Package 'oetteR'

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VignetteBuilder knitr

R topics documented:

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|--|

Description

Takes a data_ls object generated by f_clean_data() and adds boxcox transformations of all numeric variables.

Usage

```
f_boxcox(data_ls)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, numericals = < vector with column names of numerical columns>)

Details

For a boxcox transformation all values mut be > 0. The function will automatically add the abs(min(x)) + 0.0001 to all columns if they contain values <= 0

Value

returns a list

data the cleaned dataframe as tibble
categoricals vector of column names containing categorical data
numericals vector of column names containing numerical data
ids vector of column names containing ids
boxcox_names vector of column names containing boxcox transformed variables
boxcox_data tibble containing boxcox transformed variables

See Also

```
f_clean_data
```

```
data_ls = f_clean_data(mtcars)
f_manip_get_most_common_level( data_ls$data$cyl)
```

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Description

Performs a number of cleaning operations on a dataframe, detects numerical and categorical columns and returns a list containing the cleaned dataframe and vectors naming the columns with a specific data type.

Usage

```
f_clean_data(data, max_number_of_levels_factors = 10,
    min_number_of_levels_nums = 6, exclude_missing = T,
    replace_neg_values_with_zero = T, allow_neg_values = c("null"),
    id_cols = c("null"))
```

Arguments

```
data a dataframe
max_number_of_levels_factors

If a factor variable contains more then the maximum number of levels the levels
with the lowest frequency will be collapsed into 'others', Default: 10
min_number_of_levels_nums

If a numeric number contains less that the minimum of distinct values it will be
converted to a factor, Default: 6
exclude_missing

exclude observations with missing values, Default: T
replace_neg_values_with_zero
all negative values will be set to 0, Default: T
allow_neg_values
```

Details

id_cols

The list this function returns can be a bit tedious to work with. If you want to engineer a new feature you have to manually update the categoricals or the numericals vector. I suggest that you do all the feature engineering before applying this function. The advantage of this column is that when you get to the modelling or visualisation steps you have full control over which columns are used for the formula or for the type of visualisation even if you might have bloated your dataframe with some junk columns.

specify columns for which negative values are allowed, Default: c("null")

Value

```
returns a list

data the cleaned dataframe as tibble
categoricals vector of column names containing categorical data
categoricals_ordered
vector of column names containing all ordered categorical data
numericals vector of column names containing numerical data
ids vector of column names containing ids
```

specify columns containing ids.

See Also

```
f_boxcox
```

Examples

```
data_ls = f_clean_data( mtcars , id_cols = 'names')
str(data_ls)
```

```
f_clean_data_no_changes
```

wrapper for f_clean_data without modifications to data

Usage

```
f_clean_data_no_changes(data)
```

Arguments

data a dataframe

Value

returns a list

data the cleaned dataframe as tibble

categoricals vector of column names containing categorical data

categoricals_ordered

vector of column names containing all ordered categorical data

numericals vector of column names containing numerical data

ids vector of column names containing ids

See Also

```
f_clean_data
```

f_datatable_universal convert dataframe to DT:datatable inlcuding most usefull extensions and options

Description

inludes the features for excel and clipboard export, hide and unhide (column visibility), reorder columns per drag and drop, navigate table with arrow keys and prefix/suffix directed rounding of numerical values.

- count_/ _count will round to 0
- p_val will round to Default: 2
- perc_/ _perc will round to Default: 1

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Usage

```
f_datatable_universal(df, round_perc = 1, round_sign = 2,
    count_cols_as_int = T, round_other_nums = NULL)
```

Arguments

df dataframe

round_perc digits for percentages, Default: 1 round_sign digits for p_values, Default: 2

count_cols_as_int

detect count columns, Default: T

round_other_nums

round other numerical columns to that digit. Will not do anything if NULL,

Default: NULL

Value

DT:datatable

See Also

 $\verb|str_detect| data table, format Percentage, format Signif, format Round \\$

Examples

```
data_ls = f_clean_data(mtcars)
f_stat_group_counts_percentages(data_ls, 'cyl') %>%
    f_datatable_universal()

f_stat_group_mean_medians(data_ls, 'cyl') %>%
    f_datatable_universal(round_other_nums = 2)
```

f_html_breaks

create a taglist with n lines of html line breaks

Usage

```
f_html_breaks(n)
```

Arguments

n number of line breaks

Value

taglist

See Also

br,tagList

f_html_filename_2_link

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Examples

```
f_html_breaks(5)
```

```
f_html_filename_2_link
```

convert a filename + path or a file_path to a html link

Usage

```
f_html_filename_2_link(file_name = dir()[1], path = getwd(),
  file_path = NULL, link_text = file_name)
```

Arguments

```
file_name character vector, Default: dir()[1]
path character vector, Default: getwd()
file_path file_path( path, file_name)
link_text character vector
```

Value

link

See Also

```
str_replace_all
```

Examples

```
dir()[1]
f_html_filename_2_link()
dir()[1:5]
f_html_filename_2_link(dir()[1:5])
```

f_html_padding

add some padding around html objects

Usage

```
f_html_padding(obj, pad_before = 0, title = NULL, subtitle = NULL,
   caption = NULL, pad_after = 0)
```

Arguments

```
obj html object such as DT:datatable() or plotly::ggplotly()
pad_before integer, Default: 0
title character vector, Default: "
subtitle character vector, Default: "
caption character vector, Default: "
pad_after character vector, Default: 0
```

taglist

See Also

```
tagList,h3,h4,h6
```

Examples

```
f_html_padding(DT::datatable(mtcars),5,'mtcars Data','subtitle', 'caption', 8 )
```

```
f_manip_append_2_list append object to list
```

Description

convenience function to replace l[[length(1)+1]] = x

Usage

```
f_manip_append_2_list(1, x)
```

Arguments

```
listobject
```

Value

list

Examples

```
l = list('a', 'b')
l = f_manip_append_2_list(1, 'c')
str(l)
```

```
f_manip_bin_numerics bin numerical columns
```

Description

centers, scales and Yeo Johnson transforms numeric variables in a dataframe before binning into n bins of eqal range. Outliers based on boxplot stats are capped (set to min or max of boxplot stats).

Usage

```
f_manip_bin_numerics(df, bins = 5, bin_labels = c("LL", "ML", "M", "MH",
   "HH"), center = T, scale = T, transform = T)
```

Arguments

df dataframe with numeric variables

bins number of bins for numerical variables, Default: 5

bin_labels labels for the bins from low to high, Default: c("LL", "ML", "M", "MH",

"HH")#' @param center boolean, Default: T

scale boolean, Default: T transform boolean, Default: T

Value

dataframe

Examples

```
## Not run:
if(interactive()){
  #EXAMPLE1
  }
## End(Not run)
```

```
f_manip_bring_to_pos_range
```

bring vector to positice range

Description

if min < 0, add abs(min) to all values

Usage

```
f_manip_bring_to_pos_range(vec)
```

Arguments

vec numeric vector

Value

vector

```
vec = c( -2,0,2,4,6)
vec = f_manip_bring_to_pos_range( vec )
vec
```

model.matrix() creates dummy variables for factors. The names of these dummy variables however are not compatible with the formula syntax. This wrapper cleans up the names of the new variables.

Usage

```
f_manip_data_2_model_matrix_format(data, formula, scale_data = T,
   center_data = T, exclude_na_columns = T)
```

Arguments

```
data a dataframe
formula formula
scale_data boolean
center_data boolean
exclude_na_columns
boolean
```

Value

list with new dataframe and new formula

See Also

```
str_replace_all
```

```
data_ls = f_clean_data(mtcars)
data = data_ls$data
formula = hp ~ disp + am + gear
data_trans = f_manip_data_2_model_matrix_format( data, formula )
response_var =f_manip_get_response_variable_from_formula(data_trans$formula)
vars = f_manip_get_variables_from_formula(data_trans$formula)
x = as.matrix( select( data_trans$data, one_of(vars) ) )
y = data_trans$data[[response_var]]
glmnet::glmnet( x , y )
```

```
f_manip_factor_2_numeric
```

converts factor to numeric preserving numeric levels and order in character levels

Usage

```
f_manip_factor_2_numeric(vec)
```

Arguments

vec

vector

Value

vector

See Also

```
str_detect
```

Examples

```
fac_num = factor( c(1,3,8) )
fac_chr = factor( c('foo','bar') )
fac_chr_ordered = factor( c('a','b','c'), ordered = T )

f_manip_factor_2_numeric( fac_num )
f_manip_factor_2_numeric( fac_chr )
f_manip_factor_2_numeric( fac_chr_ordered )
```

Usage

```
f_manip_get_most_common_level(x)
```

factor vector

Arguments

X

Value

character vector

See Also

tidy

Examples

```
data_ls = f_clean_data(mtcars)
f_manip_get_most_common_level( data_ls$data$cyl)
```

```
\label{lem:constraint} f\_manip\_get\_response\_variable\_from\_formula \\ \textit{get response variable from formula}
```

Usage

```
f_manip_get_response_variable_from_formula(formula)
```

Arguments

formula formula

Value

character vector

See Also

```
f_manip_get_variables_from_formula
```

```
\begin{tabular}{ll} f\_manip\_get\_variables\_from\_formula \\ & get\ variables\ from\ formula \\ \end{tabular}
```

Usage

```
f_manip_get_variables_from_formula(formula)
```

Arguments

formula formula

Value

character vector

See Also

```
f_manip_get_response_variable_from_formula
```

```
f = foo~bar1 + bar2
vars = f_manip_get_variables_from_formula(f)
response_var = f_manip_get_response_variable_from_formula(f)
```

```
f_manip_matrix_2_tibble
```

converts matrices to tibble, preserving row.names

Description

row.names are added as row_names column as the first column of the tibble. Function does not fail when object cannot be converted to tibble thus can be used to map over lists with various variable types such as modells and objects.

Usage

```
f_manip_matrix_2_tibble(x)
```

Arguments

Χ

any variable

Value

a tibble or if the input variable is neither matrix dataframe or tibble the original input object.

Examples

```
mat = as.matrix(mtcars)
head( mat, 10)
f_manip_matrix_2_tibble( mat )

# convert all matrices from a list
pca = prcomp( mtcars ) %>%
   map( f_manip_matrix_2_tibble )
pca
```

```
f_manip_summarize_2_median_and_most_common_factor
```

takes a data_ls list created by f_clean_data() and returns a list with all medians for numerical and most common level for categorical variables.

Usage

```
f_manip_summarize_2_median_and_most_common_factor(data_ls)
```

Arguments

```
data_ls object generated by f_clean_data(), or a named list list( data = <dataframe>, numericals = < vector with column names of numerical columns>)
```

list

data summarized data as dataframe

data_boxcox summarized boxcox data as dataframe

Examples

```
summarized_ls = f_clean_data(mtcars) %>%
  f_boxcox() %>%
  f_manip_summarize_2_median_and_most_common_factor()
summarized_ls$data
summarized_ls$boxcox_data
```

```
f_manip_transpose_tibble
```

transpose a tibble

Description

transpose a tibble, values in first column will become column titles. Row names will be converted to first columns

Usage

```
f_manip_transpose_tibble(tib)
```

Arguments

tib

tibble

Value

tibble

```
tib = mtcars %>%
  as_tibble() %>%
  f_manip_transpose_tibble()
tib
```

```
f_model_add_predictions_2_grid_regression
                         add predictions to grid (regression models)
```

wrapper for modelr::add_predictions

Usage

```
f_model_add_predictions_2_grid_regression(grid, m, var)
```

Arguments

m

grid grid containing all variables used for the model model

character vector denoting response variable var

Value

grid

See Also

```
add_predictions
```

Examples

```
data_ls = f_clean_data(mtcars)
formula = disp~hp+mpg
m = lm(formula, data_ls$data)
grid = f_model_data_grid(data_ls, formula, 'hp', 10) %>%
  f_model_add_predictions_2_grid_regression( m, 'disp')
```

f_model_data_grid

generates a data grid based on a formula

Description

the range of one specified variable is expanded, while all other variables are set to the most common values. Similar to modelr::data_grid but it can deal with factors.

Usage

```
f_model_data_grid(col_var, data_ls, formula, n = 500, set_manual = list())
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>,

numericals = < vector with column names of numerical columns>)

formula formula

n integer, length of grid, datapoints in between range of col_var

set_manual named list, set some variables manually instead of defaulting to median or most

common factor. !! Values need to be of the same variable type as in the original

data.

Value

dataframe

Examples

```
data_ls = f_clean_data(mtcars)
formula = disp~cyl+mpg+hp
f_model_data_grid( 'mpg', data_ls, formula, 10 )
f_model_data_grid( 'mpg', data_ls, formula, 10 , set_manual = list( cyl = min(data_ls$data$cyl) ) )
```

f_model_importance

model importance

Description

supports rpart, randomForest, svm, will return NULL for other models

Usage

```
f_model_importance(m, data)
```

Arguments

m model data training data

Value

tibble

```
pl = pipelearner::pipelearner(mtcars) %>%
    pipelearner::learn_models( twidlr::rpart, disp~. ) %>%
    pipelearner::learn_models( twidlr::randomForest, disp~. ) %>%
    pipelearner::learn_models( twidlr::svm, disp~. ) %>%
    pipelearner::learn() %>%
    mutate( imp = map2(fit, train, f_model_importance) )
pl$imp
```

optimised for usage in pipelearner dataframe

Usage

```
f_model_importance_plot(importance, title, variable_color_code = NULL)
```

Arguments

Value

plotly graph

```
data_ls = f_clean_data(mtcars)
variable_color_code = f_plot_color_code_variables(data_ls)
m = twidlr::rpart(mtcars, disp~.)
imp = f_model_importance_rpart(m)
f_model_importance_plot(imp
                        , title = 'rpart'
                          variable_color_code = variable_color_code
#pipelearner
pl = pipelearner::pipelearner(data_ls$data) %>%
  pipelearner::learn_models( twidlr::rpart, disp~. ) %>%
  pipelearner::learn_models( twidlr::randomForest, disp~. ) %>%
  pipelearner::learn_models( twidlr::svm, disp~. ) %>%
  pipelearner::learn() %>%
  mutate( imp = map2(fit, train, f_model_importance)
         , title = paste( model, models.id, cv_pairs.id, train_p )
         , plot = map2( imp
                       , title
                       , f_model_importance_plot
                       , variable_color_code = variable_color_code
```

```
)
htmltools::tagList(pl$plot)
```

takes the most important variables of a model and plots a tabplot::tableplot

Usage

```
f_model_importance_plot_tableplot(data, ranked_variables, response_var,
  limit = 10, print = F, ...)
```

Arguments

```
data dataframe
ranked_variables
datafram as returned by f_model_importance()
response_var character vector denoting response variable
limit integer limit the number of variables, Default: 10
... pass kwargs to tabplot::tableplot
```

Value

tabplot::tableplot object

See Also

tableplot

```
data = f_clean_data(mtcars) %>%
    .$data
m = rpart::rpart( disp~., data)
ranked_variables = f_model_importance(m, data)
response_var = 'disp'

f_model_importance_plot_tableplot( data, ranked_variables, response_var, limit = 5 )

#pipe
form = as.formula('disp~cyl+mpg+hp')
pl = pipelearner::pipelearner(mtcars) %>%
pipelearner::learn_models( twidlr::rpart, form ) %>%
pipelearner::learn_models( twidlr::randomForest, form ) %>%
pipelearner::learn_models( twidlr::svm, form ) %>%
pipelearner::learn_models( twidlr::svm, form ) %>%
pipelearner::learn_models( twidlr::svm, form ) %>%
```

adds a bar plot of the ranked variables, a tabplot sorted by the target variable and a dependency plot (response variable vs the sequential range of one of the predictor variables while all other predictors are kept constant at mean values).

Usage

```
f_model_importance_pl_add_plots_regression(pl, data, m, ranked_variables,
  response_var, title,
  variable_color_code = f_plot_color_code_variables(data_ls), formula,
  data_ls, var_dep_limit = 10, var_dep_log_y = F, tabplot_limit = 12,
  formula_in_pl = F)
```

Arguments

```
pl
                  a dataframe containing the columns for data, m, ranked_variables, response_var
data
                  symbol (unquoted name) of data column in pl
                  symbol (unquoted name) of data column in pl
ranked_variables
                  symbol (unquoted name) of data column in pl
                  symbol (unquoted name) of data column in pl
response_var
title
                  symbol (unquoted name) of data column in pl
variable_color_code
                  dataframe created by f_plot_color_code_variables()
formula
                  fomula that was used to construct model
data_ls
                  data_ls list object containing the whole of the original data
                  number of variables to be plotted on dependency plot
var_dep_limit
var_dep_log_y
                  should y axis of dependency plot be logarithmic
                  number of variables to be plotted on tabplot
tabplot_limit
formula_in_pl
                  boolean if formula is a column in pl?
```

dataframe

See Also

f_model_importance_plot f_model_importance_plot_tableplot f_model_plot_variable_dependency_regres

Examples

```
data_ls = f_clean_data(mtcars)
form = disp~cyl+mpg+hp
variable_color_code = f_plot_color_code_variables(data_ls)
pl = pipelearner::pipelearner(data_ls$data) %>%
  pipelearner::learn_models( twidlr::rpart, form ) %>%
  pipelearner::learn_models( twidlr::randomForest, form ) %>%
  pipelearner::learn_models( twidlr::svm, form ) %>%
  pipelearner::learn() %>%
  mutate( imp = map2(fit, train, f_model_importance)
          , title = paste(model, models.id, train_p) ) %>%
  f\_model\_importance\_pl\_add\_plots\_regression( \ data
                                                                      = train
                                                                      = fit
                                                , ranked_variables = imp
                                                , title
                                                                     = title
                                                , response_var = target
                                              , variable_color_code = variable_color_code
                                                , formula
                                                                     = form
                                                , data_ls
                                                                     = data_ls
                                                , var_dep_limit
, var_dep_log_y
                                                                     = 10
                                                                      = T
                                                , tabplot_limit
                                                                      = 12 )
```

Description

should execute f_model_importance_pl_add_plots_regression() on modelling dataframe first

Usage

```
f_model_importance_pl_plots_as_html(pl, prefix = NULL)
```

Arguments

```
pl modelling dataframe containing the following columns 'imp_plot', 'imp_plot_dep', 'imp_tabplot', 'title'

prefix character vector file name prefix for html files, Default: NULL
```

html files in working directory

See Also

```
tagList
```

Examples

```
## Not run:
   data_ls = f_clean_data(mtcars)
form = disp~cyl+mpg+hp
variable_color_code = f_plot_color_code_variables(data_ls)
pl = pipelearner::pipelearner(data_ls$data) %>%
pipelearner::learn_models( twidlr::rpart, form ) %>%
pipelearner::learn_models( twidlr::randomForest, form ) %>%
pipelearner::learn_models( twidlr::svm, form ) %>%
pipelearner::learn() %>%
mutate( imp = map2(fit, train, f_model_importance)
         , title = paste(model, models.id, train_p) ) %>%
f_model_importance_pl_add_plots_regression( data
                                                                  = train
                                                                  = fit
                                             , ranked_variables = imp
                                             , title
                                                                 = title
                                             , response_var
                                                                = target
                                            , variable_color_code = variable_color_code
                                             , formula
                                                                = form
                                             , data_ls
                                                                 = data_ls
                                                                = 10
                                             , var_dep_limit
                                             , var_dep_log_y
                                                                 = T
                                             , tabplot_limit
                                                                  = 12) %>%
f_model_importance_pl_plots_as_html( prefix = 'test_oetteR_html_')
files = dir() %>%
 .[ startsWith(., 'test_oetteR_html_') ]
file.remove( files )
## End(Not run)
```

```
{\tt f\_model\_importance\_randomForest}
```

extract variable importance for randomForest model

Usage

```
f_model_importance_randomForest(m, ...)
```

Arguments

model of class randomForest

dataframe

Examples

```
#regression
m = twidlr::randomForest(mtcars, disp~.)
f_model_importance_randomForest(m)

#classification
data_ls = f_clean_data(mtcars)
m = twidlr::randomForest(data_ls$data, cyl~.)
f_model_importance_randomForest(m)
```

Usage

```
f_model_importance_rpart(m, ...)
```

Arguments

m

model of class rpart

Value

dataframe

```
#regression
m = twidlr::rpart(mtcars, disp~.)
f_model_importance_rpart(m)

#classification
data_ls = f_clean_data(mtcars)
m = twidlr::rpart(data_ls$data, cyl~.)
f_model_importance_rpart(m)
```

```
f_model_importance_svm
```

extract variable importance for svm

Usage

```
f_model_importance_svm(m, data)
```

Arguments

m model of class sym

data original training dataframe

Details

uses 1D-SA 1 dimensional sensitivity analysis using rminer::Importance()

Value

dataframe

Examples

```
#regression
m = twidlr::svm(mtcars, disp~.)
f_model_importance_svm(m, mtcars)

#classification
data = mtcars
data$cyl = factor(data$cyl, ordered = T)
m = twidlr::svm(data, cyl~.)
f_model_importance_svm(m, data)
```

Description

response variable will be plotted against the entire range of each variable staring with the most important ones. All other variables will be set to median or most common factor. This function requires a ranked list of the most important variables as returned by f_model_importance()

Usage

```
f_model_plot_variable_dependency_regression(m, ranked_variables,
  title = unlist(stringr::str_split(class(m)[1], "\\."))[1], data = NULL,
  formula, data_ls,
  variable_color_code = f_plot_color_code_variables(data_ls), limit = 12,
  log_y = F, set_manual = list(), ...)
```

Arguments

```
a regression model
ranked_variables
                  datafram as returned by f_model_importance()
                  character vector as plot title, Default: unlist(stringr::str_split(class(m)[1], "\."))[1]
title
data
                  a dataframe, only necessary if it differs from data_ls$data, Default: NULL
formula
                   the formula used to train the model
data_ls
                  data_ls object generated by f_clean_data(), or a named list list( data = <dataframe>,
                  numericals = < vector with column names of numerical columns>)
variable_color_code
                  dataframe created by f_plot_color_code_variables()
                  integer limit the number of variables to be plotted, Default: 12
limit
set_manual
                  named list, set some variables manually instead of defaulting to median or most
                  common factor. !! Values need to be of the same variable type as in the original
```

arguments passed to facet_wrap e.g. usefull for nrow, ncol

plot

See Also

Value

```
str_split
```

```
# regular version-----
data_ls
                  = f_clean_data(mtcars)
data
                  = data_ls$data
formula
                  = disp~hp+mpg+cyl
                  = randomForest::randomForest(formula, data)
ranked_variables = f_model_importance( m, data)
variable_color_code = f_plot_color_code_variables(data_ls)
                  = 12
f\_model\_plot\_variable\_dependency\_regression(\ m
                                           , ranked_variables
                            , title = unlist( stringr::str_split( class(m)[1], '\\.') )[1]
                                          , formula = formula
                                           , data_ls = data_ls
                                          , variable_color_code = variable_color_code
                                          , limit = limit
```

```
data_ls = f_clean_data(mtcars)
form = as.formula('disp~hp+cyl+wt')
variable_color_code = f_plot_color_code_variables(data_ls)
limit
                 = 10
 pl = pipelearner::pipelearner( data_ls$data ) %>%
  pipelearner::learn_models( rpart::rpart, form ) %>%
  pipelearner::learn_models( randomForest::randomForest, form ) %>%
  pipelearner::learn_models( e1071::svm, form ) %>%
  pipelearner::learn() %>%
  mutate( imp
              = map2(fit, train, f_model_importance)
         ,plot = pmap( list( m = fit, ranked_variables = imp, title = model, data = train)
                        , .f = f_model_plot_variable_dependency_regression
                        , formula = form
                        , data_ls = data_ls
                         variable_color_code = variable_color_code
                       , limit = limit
         )
  )
```

```
f_model_plot_var_dep_over_spec_var_range

plot vmodel varaible dependency over the range of a specified variable
```

Some models are able to capture relative dependencies. In order to visualise them the dataset is split into three parts. 0-25,25-75,75-100 percentile or the three most common factors. Then variable dependencies for each of the three splits are plotted. In the mtcars example below we can see that the model predicts an increase in disp if drat increases for cars with 8 cylinders, while the opposite is true for cars with only 6 cylinders.

Usage

```
f_model_plot_var_dep_over_spec_var_range(m, title, variables, range_variable,
  data, formula, data_ls, variable_color_code, log_y = F, limit = 12)
```

Arguments

```
variable_color_code
dataframe created by f_plot_color_code_variables()

log_y boolean log_scale for y axis

limit integer limit the number of variables to be plotted, Default: 12

data_ls PARAM_DESCRIPTION
```

grid can be printed with gridExtra::grid.arrange()

See Also

arrangeGrob

```
## Not run:
 # single output example ------
           = randomForest::randomForest
.f
data_ls
                  = f_clean_data(mtcars)
data
                  = data_ls$data
               - uata_is$uata
= disp~mpg+cyl+am+hp+drat+qsec+vs+gear+carb
formula
                  = .f(formula, data)
variables = f_model_importance( m, data)
title = unlist( stringr::str_split( class(m)[1], '\\.') )[1]
variable_color_code = f_plot_color_code_variables(data_ls)
limit
        = 10
log_y
                   = F
range_variable_num = data_ls$numericals[1]
range_variable_cat = data_ls$categoricals[1]
grid_num = f_model_plot_var_dep_over_spec_var_range(m
                                                 , title
                                                  , variables
                                                  , range_variable_num
                                                  , data
                                                  , formula
                                                  , data_ls
                                                  , variable_color_code
                                                  , log_y
                                                  , limit )
gridExtra::grid.arrange(grid_num)
# pipe example ------
data_ls = f_clean_data(mtcars)
form = as.formula('disp~cyl+mpg+hp+am+gear+drat+wt+vs+carb')
variable_color_code = f_plot_color_code_variables(data_ls)
grids = pipelearner::pipelearner(data_ls$data) %>%
  pipelearner::learn_models( twidlr::rpart, form ) %>%
  pipelearner::learn_models( twidlr::randomForest, form ) %>%
  pipelearner::learn_models( twidlr::svm, form ) %>%
  pipelearner::learn() %>%
```

f_model_seq_range 27

```
dplyr::mutate( imp = map2(fit, train, f_model_importance)
                 , range_var = map_chr(imp, function(x) head(x,1)row_n
                 , grid = pmap( list( m = fit
                                      , title = model
                                      , variables = imp
                                       , range_variable = range_var
                                      , data = test
                 )
                 , f_model_plot_var_dep_over_spec_var_range
                 , formula = form
                 , data_ls = data_ls
                 , variable_color_code = variable_color_code
                 , log_y = F
                 , limit = 12
                 )
 ) %>%
  .$grid
f_plot_obj_2_html( grids, type = "grids", output_file = 'test_me', title = 'Grids', height = 30 )
file.remove('test_me.html')
## End(Not run)
```

f_model_seq_range

generates sequence of variable spanning from min to max

Description

similar to modelr::seq_range but can handle categorical variables

Usage

```
f_model_seq_range(data_ls, col_var, n = 500)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>,
numericals = < vector with column names of numerical columns>)

col_var character vector denoting variable

n integer number of intermediate data points, Default: 500

Value

vector

```
data_ls = f_clean_data(mtcars)
col_var = 'disp'
f_model_seq_range( data_ls, col_var, 10)
```

28 f_pca

f_pca

calculate principle components for a dataset

Description

This function is an extended wrapper for prcomp(). I takes a data_ls object created by f_clean_data and calculates the contribution of each variable to each principle component in percent.

Usage

```
f_pca(data_ls, center = T, scale = T, use_boxcox_tansformed_vars = T,
  include_ordered_categoricals = T, threshold_vae_for_pc_perc = 2.5)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>,

numericals = < vector with column names of numerical columns>)

center boolean, Default: T
scale boolean, Default: T
use_boxcox_tansformed_vars
boolean, Default: T

Details

Blog post explaining how to calculate contributions

Value

a list with the original data complemented with the principle component vector data of each observation and an object returned by prcomp() supplemented with some extra features

data dataframe

pca pca object created by prcomp()

added features of pca:

cos2 The squared rotation vectors.A value between 0 and 1 denotes the amount of

contribution of a variable to a specific principle component

vae percent variance explained

contrib_abs_perc

The absolute contribution of one variable to the variance explained by one principle component in percent. The total contibution adds up to the total contibution

of the principle componaent in percent.

contrib_abs_perc_reduced

as above but variables contibuting less than 2.5 percent are grouped

threshold_vae_for_pc_perc

principle components that explain less percent variance than this threshold are

dropped

See Also

```
prcomp
```

Examples

```
pca_ls = f_clean_data(mtcars) %>%
f_boxcox() %>%
f_pca()
```

f_pca_plot_components plot principle components as a dot plot

Usage

```
f_pca_plot_components(pca_ls, x_axis = "PC1", y_axis = "PC2",
    group = NULL)
```

Arguments

pca_ls	list created by f_pca()
x_axis	character vector, Default: 'PC1'
y_axis	character vector, Default: 'PC2'
group	character vector denoting the grouping variable, determines dot colour, Default: NULL

Value

htmltools taglist containing a plotly graph and tow DT datatables will only show if printed in a rmarkdown document

```
## Not run:
tagls = f_clean_data(mtcars) %>%
  f_boxcox() %>%
  f_pca() %>%
  f_pca_plot_components(group = 'cyl')
## End(Not run)
```

```
\label{lem:components} f\_pca\_plot\_variance\_explained \\ plot\ varaince\ explained\ of\ principle\ components
```

Usage

```
f_pca_plot_variance_explained(pca_ls, threshold_vae_for_pc_perc = 2.5)
```

Arguments

```
pca_ls list created by f_pca()
```

Value

plotly graph

Examples

```
p = f_clean_data(mtcars) %>%
  f_boxcox() %>%
  f_pca() %>%
  f_pca_plot_variance_explained()
p
```

```
f_plot_adjust_col_vector_length

adjust length of color vector, by repeating colors
```

Usage

```
f_plot_adjust_col_vector_length(n = 74, col_vector = f_plot_col_vector74())
```

Arguments

```
n length, Default: 74
col_vector vector containing colors, Default: f_plot_col_vector74()
```

Value

vector containing colors of specified length

```
length( f_plot_adjust_col_vector_length(100) )
```

f_plot_alluvial 31

|--|

Description

plots a dataframe as an alluvial plot. All numerical variables are scaled, centered and YeoJohnson transformed before binning.

Usage

```
f_plot_alluvial(data, variables = names(data), col_id = NULL,
  max_variables = 20, bins = 5, bin_labels = c("LL", "ML", "M", "MH",
  "HH"), order_levels = NULL, fill_by = "first_variable",
  col_vector_flow = f_plot_col_vector74(faint = F, greys = F),
  col_vector_value = RColorBrewer::brewer.pal(9, "Greys")[c(3, 6, 4, 7, 5)])
```

Arguments

data	a dataframe
variables	vector denoting names and order of the plotted variables, Default: names(data)
max_variables	maximum number of variables, Default: 20
bins	number of bins for numerical variables, Default: 5
bin_labels	labels for the bins from low to high, Default: c("LL", "ML", "M", "MH", "HH")
order_levels	character vector denoting levels to be reorderer from low to high
fill_by	$one_of(c('first_variable', 'last_variable', 'all_flows', 'values')), Default: 'first_variable'$
col_vector	vector with HEX color codes, Default: RColorBrewer::brewer.pal(name = "Dark2", n = 8)

Details

DETAILS

Value

OUTPUT_DESCRIPTION

See Also

```
brewer.pal fct_relevel geom_flow,geom_stratum
```

```
## Not run:
if(interactive()){

data_ls = mtcars %>%
  f_clean_data()

data = data_ls$data
max_variables = 5
```

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```
variables = c( data_ls$categoricals[1:3], data_ls$numericals[1:3] )
f_plot_alluvial( data = data
                , variables = variables
                , max_variables = max_variables
                , fill_by = 'first_variable' )
f_plot_alluvial( data = data
                , variables = variables
                , max_variables = max_variables
                , fill_by = 'last_variable' )
f_plot_alluvial( data = data
                , variables = variables
                , max_variables = max_variables
                , fill_by = 'all_flows' )
f_plot_alluvial( data = data
                , variables = variables
                , max_variables = max_variables
                , fill_by = 'first_variable' )
# manually order variable values
f_plot_alluvial( data = data
                 , variables = variables
                 , max_variables = max_variables
                 , fill_by = 'values'
                 , order_levels = c('1', '0') )
}
## End(Not run)
```

f_plot_alluvial_1v1 plot alluvial

Description

Plots two variables of a dataframe on an alluvial plot. A third variable can be added either two the left or the right of the alluvial plot to provide coloring of the flows. All numerical variables are scaled, centered and YeoJohnson transformed before binning.

Usage

```
f_plot_alluvial_1v1(data, col_x, col_y, col_id, col_fill = NULL,
  fill_right = T, bins = 5, bin_labels = c("LL", "ML", "M", "MH", "HH"),
  order_levels_y = NULL, order_levels_x = NULL, order_levels_fill = NULL,
  complete = TRUE, fill_by = "first_variable",
  col_vector_flow = f_plot_col_vector74(faint = F, greys = F),
  col_vector_value = RColorBrewer::brewer.pal(9, "Greys")[c(3, 6, 4, 7, 5)])
```

 $f_{plot_alluvial_1v1}$ 33

Arguments

```
data
                   a dataframe
col x
                   character vector denoting column for the x axis variable
col_y
                   character vector denoting column for the y axis variable
col_id
                   character vector denoting id column
col_fill
                   character vector denoting color fill variable for flows, Default: NULL
fill_right
                   logical, TRUE fill variable is added to the right FALSE to the left, Default: T
                   number of bins for automatic binning of numerical variables, Default: 5
bins
bin_labels
                   labesl for bins, Default: c("LL", "ML", "M", "MH", "HH")
order_levels_y
                   character vector denoting order of y levels from low to high, does not have to be
                   complete can also just be used to bring levels to the front, Default: NULL
order_levels_x character vector denoting order of x levels from low to high, does not have to be
                   complete can also just be used to bring levels to the front, Default: NULL
order_levels_fill
                   character vector denoting order of color fill variable levels from low to high,
                   does not have to be complete can also just be used to bring levels to the front,
                   Default: NULL
                   add implicitly missing data, Default: TRUE
complete
fill_by
                   one_of(c('first_variable', 'last_variable', 'all_flows', 'values')), Default: 'first_variable'
col_vector_flow
                   HEX colors for flows, Default: f_plot_col_vector74(faint = F, greys = F)
col_vector_value
                   Hex colors for y levels/values, Default: RColorBrewer::brewer.pal(9, "Greys")[c(3,
                   6, 4, 7, 5
```

Value

plot

See Also

```
brewer.pal fct_relevel,fct_rev UQ geom_flow,geom_stratum
```

```
group_by(tailnum, carrier, origin, dest, qu ) %>%
 summarise( mean_arr_delay = mean(arr_delay) ) %>%
 ungroup() %>%
 mutate( mean_arr_delay = ifelse( mean_arr_delay < 10, 'on_time', 'late' ) )</pre>
levels(monthly_flights$qu) = c('Q1', 'Q2', 'Q3', 'Q4')
data = monthly_flights
col_x = 'qu'
col_y = 'mean_arr_delay'
col_fill = 'carrier'
col_id = 'tailnum'
# flow coloring variants
f_plot_alluvial_1v1( data, col_x, col_y, col_id, col_fill )
f\_plot\_alluvial\_1v1(\ data,\ col\_x,\ col\_y,\ col\_id,\ fill\_by\ =\ 'last\_variable'\ )
f_plot_alluvial_1v1( data, col_x, col_y, col_id, fill_by = 'first_variable' )
f_plot_alluvial_1v1( data, col_x, col_y, col_id, fill_by = 'all_flows' )
f_plot_alluvial_1v1( data, col_x, col_y, col_id, fill_by = 'value' )
# use same color coding for flows and y levels
f_plot_alluvial_1v1( data, col_x, col_y, col_id, fill_by = 'last_variable'
                    , col_vector_flow = f_plot_col_vector74()
                    , col_vector_value = f_plot_col_vector74() )
# move fill variable to the left
f_plot_alluvial_1v1( data, col_x, col_y, col_id, col_fill, fill_right = F )
# reorder levels
f_plot_alluvial_1v1( data, col_x, col_y, col_id, fill_by = 'first_variable'
                    , order_levels_y = c('on_time', 'late') )
f_plot_alluvial_1v1( data, col_x, col_y, col_id, fill_by = 'first_variable'
                    , order_levels_x = c('Q4', 'Q3', 'Q2', 'Q1'))
order_by_carrier_size = data %>%
 group_by(carrier) %>%
 count() %>%
 arrange( desc(n) ) %>%
 .[['carrier']]
f_plot_alluvial_1v1( data, col_x, col_y, col_id, col_fill
                    , order_levels_fill = order_by_carrier_size )
}
## End(Not run)
```

f_plot_color_code_variables

color code all variables in a data_ls list.

f_plot_col_vector74 35

Description

color coding is stable the same data_ls list gets the same coding with every function call. Assigns the same colors to the boxcox transformed and untransformed variant of a variable.

Usage

```
f_plot_color_code_variables(data_ls, col_vector = f_plot_col_vector74())
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, numericals = < vector with column names of numerical columns>)

col_vector character vector denoting Hexcode colors, Default: f_plot_col_vector74()

Value

tibble

- variable
- colorHEX code color

See Also

```
str_replace_all
```

Examples

```
f_clean_data(mtcars) %>%
  f_boxcox() %>%
  f_plot_color_code_variables() %>%
  print()
```

f_plot_col_vector74 generate a most distinctive color scale

Description

based on RColorBrewer colours of length 74 for RGB colors see rapidtables(https://www.rapidtables.com/web/color/inde

Usage

```
f_plot_col_vector74(greys = T, reds = T, blues = T, greens = T,
  faint = T, only_unique = F)
```

Arguments

colors boolean include color

Value

vector with HEX colours

36 f_plot_hist

```
f\_plot\_generate\_comparison\_pairs\\ generates\ comparison\ pairs\ for\ `ggpubr::stat\_compare\_means()`
```

Description

generates all possible pairs and filters according to t-test p_value

Usage

```
f_plot_generate_comparison_pairs(data, col_var, col_group, thresh = 0.05)
```

Arguments

data	dataframe
------	-----------

col_var character vector denoting variable column
col_group character vector denoting grouping column

thresh double, Default: 0.05

Value

list

See Also

```
str_splitUQ
```

Examples

```
f_plot_generate_comparison_pairs( mtcars, 'disp', 'cyl' )
```

Description

Function plots smart histograms for variables in a data_ls list generated by f_clean_data(). It supports three types of histograms: Bar histograms, density histograms and violin plots. We can further specify a categorical variable to group on. The function defaults to a sensible standard output if key word arguments are not applicable for variable type. Thus we can easily pipe through long lists of variables and thus generate histograms for all variables in the input (see examples).

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Usage

```
f_plot_hist(variable, data_ls, group = "None", graph_type = "violin",
  y_axis = "count", auto_range = T, n_breaks = 30, rug = T, x_min = 0,
  x_max = 100, title = "",
  col_vector = f_plot_adjust_col_vector_length(100,
  RColorBrewer::brewer.pal(name = "Dark2", n = 8)), p_val = T, add = "mean",
  ...)
```

Arguments

variable	character vector naming the variable to be plotted
data_ls	data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, numericals = < vector with column names of numerical columns>)</dataframe>
group	character vector naming the column to be used as grouping variable, Default: NULL
graph_type	one of c("violin", "bar", "line"), Default: 'violin'
y_axis	one of c("count", "density"), Default: c("count", "density")
auto_range	boolean, Default: T
n_breaks	integer, Default: 30
rug	boolean
x_min	double, requires aut_range == F, Default: 0
x_max	double, requires aut_range == F, Default: 100
title	character vector plot title
col_vector	vector with RGB colors, Default: f_plot_adjust_col_vector_length(100, RColorBrewer::brewer.pal(name = "Dark2", n = 8))
p_val	boolean, Default: T#'@param additional arguments passed to labs()
add	character vector one_of(c('mean', 'median', 'none')) , Default: 'mean'

Value

plot object

38 f_plot_pretty_points

```
#plot all variables
vars = data_ls$all_variables[ data_ls$all_variables != 'cyl' ] %>%
  map( f_plot_hist, data_ls, group = 'cyl')
vars
## End(Not run)
```

f_plot_obj_2_html

generate a separate html file from a shiny taglist

Description

lists of graphical objects like html(taglists), plots, tabplots, grids can be converted to html files

Usage

```
f_plot_obj_2_html(obj_list, type, output_file, title = "Plots", ...)
```

Arguments

```
obj_list htmltools::tagList

type one of c('taglist','plots','tabplots','grids')

output_file file_name of the html file, without .html suffix

title character vector of html document title, Default: 'Plots'
```

Examples

```
#returns a htmltools::taglist with DT::datatables and plotly plots
taglist = f_clean_data(mtcars) %>%
    f_boxcox() %>%
    f_pca() %>%
    f_pca_plot_components()

f_plot_obj_2_html(taglist, type = "taglist", output_file = 'test_me', title = 'Plots')
file.remove('test_me.html')
```

```
f_plot_pretty_points plot prettier dot plot
```

Description

color is contineoulsy scaled based on PC1 values and alpha values depend on point density.

Usage

```
f_plot_pretty_points(df, col_x, col_y, col_facet = NULL, size = 4,
   title = NULL, x_title = col_x, y_title = col_y, ...)
```

Arguments

df	datafram containing x,y pairs
col_x	character vector denoting x axis values
col_y	character vector denoting y axis values
col_facet	character vector denoting facetting column
size	size of points, Default: 4
	arguments passed to facet_wrap()

Details

Code adapted from https://drsimonj.svbtle.com/pretty-scatter-plots-with-ggplot2

Value

plot

See Also

interp.surface kde2d

Examples

```
df = ggplot2::diamonds %>%
   sample_n(2500)
col_x = 'carat'
col_y = 'price'
col_facet = 'cut'

f_plot_pretty_points(df, col_x, col_y, col_facet, title = 'price of diamonds by carat')
```

```
f_plot_profit_bars_plus_area
```

plot revenues cost and profit development over time with bars for revenue and costs and an area chart for profit.

Description

the function can graphically devide the chart into two periods e.g. past and future.

Usage

```
f_plot_profit_bars_plus_area(data, col_revenue, col_cost, col_time,
  now = max(data[, col_time]), unit_time = "years", unit_value = "CHF",
  title = "", alpha_past = 1, alpha_future = 0.5, alpha_past_area = 0.9,
  alpha_future_area = 0.7)
```

Arguments

```
datafram
data
                   character vector denoting revenue column
col_revenue
col_cost
                   character vector denoting cost column
col_time
                   character vector denoting time column
                   integer denoting a time which should be regarded as the breakpoint, Default:
now
                   max(data[, col_time])
                   character vector, will label y-axis, Default: 'years'
unit_time
                   character vector, will label x-axis, Default: 'CHF'
unit_value
title
                   character vector, will be title label, Default: "
                   double between 0 and 1 will determine alpha value for fill under the curve before
alpha_past
                   the breakpoint, Default: 1
alpha_future
                   double between 0 and 1 will determine alpha value for fill under the curve after
                   the breakpoint, Default: 0.5
alpha_past_area
                   as alpha_past but for area only, Default: 0.9
alpha_future_area
                   as alpha_future but for area only, Default: 0.7#'
```

Details

to some extent plotly compatible

Value

plot (to some extent plotly compatible)

```
data = tibble( time
                       = c(0,1,2,3,4,5,6,7,8,9,10,11,12)
              , revenue = - time^2 + time * 12
             , cost = revenue * 0.4 * -1
data[1,'cost'] = -10
data
print( f_plot_profit_bars_plus_area( data, 'revenue', 'cost', 'time') )
print( f_plot_profit_bars_plus_area( data, 'revenue', 'cost', 'time', now = 5) )
#clv figure for presenation
p = f_plot_profit_bars_plus_area( data, 'revenue', 'cost', 'time', now = 5, alpha_past_area = 0) +
  theme( panel.grid.major = element_blank()
        , panel.grid.minor = element_blank()
        , axis.text
                         = element_blank()
       )+
  labs(x = '', y = '')
print(p)
```

f_plot_profit_lines 41

f_plot_profit_lines

plot revenues cost and profit development over time as an area chart.

Description

the function can graphically devide the chart into two periods e.g. past and future.

Usage

```
f_plot_profit_lines(data, col_revenue, col_cost, col_time, now = max(data[,
    col_time]), unit_time = "years", unit_value = "CHF", title = "",
    alpha_past = 1, alpha_future = 0.5)
```

Arguments

data	datafram
col_revenue	character vector denoting revenue column
col_cost	character vector denoting cost column
col_time	character vector denoting time column
now	integer denoting a time which should be regarded as the breakpoint, Default: max(data[, col_time])
unit_time	character vector, will label y-axis, Default: 'years'
unit_value	character vector, will label x-axis, Default: 'CHF'
title	character vector, will be title label, Default: "
alpha_past	double between 0 and 1 will determine alpha value for fill under the curve before the breakpoint, Default: 1
alpha_future	double between 0 and 1 will determine alpha value for fill under the curve after the breakpoint, Default: 0.5

Details

not plotly compatibel

Value

plot (is not plotly compatibel)

42 f_plot_time

f_plot_time

plot variable distribution over time as reduced overlapping boxplots

Description

It is difficult to compare two timeerieses when you have more than one observation per timepoint without reducing all observations to a single statistical variable such as average or mean. This visualisation plots the median and the upper and lower 25 contineuos line between the medians of the timepoints.

Usage

```
f_plot_time(variable, time_variable, data_ls, time_variable_as_factor = F,
  group = NULL, normalize = F, time_unit = "day")
```

Arguments

variable character vector naming the variable to be plotted time_variable character vector naming the timevariable to be plotted data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, data_ls numericals = < vector with column names of numerical columns>) time_variable_as_factor If TRUE will convert time_variable to a factor, this will equalize the distance between timepoints on the plots and drops the connective line between timepoints, Default: F character vector naming the column to be used as grouping variable, Default: group **NULL** normalize If TRUE y variable will be divided by x variable, usefull if y variable represents a cumulated sum, Default: F time_unit character vector used as an x-axis lable, Default: 'day'

Value

plot

```
## End(Not run)
```

Description

add predictions to modelling dataframe and unnest, create a title column and a bins column

Usage

```
f_predict_plot_model_performance_regression(data)
```

Arguments

data

dataframe with the columns title, bins, resid_abs, resid_squ, ape

Value

taglist

See Also

```
tagList ggplotly datatable f_predict_pl_regression
```

```
## Not run:
form = as.formula( 'displacement~cylinders+mpg')
ISLR::Auto %>%
pipelearner::pipelearner() %>%
pipelearner::learn_models( twidlr::rpart, form ) %>%
pipelearner::learn_models( twidlr::randomForest, form ) %>%
pipelearner::learn_models( twidlr::svm, form ) %>%
pipelearner::learn() %>%
 f_predict_pl_regression( 'name' ) %>%
unnest(preds) %>%
mutate( bins = cut(target1, breaks = 3 , dig.lab = 4)
         , title = paste(models.id, cv_pairs.id, train_p, target, model) ) %>%
 f\_predict\_plot\_model\_performance\_regression() \ \%>\%
f_plot_obj_2_html(type = 'taglist', 'test_me', title = 'Model Performance')
file.remove('test_me.html')
## End(Not run)
```

```
f_predict_pl_regression
```

adds predictions to learned pipelearner dataframe

Usage

```
f_predict_pl_regression(pl, cols_id = NULL, formula = NULL,
  col_model = "fit", col_target = "target", data_test = "test",
  data_train = "train")
```

Arguments

pl learned pipelearner dataframe cols_id character vector naming id columns

Value

dataframe

See Also

```
f_predict_regression_add_predictions
```

Examples

```
form = as.formula( 'disp~cyl+mpg')

pl = mtcars %>%
   mutate(names = row.names(.)) %>%
   pipelearner::pipelearner() %>%
   pipelearner::learn_models( twidlr::rpart, form ) %>%
   pipelearner::learn_models( twidlr::randomForest, form ) %>%
   pipelearner::learn_models( twidlr::svm, form ) %>%
   pipelearner::learn_models( gamlss::gamlss, form ) %>%
   pipelearner::learn() %>%
   f_predict_pl_regression( cols_id = 'names' )
```

Description

use this function to get a quick summary of pipelearner dataframe with unnested predictions. Will group by title

Usage

```
f_predict_pl_regression_summarize(pl)
```

Arguments

pl

pipelearner dataframe with nested predictions

Value

dataframe with mape, mea, rtmse and median versions

See Also

```
f_predict_pl_regression
```

Examples

```
form = as.formula( 'disp~cyl+mpg')

pl = mtcars %>%
    mutate(names = row.names(.)) %>%
    pipelearner::pipelearner() %>%
    pipelearner::learn_models( twidlr::rpart, form ) %>%
    pipelearner::learn_models( twidlr::randomForest, form ) %>%
    pipelearner::learn_models( twidlr::svm, form ) %>%
    pipelearner::learn() %>%
    f_predict_pl_regression( 'names' ) %>%
    unnest( preds , .drop = FALSE ) %>%
    mutate( title = model ) %>%
    f_predict_pl_regression_summarize()
```

```
f_predict_regression_add_predictions
```

adds predictions, residuals, abolute residuals, squared residuals and absolute percent error to a dataframe.

Description

```
absolute percent error = (abs(resid/pred)*100)
```

Usage

```
f_predict_regression_add_predictions(data_test, m, col_target,
  data_train = NULL, cols_id = NULL, formula = NULL, ...)
```

Arguments

m	regression model
col_target	character vector naming target/response variable
cols_id	character vector naming id columns, if specified non_id columns will be dropped from dataframe, in order to be more memory efficient.
df	dataframe containing data to be used as the basis for prediction. Can also be a modelR resample object

46 f_stat_anova

Details

works with HDtweedie, randomForest, rpart, e1071::svm, glmnet, gamlss

Value

dataframe

Examples

```
df = mtcars %>%
mutate(names = row.names(.))
m = rpart::rpart(disp~., df)
pred = f_predict_regression_add_predictions(df, m, 'disp', 'names')
```

f_stat_anova

generate a dataframe with anova results from a data_ls list

Description

returns a dataframe with shapiro, anova und kruskal p values supplemented with maximum difference of means and medians between groups

Usage

```
f_stat_anova(data_ls, col_group, boxcox = F)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, data_ls numericals = < vector with column names of numerical columns>) character vector denoting grouping variable

col_group

boxcox perform analysis on boxcox-transformed or numerical variables

Value

dataframe

See Also

```
str_c map,map_dbl f_stat_anova
```

```
df_anova = data_ls = f_clean_data(mtcars) %>%
  f_stat_anova('cyl')
df_anova
```

f_stat_chi_square 47

f_stat_chi_square

generate a datatframe with chi square results from a data_ls list

Description

FUNCTION_DESCRIPTION

Usage

```
f_stat_chi_square(data_ls, col_group)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>,

numericals = < vector with column names of numerical columns>)

col_group character vector denoting grouping variable

Value

dataframe

See Also

```
is_empty,map,map_dbl
```

Examples

```
data_ls = f_clean_data(mtcars)
df_chi_squ = f_stat_chi_square(data_ls, 'cyl')
df_chi_squ
```

```
f_stat_combine_anova_with_chi_square
```

combines anova with chi square results into single dataframe

Description

keeps wither the anova or the kruskal p value depending on the results of the shapiro test (shapiro_stat $> 0.9 \text{ p_val} > 0.05$) and keeps the difference of percent of the mean. Still works if one of the input dataframes is NULL

Usage

```
f_stat_combine_anova_with_chi_square(df_anova = NULL, df_chi_square = NULL)
```

Arguments

```
df_anova dataframe created with f_stat_anova, Default: NULL df_chi_square dataframe created with f_stat_chi_square(), Default: NULL
```

Details

DETAILS

Value

```
OUTPUT_DESCRIPTION
```

Examples

```
data_ls = f_clean_data(mtcars)
df_chi_squ = f_stat_chi_square(data_ls, 'cyl')
df_anova = f_stat_anova(data_ls, 'cyl')
df_comb = f_stat_combine_anova_with_chi_square(df_anova, df_chi_squ)
df_comb
df_comb = f_stat_combine_anova_with_chi_square(df_anova)
df_comb
df_comb = f_stat_combine_anova_with_chi_square(df_chi_square = df_chi_squ)
df_comb
```

```
f_stat_diff_of_means_medians
```

calculates maximum difference in group means and medians

Description

used as a helper function for f_stat_anova

Usage

```
f_stat_diff_of_means_medians(df, col_group, col_variable)
```

Arguments

```
df dataframe

col_group character vector denoting grouping variable

col_variable character vector denoting variable
```

Value

dataframe

f_stat_group_ana 49

```
f_stat_diff_of_means_medians(df, col_group, 'v1') %>%
  bind_rows( f_stat_diff_of_means_medians(df, col_group, 'v2') ) %>%
  bind_rows( f_stat_diff_of_means_medians(df, col_group, 'v3') )
```

f_stat_group_ana

analyse group difference of dataset

Description

creates a html document with a group analysis including:

- P value table
- Dynamic Plots of all significant features
- static plots with brackets indicating statistical differences
- Tabplot
- · Alluvial Plot
- table containing means and medians for numerical variables
- table containing counts and percentages for categorical variables

The function automatically renders three html pages one for the additional static plots, one for the tableplot and one for the alluvial plots. In the same directry that can be determined by the outputfile parameter. Default behaviour will also render the entire html document returning the filepath of the new html file. The other three html files will be linked to in the document. You can modify the function to return a htmltools taglist instead. The above mentioned 3 additional html files for the other types of plots will still be rendered though with default settings. These extra plots can be switched off though.

Usage

```
f_stat_group_ana(data_ls, col_group, thresh_p_val = 0.05,
    thresh_diff_perc = 3, output_file = "group_ana", static_plots = T,
    alluvial = T, alluvial_thresh_p_val = 0.05,
    alluvial_thres_diff_perc = 7.5, max_alluvial_flows = 1500, tabplot = T,
    return_taglist = F, fig.width = 16, fig.height = 10)
```

Arguments

	data_ls	data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, numericals = < vector with column names of numerical columns>)</dataframe>
	col_group	character vector denoting grouping columns
	thresh_p_val	p value threshold for plots, Default: 0.05
thresh_diff_perc		
		minimum percent difference threshold for plots, Default: 3
	output_file	character vector containing output file name
	static_plots	boolean, render static plots indicating statistical differences with brackets, Default = TRUE
	alluvial	boolean, render alluvial plot, Default: TRUE

alluvial_thresh_p_val

double, threshold for feature to be inlcuded in alluvial plot. Features that are not highly significant and convey a large percental difference will result in a high number of flows thus cluttering the plot. It is not recommended to set these thresholds lower than the default. Default: 0.05

max_alluvial_flows

integer, maximum number of alluvial flows. Alluvial Plots can take a long time to render. Rendering an alluvial plot with the default setting of 1500 should take at least 10 min. Default 1500

tabplot boolean, render tabplot threshold for features are the same as for the dynamic

plots, Default: TRUE , static_plots = T

return_taglist boolean, return taglist instead of rendereing the final html document and return-

ing the link to the html file. Usefull if analysis should be directly included into

the current markdown document.

fig.width integer Width of Alluvial and Tabplot in inches. Default values can be comfort-

ably viewed on a 1920 x 1080 screen resolution. Default: 16

fig.height integer height of Alluvial and Tabplot in inches. Default values can be comfort-

ably viewed on a 1920 x 1080 screen resolution. Default: 10

alluvial_thresh_diff_perc

double, threshold for feature to be inleuded in alluvial plot. Features that are not highly significant and convey a large percental difference will result in a high number of flows thus cluttering the plot. It is not recommended to set these thresholds lower than the default. Default: 7.5

Value

file path to html file / or taglist

See Also

```
ggplotly tagList,h1,h2
```

Examples

```
## Not run:
    data_ls = f_clean_data(mtcars)
    f_stat_group_ana(data_ls, 'cyl', output_file = 'test_me')
    file.remove('test_me.html')
    file.remove('test_me_stat_plots.html')
    file.remove('test_me_alluvial.html')
    file.remove('test_me_tabplots.html')
## End(Not run)
```

```
f_stat_group_ana_taglist
```

analyse group difference of dataset

Description

returns a full analysis as a taglist inluding all features with p_values, medians, means, percentages and counts, as well as plots passing the treshold values

Usage

```
f_stat_group_ana_taglist(data_ls, col_group, tresh_p_val = 0.05,
    thresh_diff_perc = 3)
```

Arguments

thresh_diff_perc minimum percent difference threshold for plots, Default: 3

Value

taglist

See Also

```
ggplotly tagList,h1,h2
```

Examples

```
## Not run:
data_ls = f_clean_data(mtcars)
taglist = f_stat_group_ana_taglist(data_ls, 'cyl')
f_plot_obj_2_html(taglist, type = "taglist", output_file = 'test_me', title = 'Plots')
file.remove('test_me.html')
## End(Not run)
```

```
f_stat_group_counts_percentages
```

create a aggregated data frame with percentagers and counts for categorical variables

Usage

```
f_stat_group_counts_percentages(data_ls, col_group)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>, numericals = < vector with column names of numerical columns>)

col_group character vector denoting grouping columns

Value

dataframe

Examples

```
f_clean_data( mtcars) %>%
  f_stat_group_counts_percentages('cyl')
```

```
f_stat_group_mean_medians
```

create a aggregated data frame with means and medians for numerical variables

Usage

```
f_stat_group_mean_medians(data_ls, col_group)
```

Arguments

data_ls object generated by f_clean_data(), or a named list list(data = <dataframe>,

numericals = < vector with column names of numerical columns>)

col_group character vector denoting grouping columns

Value

dataframe

Examples

```
f_clean_data( mtcars) %>%
  f_stat_group_mean_medians('cyl')
```

```
f_stat_max_diff_of_freq
```

calculate the maximal difference in frequencies between to categorical variables

Description

used as a helper function for f_stat_chi_square

Usage

```
f_stat_max_diff_of_freq(df, col_var1, col_var2)
```

Arguments

df dataframe containing both avariables
col_var1 character vector denoting variable column 1
col_var2 character vector denoting variable column 2

Value

dataframe

f_stat_shapiro 53

Examples

```
data_ls = f_clean_data(mtcars)
df_chi_squ = f_stat_max_diff_of_freq(data_ls$data, 'cyl', 'gear')
```

f_stat_shapiro

wrapper for shapiro.test()

Description

shapiro.test i slimited to <5000 sample size and raises an error if sd(x) == 0. Wrapper samples from input vector and returns a list object with NA parameters if sd(x) == 0.

Usage

```
f_stat_shapiro(vec)
```

Arguments

vec

numeric vector

Value

```
shapiro.test object or list( statistic = NA, p.value = NA)
```

Examples

```
f_stat_shapiro( rnorm(1000, 10, 1) )
f_stat_shapiro( runif(1000, 1, 10) )
```

f_stat_stars

calculate significant level from p value

Description

```
* P:0.05, ** P:0,005, *** P:0.001
```

Usage

```
f_stat_stars(p_value)
```

Arguments

p_value

numeric

Value

character vector

f_train_lasso

Examples

```
f_stat_stars(0.06)
f_stat_stars(0.05)
f_stat_stars(0.005)
f_stat_stars(0.001)
```

f_train_lasso

wrapper for cv.glmnet and cv.HDtweedie

Description

performs lasso for different distributions, returns a list of formulas that result in the lowest rtmse for at least one of the distributions. Graphical output allows side-by-side comparison of lasso behaviour for all distributions.

Usage

```
f_train_lasso(data, formula, p = c(1, 1.25, 1.5, 1.75, 2), k = 5, family = "gaussian", ...)
```

Arguments

data	dataframe
formula	formula
p	p parameter for tweedie distributions, set p = NULL for not performing lasso for tweedie distributions, Default: $c(1, 1.25, 1.5, 1.75, 2)$
k	fold cross validation, Default: 5
family	family parameter for glmnet, can be a vector, Default: 'gaussian'
	arguments passed to cv.glmnet, cv.HDtweedie such as lambda or n_lambda
grid	grid values for lambda, Default: 10^seq(4, -4, length = 100)

Details

Columns containing NA will be removed, formula cannot be constructed with '.'

Value

list()

See Also

```
,HDtweedie ,glmnet ,cv.HDtweedie ,cv.glmnet ,pipelearner ,learn_models ,learn_cvpairs
,learn
```

Examples

```
f_train_lasso_manual_cv
```

wrapper for glmnet and HDtweedie

Description

performs lasso for different distributions, returns a list of formulas that result in the lowest rtmse for at least one of the distributions. Graphical output allows side-by-side comparison of lasso behaviour for all distributions.

Usage

```
f_train_lasso_manual_cv(data, formula, grid = 10^seq(4, -4, length = 100), p = c(1, 1.25, 1.5, 1.75, 2), k = 5, family = "gaussian")
```

Arguments

data	dataframe
formula	formula
grid	grid values for lambda, Default: 10^seq(4, -4, length = 100)
р	p parameter for tweedie distributions, set $p = NULL$ for not performing lasso for tweedie distributions, Default: $c(1, 1.25, 1.5, 1.75, 2)$
k	fold cross validation, set to 1 for testing against training data, Default: 5
family	family parameter for glmnet, can be a vector, Default: 'gaussian'

Details

Columns containing NA will be removed, formula cannot be constructed with '.'

Value

list()

See Also

HDtweedie glmnet pipelearner,learn_models,learn_cvpairs,learn

Examples

hello

Hello, World!

Description

Prints 'Hello, world!'.

Usage

hello()

Examples

hello()

```
make_container_for_function_calls

container for function calls, can be used as a progress bar
```

Description

creates a closure with a make_call() method that wraps any function call. When the wrapper is used the function call is saved and the calls are counted and the progress is being printed. Use the method set_total() to input the total number of function calls. Based on the total an ETA is estimated and a percentage calculated.

Usage

```
make_container_for_function_calls()
```

Details

DETAILS

Value

container

See Also

```
now,time_length
```

```
.f = randomForest::randomForest
call_cont = make_container_for_function_calls()
call_cont$set_total(4)
m_wr = call_cont$make_call( .f = .f, formula = disp~., data = mtcars )
#pipe version
call_cont = make_container_for_function_calls()
call_cont$set_total(5)
pl = pipelearner::pipelearner(mtcars) %>%
  pipelearner::learn_models( models = c( call_cont$make_call )
                             , formulas = c(disp^{-}.)
                             , .f = c( randomForest::randomForest )
                             , function_name = 'randomForest'
                             , print_call = c(T)
                           ) %>%
  pipelearner::learn_cvpairs( pipelearner::crossv_kfold, k = 5 ) %>%
  pipelearner::learn()
```

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