the dataset")
```

| | |
|----------------------------|--|
| mod_merge_metime_analysers | <i>Function to merge one or more metime_analyser objects</i> |
|----------------------------|--|

Description

function to merge multiple metime_analyser objects

Usage

```
mod_merge_metime_analysers(list_of_objects, annotations_index)
```

Arguments

| | |
|-------------------|---|
| list_of_objects | list of metime analyser objects that are to be merged |
| annotations_index | new list with annotations_index. Can also set to be NULL. |

Value

A merged metime_analyser object

| | |
|----------------|---|
| mod_remove_nas | <i>Function to remove NA's from data matrices</i> |
|----------------|---|

Description

A method applied on S4 object to remove NA's and change data accordingly

Usage

```
mod_remove_nas(object, which_data)
```

Arguments

| | |
|------------|---|
| object | S4 object of class metime_analyser |
| which_data | dataset/s for which the method is to be applied |

Value

S4 object with NA's removed and data manipulated accordingly

| | |
|-----------------------|---|
| mod_split_acc_to_time | <i>Function to split data according to time</i> |
|-----------------------|---|

Description

Function to split the list of dataframes into a nested list with each dataframe being split into into dataframes of different timepoints

Usage

```
mod_split_acc_to_time(object)
```

Arguments

| | |
|--------|------------------------------------|
| object | An object of class metime_analyser |
|--------|------------------------------------|

Value

list_of_data with each dataframe being broken into a list of dataframes with respect to the timepoint they belong to

Examples

```
#splitting data according to time
new_data <- mod_split_acc_to_time(object=metime_analyser_object)
```

| | |
|---------------|---|
| mod_trans_log | <i>Function to apply log transformation</i> |
|---------------|---|

Description

Function to log transform data

Usage

```
mod_trans_log(object, which_data, base)
```

Arguments

| | |
|------------|------------------------------------|
| object | An object of class metime_analyser |
| which_data | Name of the dataset to be used |
| base | base of log to be used |

Value

An object of class metime_analyser with processed data

Examples

```
# example to apply log transformation
object <- mod_logtrans(object, which_data="name of the dataset", base=2)
```

| | |
|------------------|-----------------------------------|
| mod_trans_zscore | <i>Function to scale the data</i> |
|------------------|-----------------------------------|

Description

Functions for scaling

Usage

```
mod_trans_zscore(object, which_data)
```

Arguments

| | |
|------------|------------------------------------|
| object | An object of class metime_analyser |
| which_data | Name of the dataset to be used |

Value

An object of class metime_analyser with processed data

Examples

```
# example to apply scaling
object <- mod_zscore(object, which_data="name of the dataset")
```

save_analyser_object *Function to extract analyser object data into a csv*

Description

extracts information from analyser object and saves it as a csv

Usage

```
save_analyser_object(object, which_data)
```

Arguments

| | |
|------------|-----------------------------------|
| object | An object of class metime_plotter |
| which_data | Character to specify the dataset |

Value

saves the data in the working directory as a csv and returns nothing

Examples

```
see examples here
save_analyser_object(object, which_data="dataset")
```

save_plot_from_plotter
 Function to save interactive plots

Description

extracts plot from plotter object and saves it as a widget

Usage

```
save_plot_from_plotter(object, out)
```

Arguments

| | |
|--------|--|
| object | An object of class metime_plotter |
| out | Character to specify path of the output file to save the widget in |

Value

saves the plot and returns nothing

Examples

```
save_plot_from_plotter(object)
```

| | |
|---------------------|---|
| save_plotter_object | <i>Function to extract plot data into a csv</i> |
|---------------------|---|

Description

extracts information from plotter object and saves it as a csv

Usage

```
save_plotter_object(object, out)
```

Arguments

| | |
|--------|--|
| object | An object of class metime_plotter |
| out | Character to specify path of the output file or character vector in case of visNetwork |

Value

saves the data into a csv and returns nothing

Examples

```
see examples here
Network : save_plotter_object(object, out=c("edge.csv", "node.csv", "meta.csv"))
Others : save_plotter_object(object, out="outfile.csv")
```

| | |
|--------------------|----------------------------------|
| set_parallel_cores | <i>register parallel backend</i> |
|--------------------|----------------------------------|

Description

function to run in order to perform the analysis parallely thereby saving time

Usage

```
set_parallel_cores(n_cores = NULL)
```

Arguments

n_cores A number of specified cores.

Value

set a parallel backend

show,metime_analyser-method

Setting new print definition for the metime_analyser object

Description

function to see the structure of metime_analyser object

Usage

```
## S4 method for signature 'metime_analyser'  
show(object)
```

Arguments

object S4 object of class metime_analyser

Value

structure of the S4 object

Examples

```
structure(object)
```

show,metime_plotter-method

Setting new print definition for the metime_plotter object

Description

function to see the structure of metime_plotter object

Usage

```
## S4 method for signature 'metime_plotter'  
show(object)
```

Arguments

object S4 object of class metime_plotter

Value

structure of the S4 object

Examples

```
structure(object)
```

structure,metime_plotter-method

Setting new structure definition for the metime_plotter object

Description

function to see the structure of metime_plotter object

Usage

```
## S4 method for signature 'metime_plotter'  
structure(object)
```

Arguments

object S4 object of class metime_plotter

Value

structure of the S4 object

Examples

```
structure(object)
```

| | |
|-----------|--|
| structure | <i>Setting new structure definition for the metime_analyser object</i> |
|-----------|--|

Description

function to see the structure of metime_analyser object

Usage

```
structure(object)
```

Arguments

| | |
|--------|------------------------------------|
| object | S4 object of class metime_analyser |
|--------|------------------------------------|

Value

structure of the S4 object

Examples

```
structure(object)
```

| | |
|--------------------------|--|
| viz_distribution_plotter | <i>Function for Plotting distributions of phenotypic variables</i> |
|--------------------------|--|

Description

A method to be applied onto s4 object so as to obtain distributions of various phenotypic variables

Usage

```
viz_distribution_plotter(object, colname, which_data, strats, phenotype)
```

Arguments

| | |
|------------|---|
| object | An object of class metime_analyser |
| colname | Name of the variable whose distribution is of interest |
| which_data | Name of the dataset from which the samples will be extracted |
| strats | Character vector with colnames that are to be used for stratification |
| phenotype | Logical. If true data will be collected from phenotype_data matrix else from row data |

Value

a list with either 1) density plot, mean table acc to timepoint and variable type or 2) bar plot, line plot, and variable type

Examples

```
# extracting distribuion of Age from dataset1
plot <- viz_distribution_plotter(object, colname="Age", which_data="dataset1", strats="additional columns for fac
```

| | |
|--------------------|--|
| viz_plotter_circos | <i>Setting up standard wrapper for all circos plots for a metime_plotter object.</i> |
|--------------------|--|

Description

plot function for metime_plotter object with different inputs to specialize plots. Used for all calc outputs.

Usage

```
viz_plotter_circos(object, aesthetics, outfile, layout_by)
```

Arguments

| | |
|------------|--|
| object | S4 object of class metime_plotter |
| aesthetics | list for aesthetics. eg: list(list(x="colname",y="colname",color="colname", shape="colname"), list(...)) for "dot" plot and "heatmap" plot, for heatmap: list(x="colname", y="colname", fill="colname"). Additionally two other character vectors are allowed namely .\$.vis and .\$.strats for text and for facet wrapping. |

| | |
|--------------------|--|
| viz_plotter_ggplot | <i>Setting up standard wrapper for all ggplot plots for a metime_plotter object.</i> |
|--------------------|--|

Description

plot function for metime_plotter object with different inputs to specialize plots. Used for all calc outputs.

Usage

```
viz_plotter_ggplot(object, aesthetics, interactive)
```

Arguments

| | |
|-------------|---|
| object | S4 object of class metime_plotter |
| aesthetics | list for aesthetics. eg: list(list(x="colname",y="colname",color="colname", shape="colname"), list(...)) for "dot" plot and "heatmap" plot, for heatmap: list(x="colname", y="colname", fill="colname"). Additionally two other character vectors are allowed namely .viz and .sstrats for text and for facet wrapping. |
| interactive | Flag option(Logical). If set to TRUE will generate an interactive plot else will generate a normal ggplot |

Value

metime_plotter object with updated plot

viz_plotter_visNetwork

Setting up standard wrapper for network plots from visNetwork for a metime_plotter object.

Description

plot function for metime_plotter object with different inputs to specialize plots. Used for all calc outputs.

Usage

```
viz_plotter_visNetwork(object, title, layout_by)
```

Arguments

| | |
|-----------|--|
| object | S4 object of class metime_plotter |
| title | character/string that is the title of the graph output |
| layout_by | character to define the layout style to be used |

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