# Package 'structToolbox' 

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## Type Package

Title Data processing \& analysis tools for Metabolomics and other omics

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Description An extensive set of data (pre-)processing and analysis methods and tools for metabolomics and other omics, with a strong emphasis on statistics and machine learning. This toolbox allows the user to build extensive and standardised workflows for data analysis. The methods and tools have been implemented using class-based templates provided by the struct (Statistics in R Using Class-based Templates) package. The toolbox includes pre-processing methods (e.g. signal drift and batch correction, normalisation, missing value imputation and scaling), univariate (e.g. ttest, various forms of ANOVA, Kruskal-Wallis test and more) and multivariate statistical methods (e.g. PCA and PLS, including cross-validation and permutation testing) as well as machine learning methods (e.g. Support Vector Machines). The STATistics Ontology (STATO) has been integrated and implemented to provide standardised definitions for the different methods, inputs and outputs.
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ANOVA Analysis of Variance

## Description

Analysis of Variance (ANOVA) is a univariate method used to analyse the difference among group means. Multiple test corrected p -values are computed to indicate significance for each feature.

## Usage



## Arguments

alpha (numeric) The p -value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".

```
formula (formula) A symbolic description of the model to be fitted.
ss_type (character) ANOVA sum of squares. Allowed values are limited to the follow-
        ing:
            - "I": Type I sum of squares.
            - "II": Type II sum of squares.
            - "III": Type III sum of squares.
            The default is "III".
... Additional slots and values passed to struct_class.
```


## Details

This object makes use of functionality from the following packages:

- car


## Value

A ANOVA object with the following output slots:

```
f_statistic (data.frame) The value of the calculated statistic.
    p_value (data.frame) The probability of observing the calculated statistic if the null hypothesis is true.
    significant (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold.
```


## Inheritance

A ANOVA object inherits the following struct classes:
[ANOVA]» [model]» [struct_class]

## References

Fox J, Weisberg S (2019). An R Companion to Applied Regression, Third edition. Sage, Thousand Oaks CA. https://socialsciences.mcmaster.ca/jfox/Books/Companion/.

## Examples

```
M = ANOVA(
        alpha = 0.05,
        mtc = "fdr",
        formula = y ~ x,
        ss_type = "III")
D = iris_DatasetExperiment()
M = ANOVA(formula=y~Species)
M = model_apply(M,D)
```

```
as_data_frame Convert to data.frame
```


## Description

Convert the outputs of the input model into a data.frame.

## Usage

\#\# S4 method for signature 'filter_na_count'
as_data_frame(M)
\#\# S4 method for signature 'ttest'
as_data_frame(M)
\#\# S4 method for signature 'wilcox_test'
as_data_frame(M)

## Arguments

M a model object

## Value

A data.frame of model outputs

## Examples

```
D = iris_DatasetExperiment()
M = filter_na_count(threshold=50,factor_name='Species')
M= model_apply(M,D)
df = as_data_frame(M)
```

AUC

Area under ROC curve

## Description

The area under the ROC curve of a classifier is estimated using the trapezoid method.

## Usage

AUC (...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A AUC object. This object has no output slots.

## Inheritance

A AUC object inherits the following struct classes:
[AUC]»[metric]» [struct_class]

## Examples

```
M = AUC()
D = iris_DatasetExperiment()
XCV = kfold_xval(folds=5,factor_name='Species') *
    (mean_centre() + PLSDA(number_components=2,factor_name='Species'))
MET = AUC()
XCV = run(XCV,D,MET)
```

```
autoscale Autoscaling
```


## Description

Each variable/feature is mean centred and scaled by the standard deviation. The transformed variables have zero-mean and unit-variance.

## Usage

autoscale(mode = "data", ...)

## Arguments

## mode

(character) Mode of action. Allowed values are limited to the following:

- "data": Autoscaling is applied to the data matrix only.
- "sample_meta": Autoscaling is applied to the sample_meta data only.
- "both": Autoscaling is applied to both the data matrix and the meta data.

The default is "data".
... Additional slots and values passed to struct_class.

## Value

A autoscale object with the following output slots:

| autoscaled | (DatasetExperiment) |
| :--- | :--- |
| mean_data | (numeric) |
| sd_data | (numeric) |
| mean_sample_meta | (numeric) |
| sd_sample_meta | (numeric) |

## Inheritance

A autoscale object inherits the following struct classes:
[autoscale]»[model]»[struct_class]

## Examples

```
    M = autoscale(
        mode = "data")
    D = iris_DatasetExperiment()
    M = autoscale()
    M = model_train(M,D)
    M = model_predict(M,D)
```

    balanced_accuracy Balanced Accuracy
    
## Description

Balanced Accuracy is the average proportion of correctly classified samples across all groups.

## Usage

balanced_accuracy(...)

## Arguments

Additional slots and values passed to struct_class.

## Value

A balanced_accuracy object. This object has no output slots.

## Inheritance

A balanced_accuracy object inherits the following struct classes:
[balanced_accuracy]»[metric]» [struct_class]

## Examples

```
M = balanced_accuracy()
D = iris_DatasetExperiment()
XCV = kfold_xval(folds=5,factor_name='Species') *
    (mean_centre() + PLSDA(number_components=2,factor_name='Species'))
MET = balanced_accuracy()
XCV = run(XCV,D,MET)
```

blank_filter Blank filter

## Description

A blank filter filters features by comparing the median intensity of blank samples to the median intensity of samples. Features where the relative intensity (fold change) is not large when compared to the blank are removed. The number of times a feature is detected across all blank samples may also be considered. If the feature is not detected in a high enough proportion of the blanks then it is not removed.

## Usage

```
blank_filter(
    fold_change = 20,
    blank_label = "blank",
    qc_label = "QC",
    factor_name,
    fraction_in_blank = 0,
)
```


## Arguments

fold_change (numeric) Features with fold change less than this value are removed. The default is 20.
blank_label (character) The label used to identify blank samples. The default is "blank".
qc_label (character, NULL) The label used to identify QC samples. If set to NULL then the median of the samples is used. The default is "QC".
factor_name (character) The name of a sample-meta column to use.
fraction_in_blank
(numeric) Features present in less than this proportion of the blanks are not considered for removal. The default is 0 .
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A blank_filter object with the following output slots:
filtered (DatasetExperiment) A DatasetExperiment object containing the filtered data.
flags (data.frame) A flag indicating whether the feature was rejected or not.

## Inheritance

A blank_filter object inherits the following struct classes:
[blank_filter]» [model]» [struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc.pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

```
M = blank_filter(
        fold_change = 20,
        blank_label = "Blank",
        qc_label = "QC",
        factor_name = "V1",
        fraction_in_blank = 0)
D = iris_DatasetExperiment()
M = blank_filter(fold_change=2,
    factor_name='Species',
    blank_label='setosa',
    qc_label='versicolor')
M = model_apply(M,D)
```


## Description

A histogram of the calculated fold changes for the blank filter (median samples divided by median blanks)

## Usage

blank_filter_hist(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A blank_filter_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A blank_filter_hist object inherits the following struct classes:
[blank_filter_hist]» [chart]» [struct_class]

## Examples

M = blank_filter_hist()
C = blank_filter_hist()

```
bootstrap Bootstrap resampling
```


## Description

In bootstrap resampling a subset of samples is selected at random with replacement to form a training set. Any sample not selected for training is included in the test set. This process is repeated many times, and performance metrics are computed for each repetition.

## Usage

bootstrap(number_of_repetitions = 100, collect, ...)

## Arguments

number_of_repetitions
(numeric, integer) The number of bootstrap repetitions. The default is 100 .
collect (character) The name of a model output to collect over all bootstrap repetitions, in addition to the input metric.
... Additional slots and values passed to struct_class.

## Value

A bootstrap object with the following output slots:

$$
\begin{array}{ll}
\text { results } & \text { (data.frame) } \\
\text { metric } & \text { (data.frame) } \\
\text { collected } & \text { (logical, list) }
\end{array}
$$

## Inheritance

A bootstrap object inherits the following struct classes:
[bootstrap]»[resampler]»[iterator]»[struct_class]

## Examples

$$
M=\text { bootstrap( }
$$

number_of_repetitions = 10, collect = "vip")

I = bootstrap(number_of_repetitions = 10, collect = 'vip')

```
calculate,AUC-method Calculate metric
```


## Description

Calculate metric

## Usage

```
## S4 method for signature 'AUC'
calculate(obj, Y, Yhat)
## S4 method for signature 'balanced_accuracy'
calculate(obj, Y, Yhat)
## S4 method for signature 'r_squared'
calculate(obj, Y, Yhat)
```


## Arguments

| obj | a metric object |
| :--- | :--- |
| $Y$ | the true values/group labels |
| Yhat | the predicted values/group labels |

## Value

a modified metric object

## Examples

```
MET = metric()
calculate(MET)
```

chart_plot,dfa_scores_plot, DFA-method
chart_plot method

## Description

Plots a chart object

## Usage

```
## S4 method for signature 'dfa_scores_plot,DFA'
chart_plot(obj, dobj)
## S4 method for signature 'scatter_chart,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'pca_correlation_plot,PCA'
chart_plot(obj, dobj)
## S4 method for signature 'pca_scores_plot,PCA'
chart_plot(obj, dobj)
## S4 method for signature 'pca_biplot,PCA'
chart_plot(obj, dobj)
## S4 method for signature 'pca_loadings_plot,PCA'
chart_plot(obj, dobj)
## S4 method for signature 'pca_scree_plot,PCA'
chart_plot(obj, dobj)
## S4 method for signature 'pca_dstat_plot,PCA'
```

```
chart_plot(obj, dobj)
## S4 method for signature 'plsr_prediction_plot,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'plsr_residual_hist,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'plsr_qq_plot,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'plsr_cook_dist,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'pls_scores_plot,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'plsda_predicted_plot,PLSDA'
chart_plot(obj, dobj)
## S4 method for signature 'plsda_roc_plot,PLSDA'
chart_plot(obj, dobj)
## S4 method for signature 'pls_vip_plot,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'pls_regcoeff_plot,PLSR'
chart_plot(obj, dobj)
## S4 method for signature 'blank_filter_hist,blank_filter'
chart_plot(obj, dobj)
## S4 method for signature 'confounders_lsq_barchart,confounders_clsq'
chart_plot(obj, dobj)
## S4 method for signature 'confounders_lsq_boxplot,confounders_clsq'
chart_plot(obj, dobj)
## S4 method for signature 'feature_boxplot,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'mv_histogram,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'mv_boxplot,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'DatasetExperiment_dist,DatasetExperiment'
```

```
chart_plot(obj, dobj)
## S4 method for signature 'DatasetExperiment_boxplot,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'compare_dist,DatasetExperiment'
chart_plot(obj, dobj, eobj)
## S4 method for signature 'DatasetExperiment_heatmap,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'DatasetExperiment_factor_boxplot,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'feature_profile_array,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'feature_profile,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'fold_change_plot,fold_change'
chart_plot(obj, dobj)
## S4 method for signature 'fs_line,forward_selection_by_rank'
chart_plot(obj, dobj)
## S4 method for signature 'glog_opt_plot,glog_transform'
chart_plot(obj, dobj, gobj)
## S4 method for signature 'gs_line,grid_search_1d'
chart_plot(obj, dobj)
## S4 method for signature 'hca_dendrogram,HCA'
chart_plot(obj, dobj)
## S4 method for signature 'kfoldxcv_grid,kfold_xval'
chart_plot(obj, dobj)
## S4 method for signature 'kfoldxcv_metric,kfold_xval'
chart_plot(obj, dobj)
## S4 method for signature 'kw_p_hist,kw_rank_sum'
chart_plot(obj, dobj)
## S4 method for signature 'mv_feature_filter_hist,mv_feature_filter'
chart_plot(obj, dobj)
## S4 method for signature 'mv_sample_filter_hist,mv_sample_filter'
```

```
chart_plot(obj, dobj)
## S4 method for signature 'permutation_test_plot,permutation_test'
chart_plot(obj, dobj)
## S4 method for signature 'plsda_feature_importance_plot,PLSDA'
chart_plot(obj, dobj)
## S4 method for signature 'pqn_norm_hist,pqn_norm'
chart_plot(obj, dobj)
## S4 method for signature 'resample_chart,resample'
chart_plot(obj, dobj)
## S4 method for signature 'rsd_filter_hist,rsd_filter'
chart_plot(obj, dobj)
## S4 method for signature 'feature_profile,sb_corr'
chart_plot(obj, dobj, gobj)
## S4 method for signature 'svm_plot_2d,SVM'
chart_plot(obj, dobj, gobj)
## S4 method for signature 'tSNE_scatter,tSNE'
chart_plot(obj, dobj)
## S4 method for signature 'tic_chart,DatasetExperiment'
chart_plot(obj, dobj)
## S4 method for signature 'wilcox_p_hist,wilcox_test'
chart_plot(obj, dobj)
```


## Arguments

obj a chart object
dobj a struct object
eobj a second DatasetExperiment object to compare with the first
gobj The DatasetExperiment object before signal correction was applied.

## Value

a plot object

## Examples

```
C = example_chart()
chart_plot(C,iris_DatasetExperiment())
```


## Description

In univariate classical least squares regression a line is fitted between each feature/variable and a response variable. The fitted line minimises the sum of squared differences between the true response and the predicted response. The coefficients (offset, gradient) of the fit can be tested for significance.

## Usage



## Arguments

alpha (numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc $\quad \begin{aligned} & \text { (character) Multiple test correction method. Allowed values are limited to the } \\ & \text { following: } \\ & \quad \text { • "bonferroni": Bonferroni correction in which the p-values are multiplied }\end{aligned}$ by the number of comparisons.

- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
factor_names (character, list) The column names to regress against. If a character vector then the same list is used ofr all features. If a list of character vectors is provided it is assumed there is a different set of columns for each feature.
intercept (logical) Model intercept. Allowed values are limited to the following:

- "TRUE": An intercept term is included in the model.
- "FALSE": An intercept term is not included in the model.

The default is TRUE.
... Additional slots and values passed to struct_class.

## Value

A classical_lsq object with the following output slots:
coefficients (data.frame) The regression coefficients for each term in the model.
p_value (data.frame) The probability of observing the calculated statistic if the null hypothesis is true.
significant (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold
r_squared (data.frame) The value of R Squared for the fitted model.
adj_r_squared (data.frame) The value ofAdjusted R Squared for the fitted model.

## Inheritance

A classical_lsq object inherits the following struct classes:
[classical_lsq]»[model]»[struct_class]

## Examples

```
M = classical_lsq(
        alpha = 0.05,
        mtc = "fdr",
        factor_names = "V1",
        intercept = FALSE)
D = iris_DatasetExperiment()
M = classical_lsq(factor_names = 'Species')
M = model_apply(M,D)
```

```
    compare_dist Compare distributions
```


## Description

Histograms and boxplots computed across samples and features are used to visually compare two datasets e.g. before and after filtering and/or normalisation.

## Usage

compare_dist(factor_name, ...)

## Arguments

| factor_name | (character) The name of a sample-meta column to use. |
| :--- | :--- |
| ... | Additional slots and values passed to struct_class. |

## Value

A compare_dist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A compare_dist object inherits the following struct classes:
[compare_dist]» [chart]» [struct_class]

## Examples

```
M = compare_dist(
            factor_name = "V1")
D1=MTBLS79_DatasetExperiment(filtered=FALSE)
D2=MTBLS79_DatasetExperiment(filtered=TRUE)
C = compare_dist(factor_name='Class')
chart_plot(C,D1,D2)
```

confounders_clsq Check for confounding factors

## Description

Univariate least squares regression models are used to compare models with and without potential confounding factors included. The change in coefficients (delta) is then computed for each potential confounding factor. Factors with a large delta are said to be having a large impact on the model and are therefore confounding. p-values are computed for models with confounders included to reduce potential false positives. Only suitable for main factors with 2 levels.

```
Usage
    confounders_clsq(
        alpha = 0.05,
        mtc = "fdr",
        factor_name,
        confounding_factors,
        threshold = 0.15,
    )
```


## Arguments

alpha (numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
factor_name (character) The name of the main factor with which other factors may be confounding.
confounding_factors
(character) The name(s) of factor(s) that are potential confounding factors.
threshold (numeric) Factors with a delta greater than the the threshold are considered to be confounding. The default is 0.15 .
... Additional slots and values passed to struct_class.

## Value

A confounders_clsq object with the following output slots:

| coefficients | (data.frame) |
| :--- | :--- |
| p_value | (data.frame) |
| significant | (data.frame) |
| percent_change | (data.frame) |
| potential_confounders | (list) |

## Inheritance

A confounders_clsq object inherits the following struct classes:
[confounders_clsq]» [model]» [struct_class]

## Examples

```
M = confounders_clsq(
        alpha = 0.05,
        mtc = "fdr",
        factor_name = character(0),
        confounding_factors = character(0),
        threshold = 0.15)
D = MTBLS79_DatasetExperiment()
M = filter_by_name(mode='include',dimension='variable',
        names=colnames(D$data)[1:10]) + # first 10 features
        filter_smeta(mode='exclude',levels='QC',
        factor_name='Class') + # reduce to two group comparison
    confounders_clsq(factor_name = 'Class',
        confounding_factors=c('run_order','Batch'))
M = model_apply(M,D)
```

confounders_lsq_barchart

## Description

A barchart of the relative change (delta) in regression coefficient when potential confounding factors are included, and excluded, from the model. Factors with a large delta are considered to be confounding factors.

## Usage

confounders_lsq_barchart(feature_to_plot, threshold = 10, ...)

## Arguments

feature_to_plot (numeric, character, integer) The column name of the feature to be plotted.
threshold (numeric) A horizontal line is plotted to indicate the threshold. The default is 10.

$$
\ldots \quad \text { Additional slots and values passed to struct_class. }
$$

## Value

A confounders_lsq_barchart object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A confounders_lsq_barchart object inherits the following struct classes:
[confounders_lsq_barchart]» [chart]» [struct_class]

## Examples

```
M = confounders_lsq_barchart(
            feature_to_plot = 1,
            threshold = 10)
D = MTBLS79_DatasetExperiment()
M = filter_by_name(mode='include',dimension='variable',
            names=colnames(D$data)[1:10]) + # first 10 features
        filter_smeta(mode='exclude',levels='QC',
        factor_name='Class') + # reduce to two group comparison
    confounders_clsq(factor_name = 'Class',
        confounding_factors=c('run_order','Batch'))
M = model_apply(M,D)
C = C=confounders_lsq_barchart(feature_to_plot=1,threshold=15)
chart_plot(C,M[3])
```

confounders_lsq_boxplot

Confounding factor relative change boxplot

## Description

A boxplot of the relative change (delta) in regression coefficient when potential confounding factors are included, and excluded, from the model. Factors with a large delta are considered to be confounding factors.

## Usage

confounders_lsq_boxplot(threshold = 10, ...)

## Arguments

threshold (numeric) A horizontal line is plotted to indicate the threshold. The default is 10.
... Additional slots and values passed to struct_class.

## Value

A confounders_lsq_boxplot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A confounders_lsq_boxplot object inherits the following struct classes:
[confounders_lsq_boxplot] » [chart] » [struct_class]

## Examples

```
M = confounders_lsq_boxplot(
        threshold = 10)
D = MTBLS79_DatasetExperiment()
M = filter_by_name(mode='include',dimension='variable',
        names=colnames(D$data)[1:10]) + # first 10 features
        filter_smeta(mode='exclude',levels='QC',
        factor_name='Class') + # reduce to two group comparison
    confounders_clsq(factor_name = 'Class',
        confounding_factors=c('run_order','Batch'))
M = model_apply(M,D)
C = C=confounders_lsq_boxplot(threshold=15)
chart_plot(C,M[3])
```

    constant_sum_norm Normalisation to constant sum
    
## Description

Each sample is normalised such that the total signal is equal to one (or a scaling factor if specified).

## Usage

```
    constant_sum_norm(scaling_factor = 1, ...)
```


## Arguments

scaling_factor (numeric) The scaling factor applied after normalisation. The default is 1 .
... Additional slots and values passed to struct_class.

## Value

A constant_sum_norm object with the following output slots:
normalised (DatasetExperiment) A DatasetExperiment object containing the normalised data. coeff (data.frame) The sum of each row, used to normalise the samples.

## Inheritance

A constant_sum_norm object inherits the following struct classes:
[constant_sum_norm] » [model]» [struct_class]

## Examples

$$
\begin{aligned}
& M=\text { constant_sum_norm( } \\
& \text { scaling_factor }=1) \\
& M=\text { constant_sum_norm() }
\end{aligned}
$$

```
corr_coef Correlation coefficient
```


## Description

The correlation between features and a set of continuous factor are calculated. Multiple-test corrected p-values are used to indicate whether the computed coefficients may have occurred by chance.

## Usage

corr_coef(alpha = 0.05, mtc = "fdr", factor_names, method = "spearman", ...)

## Arguments

alpha (numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
factor_names (character) The name of sample meta column(s) to use.
method (character) Type of correlation. Allowed values are limited to the following:

- "kendall": Kendall's tau is computed.
- "pearson": Pearson product moment correlation is computed.
- "spearman": Spearman's rho statistic is computed.

The default is "spearman".
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- stats


## Value

A corr_coef object with the following output slots:
coeff (data.frame) The value of the calculate statistics which is converted to a p-value when compared to a t-distrib
p_value (data.frame) The probability of observing the calculated statistic if the null hypothesis is true.
significant (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold.

## Inheritance

A corr_coef object inherits the following struct classes:
[corr_coef]»[model]»[struct_class]

## References

R Core Team (2023). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/.

## Examples

$M=$ corr_coef $($
alpha $=0.05$, mtc = "fdr", factor_names = "V1", method = "spearman")

D = MTBLS79_DatasetExperiment(filtered=TRUE)
\# subset for this example
$D=D[, 1: 10]$
\# convert to numeric for this example
D\$sample_meta\$sample_order=as.numeric(D\$sample_meta\$run_order)
D\$sample_meta\$sample_rep=as.numeric(D\$sample_meta\$Sample_Rep)
$M=$ corr_coef(factor_names=c('sample_order', 'sample_rep'))
M = model_apply (M, D)

## Description

A boxplot to visualise the distribution of values within a subset of features.

## Usage

DatasetExperiment_boxplot( factor_name, by_sample = TRUE, per_class = TRUE, number $=50$,
)

## Arguments

| factor_name | (character) The name of a sample-meta column to use. |
| :--- | :--- |
| by_sample | (logical) Plot by sample. Allowed values are limited to the following: |
|  | - "TRUE": The data is plotted across features for a subset of samples. |
|  | - "FALSE": The data is plotted across samples for a subset of features. |

The default is TRUE.
per_class (logical) Plot per class. Allowed values are limited to the following:

- "TRUE": The data is plotted for each class.
- "FALSE": The data is plotted for all samples.

The default is TRUE.
number (numeric, integer) The number of features/samples plotted. The default is 50.
$\ldots \quad$ Additional slots and values passed to struct_class.

## Value

A DatasetExperiment_boxplot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.
struct object

## Inheritance

A DatasetExperiment_boxplot object inherits the following struct classes:
[DatasetExperiment_boxplot]» [chart]»[struct_class]

## Examples

```
M = DatasetExperiment_boxplot(
    factor_name = "V1",
    by_sample = FALSE,
    per_class = FALSE,
    number = 50)
D = MTBLS79_DatasetExperiment()
C = DatasetExperiment_boxplot(factor_name='Class',number=10,per_class=FALSE)
chart_plot(C,D)
```

DatasetExperiment_dist

Feature distribution histogram

## Description

A histogram to visualise the distribution of values within features.

## Usage

DatasetExperiment_dist(factor_name, per_class = TRUE, ...)

## Arguments

factor_name (character) The name of a sample-meta column to use.
per_class (logical) Plot per class. Allowed values are limited to the following:

- "TRUE": The distributions are plotted for each class.
- "FALSE": The distribution is plotted for all samples.

The default is TRUE.
... Additional slots and values passed to struct_class.

## Value

A DatasetExperiment_dist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A DatasetExperiment_dist object inherits the following struct classes:
[DatasetExperiment_dist]» [chart]» [struct_class]

## Examples

```
M = DatasetExperiment_dist(
    factor_name = "V1",
    per_class = FALSE)
D = MTBLS79_DatasetExperiment()
C = DatasetExperiment_dist(factor_name='Class')
chart_plot(C,D)
```

DatasetExperiment_factor_boxplot
Factor boxplot

## Description

Boxplot for a feature to visualise the distribution of values within each group

## Usage

DatasetExperiment_factor_boxplot(feature_to_plot, factor_names, ...)

## Arguments

feature_to_plot (character, numeric, integer) The name of the plotted feature.
factor_names (character) The name of sample meta column(s) to use.
... Additional slots and values passed to struct_class.

## Value

A DatasetExperiment_factor_boxplot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A DatasetExperiment_factor_boxplot object inherits the following struct classes:
[DatasetExperiment_factor_boxplot]» [chart]»[struct_class]

## Examples

```
M = DatasetExperiment_factor_boxplot(
    factor_names = "V1",
    feature_to_plot = "V1")
D = iris_DatasetExperiment()
C = DatasetExperiment_factor_boxplot(factor_names='Species',feature_to_plot='Petal.Width')
chart_plot(C,D)
```

DatasetExperiment_heatmap

## DatasetExperiment heatmap

## Description

A heatmap to visualise the measured values in a data matrix.

## Usage

DatasetExperiment_heatmap(na_colour = "\#FF00E4", ...)

## Arguments

na_colour (character) The hex colour code used to plot missing values. The default is "\#FF00E4".
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- reshape2


## Value

A DatasetExperiment_heatmap object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A DatasetExperiment_heatmap object inherits the following struct classes:
[DatasetExperiment_heatmap]» [chart]» [struct_class]

## References

Wickham H (2007). "Reshaping Data with the reshape Package." Journal of Statistical Software, 21(12), 1-20. http://www.jstatsoft.org/v21/i12/.

## Examples

```
M = DatasetExperiment_heatmap(
    na_colour = "#FF00E4")
D = iris_DatasetExperiment()
C = DatasetExperiment_heatmap()
chart_plot(C,D)
```

```
DFA Discriminant Factor Analysis
```


## Description

Discriminant Factor Analysis (DFA) is a supervised classification method. Using a linear combination of the input variables, DFA finds new orthogonal axes (canonical values) to minimize the variance within each given class and maximize variance between classes.

## Usage

DFA(factor_name, number_components = 2, ...)

## Arguments

factor_name (character) The name of a sample-meta column to use.
number_components
(numeric, integer) The number of DFA components calculated. The default is 2.
... Additional slots and values passed to struct_class.

## Value

A DFA object with the following output slots:

| scores | (DatasetExperiment) |
| :--- | :--- |
| loadings | (data.frame) |
| eigenvalues | (data.frame) |
| that | (DatasetExperiment) |

## Inheritance

A DFA object inherits the following struct classes:
[DFA] » [model]» [struct_class]

## References

Manly B (1986). Multivariate Statistical Methods: A Primer. Chapman and Hall, Boca Raton.

## Examples

```
M = DFA(
    factor_name = "V1",
    number_components = 2)
D = iris_DatasetExperiment()
M = DFA(factor_name='Species')
M = model_apply(M,D)
```

```
dfa_scores_plot DFA scores plot
```


## Description

A scatter plot of the selected DFA components.

## Usage

dfa_scores_plot(
components $=c(1,2)$,
points_to_label = "none",
factor_name,
ellipse = "all",
label_filter = character(0),
label_factor = "rownames",
label_size = 3.88,
)

## Arguments

components (numeric) The components selected for plotting. The default is $c(1,2)$.
points_to_label
(character) Points to label. Allowed values are limited to the following:

- "none": No samples labels are displayed.
- "all": The labels for all samples are displayed.
- "outliers": Labels for for potential outlier samples are displayed.

The default is "none".
factor_name (character) The name of a sample-meta column to use.
ellipse (character) Plot ellipses. Allowed values are limited to the following:

- "all": Hotelling T2 ellipses ( $\mathrm{p}=0.95$ ) are plotted for all groups and all samples.
- "group": Hotelling T2 ellipses $(\mathrm{p}=0.95)$ are plotted for all groups.
- "none": Ellipses are not included on the plot.
- "sample": A Hotelling T2 ellipse ( $\mathrm{p}=0.95$ ) is plotted for all samples (ignoring group).
The default is "all".
label_filter (character) Labels are only plotted for the named groups. If zero-length then all groups are included. The default is character (0).
label_factor (character) The column name of sample_meta to use for labelling samples on the plot. "rownames" will use the row names from sample_meta. The default is "rownames".
label_size (numeric) The text size of labels. Note this is not in Font Units. The default is 3.88.
... Additional slots and values passed to struct_class.


## Details

This object makes use of functionality from the following packages:

- scales
- ggplot2

Value
A dfa_scores_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A dfa_scores_plot object inherits the following struct classes:
[dfa_scores_plot]» [chart] » [struct_class]

## References

Wickham H, Pedersen T, Seidel D (2023). scales: Scale Functions for Visualization. R package version 1.3.0, https://CRAN.R-project.org/package=scales.
Wickham H (2016). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York. ISBN 978-3-319-24277-4, https://ggplot2. tidyverse.org.

## Examples

```
M = dfa_scores_plot(
    components = c(1, 2),
    points_to_label = "none",
    factor_name = "V1",
    ellipse = "all",
    label_filter = character(0),
    label_factor = "rownames",
    label_size = 3.88)
```

```
D = iris_DatasetExperiment()
M = mean_centre() + DFA(factor_name='Species')
M = model_apply(M,D)
C = dfa_scores_plot(factor_name = 'Species')
chart_plot(C,M[2])
```

dratio_filter Dispersion ratio filter

## Description

The dispersion ratio (d-ratio) compares the standard deviation (or non-parametric equivalent) of the Quality Control (QC) samples relative to the standard deviation (or non-parametric equivalent) of the samples for each feature. If the d-ratio is greater than a predefined threshold then the observed sample variance could be due to technical variance and the feature is removed.

## Usage

```
dratio_filter(
```

    threshold \(=20\),
    qc_label = "QC",
    factor_name,
    method = "ratio",
    dispersion = "sd",
    )
    
## Arguments

threshold
(numeric) The threshold above which features are removed. The default is 20.
qc_label (character) The label used to identify QC samples. The default is "QC".
factor_name
method
(character) The name of a sample-meta column to use.
(character) dratio method. Allowed values are limited to the following:

- "ratio": Dispersion of the QCs divided by the dispersion of the samples. Corresponds to Eq 4 in Broadhurst et al (2018).
- "euclidean": Dispersion of the QCs divided by the euclidean length of the total dispersion. Total dispersion is estimated from the QC and Sample dispersion by assuming that they are orthogonal. Corresponds to Eq 5 in Broadhurst et al (2018).
The default is "ratio".
dispersion (character) Dispersion method. Allowed values are limited to the following:
- "sd": Dispersion is estimated using the standard deviation.
- "mad": Dispersion is estimated using the median absolute deviation.

The default is "sd".
... Additional slots and values passed to struct_class.

## Value

A dratio_filter object with the following output slots:

$$
\begin{array}{ll}
\text { filtered } & \text { (DatasetExperiment) A DatasetExperiment object containing the filtered data. } \\
\text { flags } & \text { (data.frame) Flag indicating whether the feature was rejected by the filter or not. } \\
\text { d_ratio } & \text { (data.frame) }
\end{array}
$$

## Inheritance

A dratio_filter object inherits the following struct classes:
[dratio_filter]» [model]»[struct_class]

## References

Broadhurst D, Goodacre R, Reinke SN, Kuligowski J, Wilson ID, Lewis MR, Dunn WB (2018). "Guidelines and considerations for the use of system suitability and quality control samples in mass spectrometry assays applied in untargeted clinical metabolomic studies." Metabolomics, 14(6).

## Examples

```
M = dratio_filter(
    threshold = 20,
    qc_label = "QC",
    factor_name = "V1",
    method = "ratio",
    dispersion = "sd")
D = MTBLS79_DatasetExperiment()
M = dratio_filter(threshold=20,qc_label='QC',factor_name='Class')
M = model_apply(M,D)
```

equal_split Equal group sized sampling

## Description

Samples are randomly chosen from each level such that the training set has equal numbers of samples for all levels. The number of samples is based on the input proportion and the smallest group size.

## Usage

equal_split(p_train = 1, factor_name, ...)

## Arguments

| p_train | (numeric) The proportion of samples selected for the training set. The default is |
| :--- | :--- |
|  | 1. |
| factor_name | (character) The name of a sample-meta column to use. |
| $\ldots$ | Additional slots and values passed to struct_class. |

## Value

A equal_split object with the following output slots:
training (DatasetExperiment) A DatasetExperiment object containing samples selected for the training set. testing (DatasetExperiment) A DatasetExperiment object containing samples selected for the testing set.

## Inheritance

A equal_split object inherits the following struct classes:
[equal_split]»[split_data]»[model]»[struct_class]

## Examples

```
    M = equal_split(
        factor_name = "V1",
        p_train = 0.75)
D = iris_DatasetExperiment()
M = equal_split(factor_name='Species')
M = model_apply(M,D)
```

    feature_boxplot Feature boxplot
    
## Description

A boxplot to visualise the distribution of values within a feature.

## Usage

```
feature_boxplot(
    label_outliers = TRUE,
    feature_to_plot,
    factor_name,
    show_counts = TRUE,
    style = "boxplot",
    jitter = FALSE,
    fill = FALSE,
    ...
)
```


## Arguments

label_outliers (logical) Label outliers. Allowed values are limited to the following:

- "TRUE": The index for outlier samples is included on the plot.
- "FALSE": No labels are displayed.

The default is TRUE.
feature_to_plot
(character, numeric, integer) The column name of the plotted feature.
factor_name (character) The name of a sample-meta column to use.
show_counts (logical) Show counts. Allowed values are limited to the following:

- "TRUE": The number of samples for each box is displayed.
- "FALSE": The number of samples for each box is not displayed.

The default is TRUE.
style (character) Plot style. Allowed values are limited to the following:

- "boxplot": Boxplot style.
- "violin": Violon plot style.

The default is "boxplot".
jitter (logical) Include points plotted with added jitter. The default is FALSE.
fill (logical) Block fill the boxes or violins with the group colour. The default is FALSE.
... Additional slots and values passed to struct_class.

## Value

A feature_boxplot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A feature_boxplot object inherits the following struct classes:
[feature_boxplot]» [chart]» [struct_class]

## Examples

M = feature_boxplot(
label_outliers = FALSE,
feature_to_plot = "V1",
factor_name = "V1",
show_counts = FALSE,
style = "boxplot",
jitter = FALSE,
fill $=$ FALSE)
D = MTBLS79_DatasetExperiment
C = feature_boxplot(factor_name='Species', feature_to_plot='Petal.Width')
chart_plot(C,D)
feature_profile Feature profile

## Description

A plot visualising the change in intensity of a feature with a continuous variable such as time, dose, or run order.

## Usage

feature_profile( run_order, qc_label, qc_column, colour_by, feature_to_plot, plot_sd = FALSE, )

## Arguments

| run_order | (character) The sample-meta column name containing run order. |
| :--- | :--- |
| qc_label | (character) The label used to identify QC samples. |
| qc_column | (character) The sample-meta column name containing the labels used to identify |
|  | QC samples. |
| colour_by | (character) The sample-meta column name to used to colour the plot. |

feature_to_plot
(numeric, character, integer) The name or column id of the plotted feature.
plot_sd (logical) Plot standard deviation. Allowed values are limited to the following:

- "TRUE": Standard deviation of samples and QCs are included on the plot.
- "FALSE": Standard deviation is not plotted.

The default is FALSE.
... Additional slots and values passed to struct_class.

## Value

A feature_profile object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A feature_profile object inherits the following struct classes:
[feature_profile]»[chart]»[struct_class]

## Examples

```
M = feature_profile(
    run_order = character(0),
    qc_label = character(0),
    qc_column = character(0),
    colour_by = character(0),
    feature_to_plot = numeric(0),
    plot_sd = FALSE)
D = MTBLS79_DatasetExperiment()
C = feature_profile(run_order='run_order',
    qc_label='QC',
    qc_column='Class',
    colour_by='Class',
    feature_to_plot=1)
chart_plot(C,D)
```

feature_profile_array Feature profile

## Description

A plot visualising the change in intensity of a feature with a continuous variable such as time, dose, or run order.

## Usage

feature_profile_array( run_order, qc_label, qc_column, colour_by, feature_to_plot, nrow $=5$, log = TRUE,

$$
\ldots
$$

)

## Arguments

| run_order | (character) The sample-meta column name containing run order. |
| :---: | :---: |
| qc_label | (character) The label used to identify QC samples. |
| qc_column | (character) The sample-meta column name containing the labels used to identify QC samples. |
| colour_by | (character) The sample-meta column name to used to colour the plot. |
| feature_to_plot |  |
|  | (numeric, character, integer) The name or column id of the plotted feature. |
| nrow | (numeric, integer) The number of rows in the plot. The default is 5 . |
| $\log$ | (logical) Log transform. Allowed values are limited to the following: |
|  | - "TRUE": The data is log tranformed before plotting. |
|  | - "FALSE": The data is not transformed before plotting. |

The default is TRUE.
... Additional slots and values passed to struct_class.

## Value

A feature_profile_array object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A feature_profile_array object inherits the following struct classes:
[feature_profile_array]» [chart]» [struct_class]

## Examples

$$
\begin{aligned}
& \text { M = feature_profile_array }( \\
& \text { run_order }=\text { character }(0), \\
& \text { qc_label }=\text { character }(0),
\end{aligned}
$$

```
        qc_column = character(0),
        colour_by = character(0),
        feature_to_plot = numeric(0),
        nrow = 1,
        log = FALSE)
    D = MTBLS79_DatasetExperiment()
    C = feature_profile_array(
    run_order='run_order',
    qc_label='QC',
    qc_column='Class',
    colour_by='Class',
    feature_to_plot=1:3,
    nrow=1,
    log=TRUE)
chart_plot(C,D)
```

filter_by_name Filter by name

## Description

Filter samples/variables by row/column name, index or logicals.

## Usage

filter_by_name(mode = "exclude", dimension = "sample", names, ...)

## Arguments

| mode | (character) The filtering mode controls whether samples/features are mode="included" <br> or mode="excluded" based on their name. The default is "exclude". <br> (character) The filtering dimensions controls whether dimension="sample" or <br> dimension="variable" are filtered based on their name. The default is "sample". <br> (character, numeric, logical) The name of features/samples to be filtered. Must <br> be an exact match. Can also provide indexes (numeric) or logical. |
| :--- | :--- |
| names | Additional slots and values passed to struct_class. |
| $\ldots$ |  |

## Value

A filter_by_name object with the following output slots:
filtered (DatasetExperiment)

## Inheritance

A filter_by_name object inherits the following struct classes:
[filter_by_name]» [model]» [struct_class]

## Examples

```
M = filter_by_name(
    mode = "exclude",
    dimension = "sample",
    names = character(0))
D = MTBLS79_DatasetExperiment()
M = filter_by_name(mode='exclude',dimension='variable',names=c(1, 2, 3))
M = model_apply(M,D)
```

filter_na_count Minimum number of measured values filter

## Description

The number of measured values is counted for each feature, and any feature with less than a predefined minimum number of values in each group is removed. If there are several factors, then the threshold is applied so that the minimum number of samples is present for all combinations (interactions) of groups.

## Usage

filter_na_count(threshold, factor_name, ...)

## Arguments

threshold (numeric) The minimum number of samples in each group/interaction.
factor_name (character) The name of a sample-meta column to use.
... Additional slots and values passed to struct_class.

## Value

A filter_na_count object with the following output slots:
filtered (DatasetExperiment) A DatasetExperiment object containing the filtered data.
count (data.frame) The number of measured values in each group/interaction.
na_count (data.frame) The number of missing values in each group/interaction.
flags (data.frame) Flags to indicate which features were removed.

## Inheritance

A filter_na_count object inherits the following struct classes:
[filter_na_count]» [model]»[struct_class]

## Examples

```
    M = filter_na_count(
    threshold = 2,
    factor_name = "V1")
D = MTBLS79_DatasetExperiment()
M = filter_na_count(threshold=3,factor_name='Class')
M = model_apply(M,D)
```

    filter_smeta Filter by sample meta data
    
## Description

The data is filtered by so that the named levels of a factor are included/excluded from the dataset.

## Usage

filter_smeta(mode = "include", levels, factor_name, ...)

## Arguments

mode (character) Mode of action. Allowed values are limited to the following:

- "include": Samples in the specified levels are retained.
- "exclude": Samples in the specified levels are excluded.

The default is "include".
levels (character) The level name(s) for filtering.
factor_name (character) The name of a sample-meta column to use.
... Additional slots and values passed to struct_class.

## Value

A filter_smeta object with the following output slots:

> filtered (DatasetExperiment)

## Inheritance

A filter_smeta object inherits the following struct classes:
[filter_smeta]»[model]»[struct_class]

## Examples

```
M = filter_smeta(
        mode = "include",
        levels = character(0),
        factor_name = "V1")
D = MTBLS79_DatasetExperiment()
M = filter_smeta(mode='exclude',levels='QC',factor_name='QC')
M = model_apply(M,D)
```

    fisher_exact Fisher Exact Test
    
## Description

A fisher exact test is used to analyse contingency tables by comparing the number of correctly/incorrectly predicted group labels. A multiple test corrected p-value indicates whether the number of measured values is significantly different between groups.

## Usage

fisher_exact(alpha = 0.05, mtc = "fdr", factor_name, factor_pred, ...)

## Arguments

| alpha | (numeric) The p-value cutoff for determining significance. The default is 0.05. |
| :---: | :---: |
| mtc | (character) Multiple test correction method. Allowed values are limited to the following: |
|  | - "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons. <br> - "fdr": Benjamini and Hochberg False Discovery Rate correction. <br> - "none": No correction. |
|  | The default is "fdr". |
| factor_name | (character) The name of a sample-meta column to use. |
| factor_pred | (data.frame) A data.frame, where each column is a factor of predicted group labels to compare with the true groups labels. |
|  | Additional slots and values passed to struct_class. |

## Value

A fisher_exact object with the following output slots:
p_value (data.frame) The probability of observing the calculated statistic if the null hypothesis is true.
significant (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold.

## Inheritance

A fisher_exact object inherits the following struct classes:
[fisher_exact]» [model]» [struct_class]

## Examples

```
M = fisher_exact(
    alpha = 0.05,
    mtc = "fdr",
    factor_name = "V1",
    factor_pred = data.frame(id=NA))
# load some data
D=MTBLS79_DatasetExperiment()
# prepare predictions based on NA
pred=as.data.frame(is.na(D$data))
pred=lapply(pred,factor,levels=c(TRUE,FALSE))
pred=as.data.frame(pred)
# apply method
M = fisher_exact(alpha=0.05,mtc='fdr',factor_name='Class',factor_pred=pred)
M=model_apply(M,D)
```

fold_change Fold change

## Description

Fold change is the relative change in mean (or non-parametric equivalent) intensities of a feature between all pairs of levels in a factor.

## Usage

```
fold_change(
    factor_name,
    paired = FALSE,
    sample_name = character(0),
    threshold = 2,
    control_group = character(0),
    method = "geometric",
    conf_level = 0.95,
)
```


## Arguments



## Value

A fold_change object with the following output slots:
fold_change (data.frame) The fold change between groups.
lower_ci (data.frame) Lower confidence interval for fold change.
upper_ci (data.frame) Upper confidence interval for fold change.
significant (data.frame) A logical indictor of whether the calculated fold change including the estimated confidence limit

## Inheritance

A fold_change object inherits the following struct classes:

```
[fold_change]»[model]»[struct_class]
```


## References

Price Jr RM, Bonett DG (2020). "Confidence Intervals for Ratios of Means and Medians." Journal of Educational and Behavioral Statistics, 45(6), 750-770.

## Examples

```
M = fold_change(
    factor_name = "V1",
    sample_name = character(0),
    paired = FALSE,
    threshold = 2,
    control_group = character(0),
    method = "geometric",
    conf_level = 0.95)
D = MTBLS79_DatasetExperiment()
M = fold_change(factor_name='Class')
M = model_apply(M,D)
```

    fold_change_int Fold change for interactions between factors
    
## Description

For more than one factor the fold change calculation is extended to include all combinations of levels (interactions) of all factors. Paired fold changes are not possible for this computation.

## Usage

```
fold_change_int(
    factor_name,
    threshold = 2,
    control_group = character(0),
    method = "geometric",
    conf_level = 0.95,
    )
```


## Arguments

\(\left.\begin{array}{ll}factor_name (character) The name of a sample-meta column to use. <br>
threshold <br>
(numeric) The fold change threshold for labelling features as significant. The <br>

default is 2.\end{array}\right]\)| control_group (character) The level names of the groups used in the denominator (where pos- |
| :--- |
| sible) when computing fold change. One level for each factor, assumed to be in |
| the same order as factor_name. The default is character (0). |
| (character) Fold change method. Allowed values are limited to the following: |
| - "geometric": A log transform is applied before using group means to cal- |
| culate fold change. In the non-tranformedspace this is equivalent to using |
| geometric group means. Confidence intervals for independant and paired |
| sampling are estimated using standard error of the mean in log transformed |
| space before being transformed back to the original space. |
| - "median": The group medians and the method described by Price and |
| Bonett is used to estimate confidence intervals. For paired data standard |
| error of the median is used to estimate confidence intervals from the me- |
| dian fold change of all pairs. |
| - "mean": The group means and the method described by Price and Bonnet is |
| used to estimate confidence intervals. For paired data standard error of the |
| mean is used to estimate confidence intervals from the mean fold change of |

## Value

A fold_change_int object with the following output slots:
fold_change (data.frame) The fold change between groups.
lower_ci (data.frame) Lower confidence interval for fold change.
upper_ci (data.frame) Upper confidence interval for fold change.
significant (data.frame) A logical indictor of whether the calculated fold change including the estimated confidence limit

## Inheritance

A fold_change_int object inherits the following struct classes:
[fold_change_int]»[fold_change]» [model] » [struct_class]

## References

Lloyd GR, Jankevics A, Weber RJM (2020). "struct: an R/Bioconductor-based framework for standardized metabolomics data analysis and beyond." Bioinformatics, 36(22-23), 5551-5552. https:
//doi.org/10.1093/bioinformatics/btaa1031.

## Examples

```
    M = fold_change_int(
        factor_name = "V1",
        sample_name = character(0),
    threshold = 2,
    control_group = character(0),
    method = "geometric",
    paired = FALSE,
    conf_level = 0.95)
D = MTBLS79_DatasetExperiment()
D=D[,1:10,drop=FALSE]
M = filter_smeta(mode='exclude',levels='QC',factor_name='Class') +
    fold_change_int(factor_name=c('Class','Batch'))
M = model_apply(M,D)
```

fold_change_plot Fold change plot

## Description

A plot of fold changes calculated for a chosen subset of features. A predefined fold change threshold is indicated by shaded regions.

## Usage

fold_change_plot(number_features = 20, orientation = "portrait", ...)

## Arguments

number_features
(numeric) The number randomly selected features to plot, or a list of column numbers. The default is 20 .
orientation (character) Plot orientation. Allowed values are limited to the following:

- "landscape": Features are plotted on the y-axis.
- "portrait": Features are plotted on the x-axis.

The default is "portrait".
... Additional slots and values passed to struct_class.

## Value

A fold_change_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A fold_change_plot object inherits the following struct classes:
[fold_change_plot]» [chart]» [struct_class]

## Examples

M = fold_change_plot (
number_features = 10 ,
orientation = "portrait")
C = fold_change_plot()
forward_selection_by_rank

> Forward selection by rank

## Description

A model is trained and performance metric computed by including increasing numbers of features in the model. The features to be included in each step are defined by their rank, which is computed from another variable e.g. VIP score. An "optimal"subset of features is suggested by minimising the input performance metric.

## Usage

forward_selection_by_rank(
min_no_vars = 1, max_no_vars = 100, step_size = 1, factor_name, variable_rank,
)

## Arguments

min_no_vars
max_no_vars (numeric) The maximum number of variables to include in the model. The default is 100 .
step_size (numeric) The incremental change in number of features in the model. The default is 1 .

| factor_name | (character) The name of a sample-meta column to use. |
| :--- | :--- |
| variable_rank | (numeric, integer) The values used to rank the features. |
| $\ldots$ | Additional slots and values passed to struct_class. |

## Value

A forward_selection_by_rank object with the following output slots:
$\begin{array}{ll}\text { metric } & \text { (data.frame) The value of the computed metric for each model. For nested models the metric is averaged. } \\ \text { results } & \text { (data.frame) The predicted outputs from collated from all models computed during forward selection. } \\ \text { chosen_vars } & \text { (numeric, integer) The column number of the variables chosen for the best performing model. } \\ \text { smoothed } & \text { (numeric) The value of the performance metric for each evaluated model after smoothing. } \\ \text { searchlist } & \text { (numeric) The maxmimum rank of features included in each model. }\end{array}$

## Inheritance

A forward_selection_by_rank object inherits the following struct classes:
[forward_selection_by_rank]» [resampler]» [iterator]» [struct_class]

## Examples

```
M = forward_selection_by_rank(
    min_no_vars = 1,
    max_no_vars = 100,
    step_size = 1,
    factor_name = "V1",
    variable_rank = 1)
# some data
D = MTBLS79_DatasetExperiment(filtered=TRUE)
# normalise, impute and scale then remove QCs
P = pqn_norm(qc_label='QC',factor_name='Class') +
    knn_impute(neighbours=5) +
    glog_transform(qc_label='QC',factor_name='Class') +
    filter_smeta(mode='exclude',levels='QC',factor_name='Class')
P = model_apply(P,D)
D = predicted(P)
# forward selection using a PLSDA model
M = forward_selection_by_rank(factor_name='Class',
                                    min_no_vars=2,
                                    max_no_vars=11,
                                    variable_rank=1:2063) *
    (mean_centre() + PLSDA(number_components=1,
    factor_name='Class'))
M = run(M,D,balanced_accuracy())
```

fs_line Forward selection line plot

## Description

A line plot for forward selection. The computed model performance metric is plotted against the number of features included in the model.

## Usage

fs_line(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A fs_line object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A fs_line object inherits the following struct classes:
[fs_line]» [chart]»[struct_class]

## Examples

```
M = fs_line()
# some data
D = MTBLS79_DatasetExperiment(filtered=TRUE)
# normalise, impute and scale then remove QCs
P = pqn_norm(qc_label='QC',factor_name='Class') +
    knn_impute(neighbours=5) +
    glog_transform(qc_label='QC',factor_name='Class') +
    filter_smeta(mode='exclude',levels='QC',factor_name='Class')
P = model_apply(P,D)
D = predicted(P)
# forward selection using a PLSDA model
M = forward_selection_by_rank(factor_name='Class',
                                    min_no_vars=2,
                                    max_no_vars=11,
                                    variable_rank=1:2063) *
    (mean_centre() + PLSDA(number_components=1,
    factor_name='Class'))
```

```
M = run(M,D,balanced_accuracy())
# chart
C = fs_line()
chart_plot(C,M)
```

glog_opt_plot Glog optimisation

## Description

A plot of the sum of squares error (SSE) vs different values of lambda for the glog transform. The indicated optimum value for lambda minimises the SSE.

## Usage

glog_opt_plot(plot_grid $=100, \ldots$ )

## Arguments

plot_grid (numeric) The default is 100.
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A glog_opt_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A glog_opt_plot object inherits the following struct classes:
[glog_opt_plot]» [chart]» [struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc.pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

```
M = glog_opt_plot(
        plot_grid = numeric(0))
D = iris_DatasetExperiment()
M = glog_transform(qc_label='versicolor',factor_name='Species')
M = model_apply(M,D)
C = glog_opt_plot()
chart_plot(C,M,D)
```

glog_transform Generalised logarithmic transform

## Description

The generalised logarithm (glog) transformation applies a log transformation while applying an offset to account for technical variation.

## Usage

glog_transform(qc_label = "QC", factor_name, lambda = NULL, ...)

## Arguments

| qc_label | (character) The label used to identify QC samples. The default is "QC". |
| :--- | :--- |
| factor_name | (character) The name of a sample-meta column to use. |
| lambda | (numeric, NULL) The value of lambda to use. If NULL then the pmp package <br> will be used to determine an "optimal" value for lambda. The default is NULL. |
| $\ldots$ | Additional slots and values passed to struct_class. |

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A glog_transform object with the following output slots:
transformed (DatasetExperiment) A DatasetExperiment object containing the glog transformed data.
error_flag (logical) A logical indicating whether the glog optimisation for lambda was successful. If not then PMP return

## Inheritance

A glog_transform object inherits the following struct classes:
[glog_transform]» [model]» [struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc. pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.
Durbin B, Hardin J, Hawkins D, Rocke D (2002). "A variance-stabilizing transformation for geneexpression microarray data." Bioinformatics, 18 (Suppl 1), S105-S110.

Parsons HM, Ludwig C, Gunther UL, Viant MR (2007). "Improved classification accuracy in 1- and ', '2-dimensional NMR metabolomics data using the variance ', 'stabilising generalised logarithm transformation." Bioinformatics, 8(1), 234.

## Examples

```
M = glog_transform(
    qc_label = "QC",
    factor_name = "V1",
    lambda = NULL)
D = iris_DatasetExperiment()
M = glog_transform(qc_label='versicolor',factor_name='Species')
M = model_apply(M,D)
```

```
grid_search_1d One dimensional grid search
```


## Description

A one dimensional grid search calculates a performance metric for a model at evenly spaced values for a model input parameter. The "optimum" value for the parameter is suggested as the one which maximises performance, or minimises error (whichever is appropriate for the chosen metric)

## Usage

```
grid_search_1d(
    param_to_optimise,
    search_values,
    model_index,
    factor_name,
    max_min = "min",
)
```


## Arguments

param_to_optimise
(character) The name of the model input parameter that is the focus of the search.
search_values (ANY) The values of the input parameter being optimised.

model_index | (numeric, integer) The index of the model in the sequence that uses the parame- |
| :--- |
| ter being optimised. |
| factor_name (character) The name of a sample-meta column to use. |
| max_min |
| (character) Maximise or minimise. Allowed values are limited to the following: |
| - "max": The optimium parameter value is suggested based on maximising |
| the performance metric. |
| • "min": The optimium parameter value is suggested based on minimising |
| the performance metric. |

The default is "min".
Additional slots and values passed to struct_class.

## Value

A grid_search_1d object with the following output slots:

| results | (data.frame) |
| :--- | :--- |
| metric | (data.frame) |
| optimum_value | (numeric) |

## Inheritance

A grid_search_1d object inherits the following struct classes:
[grid_search_1d]» [resampler]» [iterator]» [struct_class]

## Examples

```
M = grid_search_1d(
    param_to_optimise = character(0),
    search_values = numeric(0),
    model_index = numeric(0),
    factor_name = "V1",
    max_min = "min")
D = MTBLS79_DatasetExperiment()
# some preprocessing
M = pqn_norm(qc_label='QC',factor_name='Class') +
    knn_impute() +
    glog_transform(qc_label='QC',factor_name='Class') +
    filter_smeta(factor_name='Class',levels='QC',mode='exclude')
M=model_apply(M,D)
D=predicted(M)
# reduce number of features for this example
D=D[,1:10]
# optmise number of components for PLS model
I = grid_search_1d(param_to_optimise='number_components',search_values=1:5,
```

```
    model_index=2,factor_name='Class') *
    (mean_centre()+PLSDA(factor_name='Class'))
    I = run(I,D,balanced_accuracy())
```

    gs_line Grid search line plot
    
## Description

A plot of the calculated performance metric against the model input parameter values used to train the model. The optimum parameter value is indicated based on minimising (or maximising) the chosen metric.

## Usage

gs_line(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A gs_line object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A gs_line object inherits the following struct classes:
[gs_line]» [chart]» [struct_class]

## Examples

$$
\begin{aligned}
& \text { M = gs_line() } \\
& \text { C = gs_line() }
\end{aligned}
$$

## HCA Hierarchical Cluster Analysis

## Description

Hierarchical Cluster Analysis is a numerical technique that uses agglomerative clustering to identify clusters or groupings of samples.

```
Usage
    HCA(
        dist_method = "euclidean",
        cluster_method = "complete",
        minkowski_power = 2,
        factor_name,
    )
```


## Arguments

dist_method
(character) Distance measure. Allowed values are limited to the following:

- "euclidean": The euclidean distance (2 norm).
- "maximum": The maximum distance.
- "manhattan": The absolute distance (1 norm).
- "canberra": A weighted version of the mahattan distance.
- "minkowski": A generalisation of manhattan and euclidean distance to nth norm.
The default is "euclidean".
cluster_method (character) Agglomeration method. Allowed values are limited to the following:
- "ward.D": Ward clustering.
- "ward.D2": Ward clustering using sqaured distances.
- "single": Single linkage.
- "complete": Complete linkage.
- "average": Average linkage (UPGMA).
- "mcquitty": McQuitty linkage (WPGMA).
- "median": Median linkage (WPGMC).
- "centroid": Centroid linkage (UPGMC).

The default is "complete".
minkowski_power
(numeric) The default is 2.
factor_name (character) The name of a sample-meta column to use.
Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- stats


## Value

A HCA object with the following output slots:

```
dist_matrix (dist) An object containing pairwise distance information between samples.
hclust (hclust) An object of class hclust which describes the tree produced by the clustering process.
factor_df (data.frame)
```


## Inheritance

A HCA object inherits the following struct classes:
[HCA]» [model]»[struct_class]

## References

R Core Team (2023). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/.

## Examples

```
M = HCA(
    dist_method = "euclidean",
    cluster_method = "complete",
    minkowski_power = numeric(0),
    factor_name = "V1")
D = iris_DatasetExperiment()
M = HCA(factor_name='Species')
M = model_apply(M,D)
```

hca_dendrogram HCA dendrogram

## Description

A dendrogram visualising the clustering by HCA.

## Usage

hca_dendrogram(...)

## Arguments

... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- ggdendro

Value
A hca_dendrogram object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A hca_dendrogram object inherits the following struct classes:
[hca_dendrogram]» [chart]» [struct_class]

## References

de Vries A, Ripley BD (2022). ggdendro: Create Dendrograms and Tree Diagrams Using 'ggplot2'. R package version 0.1 .23 , https://CRAN.R-project.org/package=ggdendro.

## Examples

M = hca_dendrogram()
C = hca_dendrogram()

## Description

Tukey's HSD post hoc test is a modified t-test applied for all features to all pairs of levels in a factor. It is used to determine which groups are different (if any). A multiple test corrected p-value is computed to indicate which groups are significantly different to the others for each feature.

## Usage

HSD(alpha $=0.05$, mtc $=" f d r "$, formula, unbalanced $=$ FALSE,.. )

## Arguments

alpha
(numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
formula (formula) A symbolic description of the model to be fitted.
unbalanced (logical) Unbalanced model. Allowed values are limited to the following:

- "TRUE": A correction is applied for unbalanced designs.
- "FALSE": No correction is applied for unbalanced designs.

The default is FALSE.
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- agricolae


## Value

A HSD object with the following output slots:

```
difference (data.frame)
UCL (data.frame)
LCL (data.frame)
p_value (data.frame) The probability of observing the calculated statistic if the null hypothesis is true.
significant (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold.
```


## Inheritance

A HSD object inherits the following struct classes:
[HSD]» [model]» [struct_class]

## References

de Mendiburu F (2023). agricolae: Statistical Procedures for Agricultural Research. R package version 1.3-7, https://CRAN.R-project.org/package=agricolae.

## Examples

```
M = HSD(
    alpha = 0.05,
    mtc = "fdr",
    formula = y ~ x,
    unbalanced = FALSE)
D = iris_DatasetExperiment()
M = HSD(formula=y~Species)
M = model_apply(M,D)
```

```
HSDEM
```

Tukey's Honest Significant Difference using estimated marginal means

## Description

Tukey's HSD post hoc test is a modified t-test applied for all features to all pairs of levels in a factor. It is used to determine which groups are different (if any). A multiple test corrected p-value is computed to indicate which groups are significantly different to the others for each feature. For mixed effects models estimated marginal means are used.

## Usage

HSDEM(alpha = 0.05, mtc = "fdr", formula, ...)

## Arguments

alpha (numeric) The p -value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
formula (formula) A symbolic description of the model to be fitted.
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- emmeans
- nlme

Value
A HSDEM object with the following output slots:
$\begin{array}{ll}\text { p_value } & \text { (data.frame) The probability of observing the calculated statistic if the null hypothesis is true. } \\ \text { significant } & \text { (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold. }\end{array}$

## Inheritance

A HSDEM object inherits the following struct classes:
[HSDEM] » [model] » [struct_class]

## References

Lenth R (2023). emmeans: Estimated Marginal Means, aka Least-Squares Means. R package version 1.9.0, https://CRAN.R-project.org/package=emmeans.
Pinheiro J, Bates D, R Core Team (2023). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-164, https://CRAN.R-project.org/package=nlme.

Pinheiro JC, Bates DM (2000). Mixed-Effects Models in S and S-PLUS. Springer, New York. doi:10.1007/b98882 https://doi.org/10.1007/b98882.

## Examples

```
M = HSDEM(
        alpha = 0.05,
        mtc = "fdr",
        formula = y ~ x)
D = iris_DatasetExperiment()
D$sample_meta$id=rownames(D) # dummy id column
M = HSDEM(formula = y~Species+ Error(id/Species))
M = model_apply(M,D)
```

kfoldxcv_grid $\quad k$-fold cross validation plot

## Description

A graphic for visualising the true class and the predicted class of samples in all groups for all cross-validation folds.

## Usage

kfoldxcv_grid(factor_name, level, ...)

## Arguments

| factor_name | (character) The name of a sample-meta column to use. |
| :--- | :--- |
| level | (character) The level/group to plot. |
| $\ldots$ | Additional slots and values passed to struct_class. |

## Value

A kfoldxcv_grid object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A kfoldxcv_grid object inherits the following struct classes:
[kfoldxcv_grid]»[chart]»[struct_class]

## Examples

```
M = kfoldxcv_grid(
        factor_name = "V1",
        level = "level_1")
    D = iris_DatasetExperiment()
    I = kfold_xval(factor_name='Species') *
        (mean_centre() + PLSDA(factor_name='Species'))
    I = run(I,D,balanced_accuracy())
    C = kfoldxcv_grid(factor_name='Species',level='setosa')
    chart_plot(C,I)
```

    kfoldxcv_metric kfoldxcv metric plot
    
## Description

A boxplot of the performance metric computed for each fold of a k-fold cross-validation.

## Usage

kfoldxcv_metric(...)

## Arguments

... Additional slots and values passed to struct_class.

Value
A kfoldxcv_metric object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A kfoldxcv_metric object inherits the following struct classes:
[kfoldxcv_metric]»[chart]»[struct_class]

## Examples

M = kfoldxcv_metric()
C = kfoldxcv_metric()
kfold_xval $\quad k$-fold cross-validation

## Description

k -fold cross-validation is an iterative approach applied to validate models. The samples are divided into k "folds", or subsets. Each subset is excluded from model training and used for model validation once, resulting in a single left-out prediction for each sample. Model performance metrics are then computed for the training and test sets across all folds.

## Usage

kfold_xval(folds = 10, method = "venetian", factor_name, collect = NULL, ...)

## Arguments

folds (numeric, integer) The number of cross-validation folds. The default is 10.
method (character) Fold selection method. Allowed values are limited to the following:

- "venetian": Every nth sample is assigned to the same fold, where n is the number of folds.
- "blocks": Blocks of adjacent samples are assigned to the same fold.
- "random": Samples are randomly assigned to a fold.

The default is "venetian".
factor_name (character) The name of a sample-meta column to use.
collect
(NULL, character) The name of a model output to collect over all bootstrap repetitions, in addition to the input metric. The default is NULL.
... Additional slots and values passed to struct_class.

## Value

A kfold_xval object with the following output slots:

| results | (data.frame) |
| :--- | :--- |
| metric | (data.frame) |
| metric.train | (numeric) |
| metric.test | (numeric) |
| collected | (list) |

## Inheritance

A kfold_xval object inherits the following struct classes:
[kfold_xval]»[resampler]» [iterator]» [struct_class]

## Examples

```
    M = kfold_xval(
        folds = 5,
        method = "random",
        factor_name = "V1",
        collect = NULL)
    D = iris_DatasetExperiment()
    I = kfold_xval(factor_name='Species') *
    (mean_centre() + PLSDA(factor_name='Species'))
    I = run(I,D,balanced_accuracy())
```

    knn_impute \(\quad k N N\) missing value imputation
    
## Description

k -nearest neighbour missing value imputation replaces missing values in the data with the average of a predefined number of the most similar neighbours for which the value is present

## Usage

knn_impute( neighbours $=5$, sample_max = 50, feature_max $=50$, by = "features",
)

## Arguments

neighbours (numeric) The number of neighbours ( $k$ ) to use for imputation. The default is 5 .
sample_max (numeric) The maximum percent missing values per sample. The default is 50 .
feature_max (numeric) The maximum percent missing values per feature. The default is 50 .
by (character) Impute using similar "samples" or "features". Default features. The default is "features".
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A knn_impute object with the following output slots:
imputed (DatasetExperiment) A DatasetExperiment object containing the data where missing values have been imputed.

## Inheritance

A knn_impute object inherits the following struct classes:
[knn_impute] » [model] » [struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc. pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

```
M = knn_impute(
    neighbours = 5,
    feature_max = 50,
    sample_max = 50,
    by = "features")
M = knn_impute()
```

```
kw_p_hist Histogram of p values
```


## Description

A histogram of the p-values computed by the kruskal-wallis method

## Usage

```
    kw_p_hist(...)
```


## Arguments

... Additional slots and values passed to struct_class.

## Value

A kw_p_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A kw_p_hist object inherits the following struct classes:
[kw_p_hist]»[chart]»[struct_class]

## Examples

M = kw_p_hist()
C = kw_p_hist()
kw_rank_sum Kruskal-Wallis rank sum test

## Description

The Kruskal-Wallis test is a univariate hypothesis testing method that allows multiple ( $\mathrm{n}>=2$ ) groups to be compared without making the assumption that values are normally distributed. It is the nonparametric equivalent of a 1-way ANOVA. The test is applied to all variables/features individually, and multiple test corrected p-values are computed to indicate the significance of variables/features.

## Usage

kw_rank_sum(alpha $=0.05$, mtc $\left.=" f d r ", f a c t o r \_n a m e s, \ldots\right)$

## Arguments

> alpha (numeric) The p-value cutoff for determining significance. The default is 0.05. mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
factor_names (character) The name of sample meta column(s) to use.
... Additional slots and values passed to struct_class.

## Value

A kw_rank_sum object with the following output slots:

```
test_statistic (data.frame) The value of the calculated statistic which is converted to a p-value when compared to a chi2-
p_value (data.frame) The probability of observing the calculated statistic.
dof (numeric) The number of degrees of freedom used to calculate the test statistic.
significant (data.frame) TRUE if the calculated p-value is less than the supplied threhold (alpha).
estimates (data.frame)
```


## Inheritance

A kw_rank_sum object inherits the following struct classes:
[kw_rank_sum] » [model] » [struct_class]

## Examples

```
M = kw_rank_sum(
    alpha = 0.05,
    mtc = "fdr",
    factor_names = "V1")
D = iris_DatasetExperiment()
M = kw_rank_sum(factor_names='Species')
M = model_apply(M,D)
```


## Description

Linear models can be used to carry out regression, single stratum analysis of variance and analysis of covariance.

## Usage

linear_model(formula, na_action = "na.omit", contrasts = list(), ...)

## Arguments

formula (formula) A symbolic description of the model to be fitted.
na_action (character) NA action. Allowed values are limited to the following:

- "na.omit": Incomplete cases are removed.
- "na.fail": An error is thrown if NA are present.
- "na.exclude": Incomplete cases are removed, and the output result is padded to the correct size using NA.
- "na. pass": Does not apply a linear model if NA are present.

The default is "na.omit".
contrasts (list) The contrasts associated with a factor. The default is list().
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- stats


## Value

A linear_model object with the following output slots:

| $l m$ | $(\operatorname{lm})$ The 1 lm object for this model_. |
| :--- | :--- |
| coefficients | (numeric) The coefficients for the fitted model_. |
| residuals | (numeric) The residuals for the fitted model_. |
| fitted_values | (numeric) The fitted values for the data used to train the model_. |
| predicted_values | (numeric) The predicted values for new data using the fitted model_. |
| r_squared | (numeric) The value of R Squared for the fitted model_. |
| adj_r_squared | (numeric) The value ofAdjusted R Squared for the fitted model_. |

## Inheritance

A linear_model object inherits the following struct classes:
[linear_model]» [model]» [struct_class]

## References

R Core Team (2023). R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/.

## Examples

```
M = linear_model(
    formula = y ~ x,
    na_action = "na.omit",
    contrasts = list())
D = iris_DatasetExperiment()
M = linear_model(formula = y~Species)
```

log_transform logarithm transform

## Description

A logarithmic transform is applied to all values in the data matrix.

## Usage

log_transform(base $=10, \ldots$ )

## Arguments

base (numeric) The base of the logarithm used for the transform. The default is 10.
... Additional slots and values passed to struct_class.

## Value

A log_transform object with the following output slots:
transformed (DatasetExperiment) A DatasetExperiment object containing the log transformed data.
struct object

## Inheritance

A log_transform object inherits the following struct classes:
[log_transform]» [model]»[struct_class]

## Examples

M = log_transform( base = 10)
$M=$ log_transform()
mean_centre Mean centre

## Description

The mean sample is subtracted from all samples in the data matrix. The features in the centred matrix all have zero mean.

## Usage

mean_centre(mode = "data", ...)

## Arguments

mode (character) Mode of action. Allowed values are limited to the following:

- "data": Centring is applied to the data block.
- "sample_meta": Centring is applied to the sample_meta block.
- "both": Centring is applied to both the data and the sample_meta blocks.

The default is "data".
... Additional slots and values passed to struct_class.

## Value

A mean_centre object with the following output slots:

| centred | (DatasetExperiment) |
| :--- | :--- |
| mean_data | (numeric) |
| mean_sample_meta | (numeric) |

## Inheritance

A mean_centre object inherits the following struct classes:
[mean_centre]» [preprocess]» [model]»[struct_class]

## Examples

$$
\begin{aligned}
& M=\text { mean_centre }( \\
& \text { mode }=\text { "data") } \\
& M=\text { mean_centre }()
\end{aligned}
$$

```
mean_of_medians Mean of medians
```


## Description

The data matrix is normalised by the mean of the median of each factor level.

## Usage

mean_of_medians(factor_name, ...)

## Arguments

$$
\begin{array}{ll}
\text { factor_name } & \text { (character) The name of a sample-meta column to use. } \\
\ldots & \text { Additional slots and values passed to struct_class. }
\end{array}
$$

## Value

A mean_of_medians object with the following output slots: transformed (DatasetExperiment) Data after the tranformation has been applied.

## Inheritance

A mean_of_medians object inherits the following struct classes:

```
[mean_of_medians]» [model] »[struct_class]
```


## Examples

```
M = mean_of_medians(
    factor_name = "V1")
D = iris_DatasetExperiment()
M = mean_of_medians(factor_name='Species')
M = model_apply(M,D)
```

```
mixed_effect Mixed effects model
```


## Description

A mixed effects model is an extension of ANOVA where there are both fixed and random effects.

## Usage

mixed_effect(alpha = 0.05, mtc $=$ "fdr", formula, ss_type = "marginal", ...)

## Arguments

alpha (numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
formula (formula) A symbolic description of the model to be fitted.
ss_type (character) Sum of squares type. Allowed values are limited to the following:

- "marginal": Type III sum of squares.
- "sequential": Type II sum of squares.

The default is "marginal".
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- nlme
- emmeans


## Value

A mixed_effect object with the following output slots:

```
f_statistic (data.frame) The value of the calculated statistic.
    p_value (data.frame) The probability of observing the calculated statistic if the null hypothesis is true.
    significant (data.frame) True/False indicating whether the p-value computed for each variable is less than the threshold.
```


## Inheritance

A mixed_effect object inherits the following struct classes:
[mixed_effect]» [ANOVA]» [model]» [stato]» [struct_class]

## References

Pinheiro J, Bates D, R Core Team (2023). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-164, https://CRAN.R-project.org/package=nlme.
Pinheiro JC, Bates DM (2000). Mixed-Effects Models in S and S-PLUS. Springer, New York. doi:10.1007/b98882 https://doi.org/10.1007/b98882.

Lenth R (2023). emmeans: Estimated Marginal Means, aka Least-Squares Means. R package version 1.9.0, https://CRAN.R-project.org/package=emmeans.
Fox J, Weisberg S (2019). An R Companion to Applied Regression, Third edition. Sage, Thousand Oaks CA. https://socialsciences.mcmaster.ca/jfox/Books/Companion/.

## Examples

```
M = mixed_effect(
        alpha = 0.05,
        mtc = "fdr",
        formula = y ~ x,
        ss_type = "marginal")
D = iris_DatasetExperiment()
D$sample_meta$id=rownames(D) # dummy id column
M = mixed_effect(formula = y~Species+ Error(id/Species))
M = model_apply(M,D)
```

model_apply, ANOVA, DatasetExperiment-method
Apply method

## Description

Applies method to the input DatasetExperiment

## Usage

```
## S4 method for signature 'ANOVA,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'HSD,DatasetExperiment'
model_apply(M, D)
    ## S4 method for signature 'mixed_effect,DatasetExperiment'
    model_apply(M, D)
```

```
## S4 method for signature 'HSDEM,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'classical_lsq,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'confounders_clsq,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'constant_sum_norm,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'corr_coef,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'split_data,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'equal_split,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'filter_smeta,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'fisher_exact,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'fold_change,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'fold_change_int,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'HCA,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'knn_impute,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'kw_rank_sum,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'log_transform,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'mean_of_medians,DatasetExperiment'
model_apply(M, D)
```

```
## S4 method for signature 'nroot_transform,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'pairs_filter,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'prop_na,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'rsd_filter,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'sb_corr,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'stratified_split,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'tSNE,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'ttest,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'vec_norm,DatasetExperiment'
model_apply(M, D)
## S4 method for signature 'wilcox_test,DatasetExperiment'
model_apply(M, D)
```


## Arguments

M a method object
D another object used by the first

## Value

Returns a modified method object

## Examples

```
M=model()
model_apply(M,DatasetExperiment())
```

```
model_predict,DFA,DatasetExperiment-method
    Model prediction
```


## Description

Apply a model using the input DatasetExperiment. Assumes the model is trained first.

## Usage

```
    ## S4 method for signature 'DFA,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'PCA,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'PLSR,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'PLSDA,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'autoscale,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'blank_filter,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'constant_sum_norm,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'dratio_filter,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'filter_by_name,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'filter_na_count,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'filter_smeta,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'glog_transform,DatasetExperiment'
    model_predict(M, D)
    ## S4 method for signature 'linear_model,DatasetExperiment'
```

```
model_predict(M, D)
## S4 method for signature 'mean_centre,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'mv_feature_filter,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'mv_sample_filter,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'OPLSR,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'OPLSDA,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'pareto_scale,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'pqn_norm,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'SVM,DatasetExperiment'
model_predict(M, D)
## S4 method for signature 'vec_norm,DatasetExperiment'
model_predict(M, D)
```


## Arguments

| M | a model object |
| :--- | :--- |
| D | a DatasetExperiment object |

## Value

Returns a modified model object

## Examples

```
M = example_model()
M = model_predict(M,iris_DatasetExperiment())
```

    model_reverse, autoscale, DatasetExperiment-method
                        Reverse preprocessing
    
## Description

Reverse the effect of a preprocessing step on a DatasetExperiment.

## Usage

```
## S4 method for signature 'autoscale,DatasetExperiment'
model_reverse(M, D)
## S4 method for signature 'mean_centre,DatasetExperiment'
model_reverse(M, D)
```


## Arguments

| M | a model object |
| :--- | :--- |
| D | a DatasetExperiment object |

## Value

Returns a modified DatasetExperiment object

## Examples

$M=$ example_model()
D = model_reverse(M,iris_DatasetExperiment())

```
model_train,DFA,DatasetExperiment-method
Train a model
```


## Description

Trains a model using the input DatasetExperiment

## Usage

\#\# S4 method for signature 'DFA,DatasetExperiment' model_train(M, D)
\#\# S4 method for signature 'PCA,DatasetExperiment'
model_train(M, D)
\#\# S4 method for signature 'PLSR,DatasetExperiment'
model_train(M, D)
\#\# S4 method for signature 'PLSDA,DatasetExperiment'
model_train(M, D)

```
## S4 method for signature 'autoscale,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'blank_filter,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'constant_sum_norm,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'dratio_filter,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'filter_by_name,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'filter_na_count,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'filter_smeta,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'glog_transform,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'linear_model,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'mean_centre,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'mv_feature_filter,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'mv_sample_filter,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'OPLSR,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'OPLSDA,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'pareto_scale,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'pqn_norm,DatasetExperiment'
model_train(M, D)
```

```
## S4 method for signature 'SVM,DatasetExperiment'
model_train(M, D)
## S4 method for signature 'vec_norm,DatasetExperiment'
model_train(M, D)
```


## Arguments

| M | a model object |
| :--- | :--- |
| D | a DatasetExperiment object |

## Value

Returns a modified model object

## Examples

M = example_model()
$M=$ model_train(M,iris_DatasetExperiment())

MTBLS79_DatasetExperiment
MTBLS79: Direct infusion mass spectrometry metabolomics dataset: a benchmark for data processing and quality control

## Description

Direct-infusion mass spectrometry (DIMS) metabolomics is an important approach for characterising molecular responses of organisms to disease, drugs and the environment. Increasingly large-scale metabolomics studies are being conducted, necessitating improvements in both bioanalytical and computational workflows to maintain data quality. This dataset represents a systematic evaluation of the reproducibility of a multi-batch DIMS metabolomics study of cardiac tissue extracts. It comprises of twenty biological samples (cow vs. sheep) that were analysed repeatedly, in 8 batches across 7 days, together with a concurrent set of quality control (QC) samples. Data are presented from each step of the workflow and are available in MetaboLights (https://www.ebi.ac.uk/metabolights/MTBLS79)

## Usage

MTBLS79_DatasetExperiment(filtered = FALSE)

## Arguments

filtered TRUE to load data with quality control filters already applied, or FALSE to load the unfiltered data. Default is FALSE. The raw data is available from (https://www.ebi.ac.uk/metabolights/MTBLS79) and as an R dataset in the pmp package, available on Bioconductor.

## Value

DatasetExperiment object

## Examples

```
D = MTBLS79_DatasetExperiment()
summary(D)
```

```
mv_boxplot Missing value boxplots
```


## Description

Boxplots of the number of missing values per sample/feature.

## Usage

```
    mv_boxplot(
        label_outliers = TRUE,
        by_sample = TRUE,
        factor_name,
        show_counts = TRUE,
    ...
    )
```


## Arguments

label_outliers (logical) Label outliers. Allowed values are limited to the following:

- "TRUE": Sample labels for potential outliers are displayed on the plot.
- "FALSE": Sample labels are not included on the plot.

The default is TRUE.
by_sample (logical) Plot by sample or by feature. Allowed values are limited to the following:

- "TRUE": Missing values are plotted per sample.
- "FALSE": Missing values are plotted per feature.

The default is TRUE.
factor_name (character) The name of a sample-meta column to use.
show_counts (logical) Show counts. Allowed values are limited to the following:

- "TRUE": The number of samples for each box is displayed.
- "FALSE": The number of samples for each box is not displayed.

The default is TRUE.
... Additional slots and values passed to struct_class.

## Value

A mv_boxplot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A mv_boxplot object inherits the following struct classes:
[mv_boxplot]» [chart]» [struct_class]

## Examples

```
M = mv_boxplot(
        label_outliers = FALSE,
        by_sample = FALSE,
        factor_name = "V1",
        show_counts = FALSE)
D = MTBLS79_DatasetExperiment()
C = mv_boxplot(factor_name='Class')
chart_plot(C,D)
```

mv_feature_filter Filter features by missing values

## Description

Removes features where the percentage of non-missing values falls below a threshold.

## Usage

mv_feature_filter(
threshold = 20,
qc_label = "QC",
method = "QC",
factor_name,
...
)

## Arguments

threshold (numeric) The minimum percentage of non-missing values. The default is 20.
qc_label (character) The label used to identify QC/group samples when using the "QC" (within a named group) filtering method. The default is "QC".
method (character) Filtering method. Allowed values are limited to the following:

- "within_all": Features are removed if the threshold for non-missing values is not met for all groups.
- "within_one": Features are removed if the threshold for non-missing values is not met for any group.
- "QC": Features are removed if the threshold for non-missing values is not met for the named group.
- "across": The filter is applied ignoring sample group.

The default is "QC".
factor_name (character) The name of a sample-meta column to use.
Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A mv_feature_filter object with the following output slots:
filtered (DatasetExperiment) A DatasetExperiment object containing the filtered data.
flags (data.frame) \% missing values and a flag indicating whether the sample was rejected. $0=$ rejected.

## Inheritance

A mv_feature_filter object inherits the following struct classes:
[mv_feature_filter]» [model]»[struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc.pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

```
M = mv_feature_filter(
    threshold = 20,
    qc_label = "QC",
    method = "QC",
    factor_name = "V1")
D = iris_DatasetExperiment()
M = mv_feature_filter(factor_name='Species',qc_label='versicolor')
M = model_apply(M,D)
```

```
mv_feature_filter_hist
Histogram of missing values per feature
```


## Description

A histogram of the proportion of missing values per feature.

## Usage

mv_feature_filter_hist(...)

## Arguments

$$
\ldots \quad \text { Additional slots and values passed to struct_class. }
$$

## Value

A mv_feature_filter_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A mv_feature_filter_hist object inherits the following struct classes:
[mv_feature_filter_hist]» [chart]» [struct_class]

## Examples

$M=$ mv_feature_filter_hist()
C = mv_feature_filter_hist()
mv_histogram Missing value histogram

## Description

A histogram of the numbers of missing values per sample/feature

## Usage

mv_histogram(label_outliers = TRUE, by_sample = TRUE, ...)

## Arguments

label_outliers (logical) Label outliers. Allowed values are limited to the following:

- "TRUE": Sample labels for potential outliers are displayed on the plot.
- "FALSE": Sample labels are not included on the plot.

The default is TRUE.
by_sample (logical) Plot by sample or by feature. Allowed values are limited to the following:

- "TRUE": Missing values are plotted per sample.
- "FALSE": Missing values are plotted per feature.

The default is TRUE.
.. additional slots and values passed to struct_class

## Value

A mv_histogram object. This object has no output slots. See chart_plot in the struct package to plot this chart object.
struct object

## Inheritance

A mv_histogram object inherits the following struct classes:
[mv_histogram]» [chart]» [struct_class]

## Examples

```
M = mv_histogram(
    label_outliers = FALSE,
    by_sample = FALSE)
D = MTBLS79_DatasetExperiment()
C = mv_histogram(label_outliers=FALSE,by_sample=FALSE)
chart_plot(C,D)
```

mv_sample_filter Missing value sample filter

## Description

Removes samples where the percent number of missing values exceeds a threshold.

## Usage

mv_sample_filter(mv_threshold = 20, ...)

## Arguments

mv_threshold (numeric) The maximum percentage of features with missing values in a sample. The default is 20 .
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A mv_sample_filter object with the following output slots:
filtered (DatasetExperiment) A DatasetExperiment object containing the filtered data. flags (data.frame) A flag indicating whether the sample was rejected. $0=$ rejected. percent_missing (data.frame) \% missing values for each sample.

## Inheritance

A mv_sample_filter object inherits the following struct classes:
[mv_sample_filter]» [model]» [struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc.pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

$$
\begin{gathered}
M=\text { mv_sample_filter }( \\
\text { mv_threshold }=20) \\
C=\text { mv_sample_filter }()
\end{gathered}
$$

```
mv_sample_filter_hist Histogram of missing values per sample
```


## Description

A histogram of the the proportion of missing values per sample

## Usage

mv_sample_filter_hist(...)

## Arguments

$$
\ldots \quad \text { Additional slots and values passed to struct_class. }
$$

## Value

A mv_sample_filter_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A mv_sample_filter_hist object inherits the following struct classes:
[mv_sample_filter_hist]» [chart]»[struct_class]

## Examples

M = mv_sample_filter_hist()
C = mv_sample_filter_hist()

```
nroot_transform nth root transform
```


## Description

All values in the data matrix are transformed by raising them to the power of $1 / n$.

## Usage

nroot_transform(root = 2, ...)
ontology_cache

## Arguments

root (numeric) The nth root used for the transform. The default is 2.
$\ldots \quad$ Additional slots and values passed to struct_class.

## Value

A nroot_transform object with the following output slots:
transformed (DatasetExperiment) A DatasetExperiment object containing the nth root transformed data.

## Inheritance

A nroot_transform object inherits the following struct classes:
[nroot_transform]» [model]» [struct_class]

## Examples

$$
M=\text { nroot_transform( }
$$

root = 2)
$M=$ nroot_transform()

```
ontology_cache ontology cache
```


## Description

A cached list of ontology terms obtained from the ontology lookup service (OLS) for ontology terms specified for objects in structToolbox.

## Usage

ontology_cache()

## Value

list of cached ontology terms

## See Also

> ontology

## Examples

```
cache = ontology_cache()
```

OPLSDA Orthogonal Partial Least Squares regression

## Description

OPLS splits a data matrix into two parts. One part contains information orthogonal to the input vector, and the other is non-orthogonal.

## Usage

OPLSDA(number_components = 1, factor_name, ...)

## Arguments

number_components
(numeric, integer) The number of orthgonal components. The default is 1.
factor_name (character) The name of a sample-meta column to use.
... Additional slots and values passed to struct_class.

## Value

A OPLSDA object with the following output slots:

```
opls_model (list)
filtered (DatasetExperiment)
orthogonal (DatasetExperiment)
```


## Inheritance

A OPLSDA object inherits the following struct classes:
[OPLSDA]» [OPLSR] » [model] » [struct_class]

## Examples

```
M = OPLSDA(
    number_components = 2,
    factor_name = "V1")
M = OPLSR('number_components'=2,factor_name='Species')
```

OPLSR
Orthogonal Partial Least Squares regression

## Description

OPLS splits a data matrix into two parts. One part contains information orthogonal to the input vector, and the other is non-orthogonal.

## Usage

OPLSR(number_components = 2, factor_name, ...)

## Arguments

number_components
(numeric, integer) The number of orthgonal components. The default is 2.
factor_name (character) The name of a sample-meta column to use.
... Additional slots and values passed to struct_class.

## Value

A OPLSR object with the following output slots:

```
opls_model (list)
filtered (DatasetExperiment)
orthogonal (DatasetExperiment)
```


## Inheritance

A OPLSR object inherits the following struct classes:

```
[OPLSR]» [model]»[struct_class]
```


## Examples

```
M = OPLSR(
    number_components = 2,
    factor_name = "V1")
M = OPLSR('number_components'=2,factor_name='Species')
```

```
pairs_filter Pairs filter
```


## Description

This filter is used for study designs with paired sampling to ensure that measurements from the same source (e.g. patient) are represented in all factor levels and interactions.

## Usage

pairs_filter(factor_name, sample_id, ...)

## Arguments

factor_name (character) The name of a sample-meta column to use.
sample_id (character) Name of sample meta column containing sample identifiers.
... Additional slots and values passed to struct_class.

## Value

A pairs_filter object with the following output slots:
filtered (DatasetExperiment) A DatasetExperiment object after the filter has been applied.
flags (data.frame) A data.frame indicating whether features were filtered from the DatasetExperiment.
struct object

## Inheritance

A pairs_filter object inherits the following struct classes:
[pairs_filter]» [model]» [struct_class]

## Examples

```
M = pairs_filter(
    factor_name = "V1",
    sample_id = "V1")
M=pairs_filter(factor_name='Class',sample_id='ids')
```

```
pareto_scale Pareto scaling
```


## Description

The mean sample is subtracted from all samples and then scaled by the square root of the standard deviation. The transformed data has zero mean.

## Usage

pareto_scale(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A pareto_scale object with the following output slots:

| scaled | (DatasetExperiment) |
| :--- | :--- |
| mean | (numeric) |
| sd | (numeric) |

## Inheritance

A pareto_scale object inherits the following struct classes:
[pareto_scale]» [model]» [struct_class]

## Examples

```
M = pareto_scale()
D = iris_DatasetExperiment()
M = pareto_scale()
M = model_train(M,D)
M = model_predict(M,D)
```


## Description

PCA is a multivariate data reduction technique. It summarises the data in a smaller number of Principal Components that maximise variance.

## Usage

PCA(number_components = 2, ...)

## Arguments

number_components
(numeric, integer) The number of Principal Components calculated. The default is 2 .
... Additional slots and values passed to struct_class.

## Value

A PCA object with the following output slots:

| scores | (DatasetExperiment) A matrix of PCA scores where each column corresponds to a Principal Component. |
| :--- | :--- |
| loadings | (data.frame) |
| eigenvalues | (data.frame) |
| ssx | (numeric) |
| correlation | (data.frame) |
| that | (DatasetExperiment) |

## Inheritance

A PCA object inherits the following struct classes:
[PCA]» [model]» [struct_class]

## Examples

```
M = PCAC
    number_components = 2)
```

```
pca_biplot PCA biplot
```


## Description

A scatter plot of the selected principal component scores overlaid with the corresponding principal component loadings.

## Usage

pca_biplot(
components $=c(1,2)$,
points_to_label = "none",
factor_name,
scale_factor = 0.95,
style = "points",
label_features = FALSE,
)

## Arguments

components (numeric) The principal components used to generate the plot. The default is $c(1,2)$.
points_to_label
(character) points_to_label. Allowed values are limited to the following:

- "none": No samples are labelled on the plot.
- "all": All samples are labelled on the plot.
- "outliers": Potential outliers are labelled on the plot.

The default is "none".
factor_name (character) The name of a sample-meta column to use.
scale_factor (numeric) The scaling factor applied to the loadings. The default is 0.95 .
style (character) Plot style. Allowed values are limited to the following:

- "points": Loadings and scores are plotted as a scatter plot.
- "arrows": The loadings are plotted as arrow vectors.

The default is "points".
label_features (logical) Add feature labels. Allowed values are limited to the following:

- "TRUE": Features are labelled.
- "FALSE": Features are not labelled.

The default is FALSE.
... Additional slots and values passed to struct_class.

## Value

A pca_biplot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pca_biplot object inherits the following struct classes:
[pca_biplot]»[chart]»[struct_class]

## Examples

```
M = pca_biplot(
            components = c(1, 2),
        points_to_label = "none",
        factor_name = "V1",
        scale_factor = 0.95,
        style = "points",
        label_features = FALSE)
    C = pca_biplot(factor_name='Species')
```

    pca_correlation_plot PCA correlation plot
    
## Description

A plot of the correlation between the variables/features and the selected principal component scores. Features with high correlation are well represented by the selected component(s)

## Usage

pca_correlation_plot(components = c(1, 2), ...)

## Arguments

components (numeric) The Principal Components used to generate the plot. The default is $c(1,2)$.
... Additional slots and values passed to struct_class.

## Value

A pca_correlation_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pca_correlation_plot object inherits the following struct classes:
[pca_correlation_plot]» [chart]» [struct_class]

## Examples

```
M = pca_correlation_plot(
    components = c(1, 2))
C = pca_correlation_plot()
```

```
pca_dstat_plot d-statistic plot
```


## Description

A bar chart of the d-statistics for samples in the input PCA model. Samples above the indicated threshold are considered to be outlying.

## Usage

pca_dstat_plot(number_components $=2$, alpha $=0.05, \ldots$ )

## Arguments

number_components
(numeric) The number of principal components to use. The default is 2.
alpha (numeric) A confidence threshold for rejecting samples based on the d-statistic. The default is 0.05 .
... Additional slots and values passed to struct_class.

## Value

A pca_dstat_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pca_dstat_plot object inherits the following struct classes:
[pca_dstat_plot]»[chart]» [struct_class]

## Examples

$$
\begin{aligned}
& M=\text { pca_dstat_plot }( \\
& \text { number_components }=2, \\
& \text { alpha }=0.95) \\
& C=\text { pca_dstat_plot }()
\end{aligned}
$$

```
pca_loadings_plot PCA loadings plot
```


## Description

A barchart (one component) or scatter plot (two components) of the selected principal component loadings.

## Usage

pca_loadings_plot(
components $=c(1,2)$, style = "points", label_features $=$ NULL,
)

## Arguments

components (numeric) The principal components used to generate the plot. The default is $c(1,2)$.
style (character) Plot style. Allowed values are limited to the following:

- "points": Loadings and scores are plotted as a scatter plot.
- "arrows": The loadings are plotted as arrow vectors.

The default is "points".
label_features (character, NULL) Feature labels. Allowed values are limited to the following:

- "character ()": A vector of labels for the features.
- "NULL": No labels.
- "row. names": Labels will be extracted from the column names of the data matrix.
The default is NULL.
... Additional slots and values passed to struct_class.


## Value

A pca_loadings_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pca_loadings_plot object inherits the following struct classes:
[pca_loadings_plot]»[chart]»[struct_class]

## Examples

M = pca_loadings_plot(

$$
\text { components }=c(1,2),
$$

style = "points",
label_features = NULL)

C = pca_loadings_plot()

```
pca_scores_plot PCA scores plot
```


## Description

Plots a 2d scatter plot of the selected components

## Usage

```
pca_scores_plot(
    xcol = "PC1",
    ycol = "PC2",
    points_to_label = "none",
    factor_name,
    ellipse = "all",
    ellipse_type = "norm",
    ellipse_confidence = 0.95,
    label_filter = character(0),
    label_factor = "rownames",
    label_size = 3.88,
    components = NULL,
    ...
)
```


## Arguments

xcol (numeric, integer, character) The column name, or index, of data to plot on the x -axis. The default is "PC1".
ycol (numeric, integer, character) The column name, or index, of data to plot on the $y$-axis. The default is "PC2".
points_to_label
(character) Points to label. Allowed values are limited to the following:

- "none": No samples labels are displayed.
- "all": The labels for all samples are displayed.
- "outliers": Labels for for potential outlier samples are displayed.

The default is "none".
$\begin{array}{ll}\text { factor_name } & \text { (character) The name of a sample-meta column to use. } \\ \text { ellipse } & \text { (character) Plot ellipses. Allowed values are limited to the following: }\end{array}$

- "all": Ellipses are plotted for all groups and all samples.
- "group": Ellipses are plotted for all groups.
- "none": Ellipses are not included on the plot.
- "sample": An ellipse is plotted for all samples (ignoring group).

The default is "all".
ellipse_type (character) Type of ellipse. Allowed values are limited to the following:

- "norm": Multivariate normal ( $\mathrm{p}=0.95$ ).
- "t": Multivariate $\mathrm{t}(\mathrm{p}=0.95)$.

The default is "norm".
ellipse_confidence
(numeric) The confidence level for plotting ellipses. The default is 0.95 .
label_filter (character) Labels are only plotted for the named groups. If zero-length then all groups are included. The default is character (0).
label_factor (character) The column name of sample_meta to use for labelling samples on the plot. "rownames" will use the row names from sample_meta. The default is "rownames".
label_size (numeric) The text size of labels. Note this is not in Font Units. The default is 3.88.
components (numeric, integer, NULL) The principal components used to generate the plot. If provided this parameter overrides xcol and ycol params. The default is NULL.
... Additional slots and values passed to struct_class.

## Value

A pca_scores_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pca_scores_plot object inherits the following struct classes:
[pca_scores_plot]» [scatter_chart]» [chart]» [struct_class]

## Examples

```
M = pca_scores_plot(
    components = NULL,
    xcol = 1,
    ycol = 2,
    points_to_label = "none",
    factor_name = "V1",
    ellipse = "all",
    label_filter = character(0),
    label_factor = "rownames",
    label_size = 3.88,
    ellipse_type = "norm",
    ellipse_confidence = 0.95)
D = iris_DatasetExperiment()
M = mean_centre() + PCA()
M = model_apply(M,D)
C = pca_scores_plot(factor_name = 'Species')
chart_plot(C,M[2])
```

    pca_scree_plot Scree plot
    
## Description

A plot of the percent variance and cumulative percent variance for the components of a PCA model.

## Usage

pca_scree_plot(max_pc = 15, ...)

## Arguments

max_pc (numeric, integer) The maximum number of components to include in the plot. The default is 15 .
... Additional slots and values passed to struct_class.

## Value

A pca_scree_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.
struct object

## Inheritance

A pca_scree_plot object inherits the following struct classes:
[pca_scree_plot]» [chart]» [struct_class]

## Examples

$$
\begin{gathered}
M=\text { pca_scree_plot }\left(\begin{array}{c}
\text { max_pc }=15) \\
C=
\end{array}\right. \\
\text { pca_scree_plot }()
\end{gathered}
$$

```
permutation_test Permutation test
```


## Description

A permutation test generates a "null" model by randomising the response (for regression models) or group labels (for classification models). This is repeated many times to generate a distribution of performance metrics for the null model. This distribution can then be compared to the performance of the true model. If there is overlap between the true and null model performances then the model is overfitted.

## Usage

permutation_test(number_of_permutations = 50, factor_name, ...)

## Arguments

number_of_permutations
(numeric, integer) The number of permutations. The default is 50 .
factor_name (character) The name of a sample-meta column to use.
... Additional slots and values passed to struct_class.

## Value

A permutation_test object with the following output slots:

```
results.permuted (data.frame)
results.unpermuted (data.frame)
metric (data.frame)
```


## Inheritance

A permutation_test object inherits the following struct classes:

```
[permutation_test]»[resampler]»[iterator]»[struct_class]
```


## Examples

```
M = permutation_test(
    number_of_permutations = 100,
    factor_name = "V1")
I=permutation_test(factor_name='Species')
```

```
permutation_test_plot permutation_test_plot class
```


## Description

Plots the results of a permutation test.

## Usage

permutation_test_plot(style = "boxplot", binwidth = 0.05, ...)

## Arguments

| style | The plot style. One of 'boxplot', 'violin', 'histogram', 'density' or 'scatter'. |
| :--- | :--- |
| binwidth | Binwidth for the "histogram" style. Ignored for all other styles. |
| $\ldots$ | additional slots and values passed to struct_class |

## Value

struct object

## Examples

C = permutation_test_plot(style='boxplot')

```
permute_sample_order Permute Sample Order
```


## Description

The order of samples in the data matrix is randomly permuted. The relationship between the samples and the sample meta data is maintained.

## Usage

permute_sample_order(number_of_permutations = 10, ...)

## Arguments

number_of_permutations
(numeric, integer) The number of times the sample order is permuted. The default is 10.
... Additional slots and values passed to struct_class.

## Value

A permute_sample_order object with the following output slots:

| results | (data.frame) |
| :--- | :--- |
| metric | (data.frame) |
| metric.train | (numeric) |

## Inheritance

A permute_sample_order object inherits the following struct classes:

```
[permute_sample_order]»[resampler]» [iterator]»[struct_class]
```


## Examples

M = permute_sample_order(
number_of_permutations = 100)
$C=$ permute_sample_order()
PLSDA Partial least squares discriminant analysis

## Description

PLS is a multivariate regression technique that extracts latent variables maximising covariance between the input data and the response. The Discriminant Analysis variant uses group labels in the response variable. For $>2$ groups a 1-vs-all approach is used. Group membership can be predicted for test samples based on a probability estimate of group membership, or the estimated y-value.

## Usage

PLSDA(number_components = 2, factor_name, pred_method = "max_prob", ...)

## Arguments

number_components
(numeric, integer) The number of PLS components. The default is 2 .
factor_name (character) The name of a sample-meta column to use.
pred_method (character) Prediction method. Allowed values are limited to the following:

- "max_yhat": The predicted group is selected based on the largest value of y_hat.
- "max_prob": The predicted group is selected based on the largest probability of group membership.

The default is "max_prob".
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pls


## Value

A PLSDA object with the following output slots:

```
scores (DatasetExperiment)
loadings (data.frame)
yhat (data.frame)
design_matrix (data.frame)
y (data.frame)
reg_coeff (data.frame)
probability (data.frame)
vip (data.frame)
pls_model (list)
pred (data.frame)
threshold (numeric)
sr
sr_pvalue
(data.frame) Selectivity ratio for a variable represents a measure of a variable's importance in the PLS mod (data.frame) A p-value computed from the Selectivity Ratio based on an F-distribution.
```


## Inheritance

A PLSDA object inherits the following struct classes:
[PLSDA]» [PLSR]»[model]» [struct_class]

## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.

Perez NF, Ferre J, Boque R (2009). "Calculation of the reliability of classification in discriminant partial least-squares binary classification." Chemometrics and Intelligent Laboratory Systems, 95(2), 122-128.
Barker M, Rayens W (2003). "Partial least squares for discrimination." Journal of Chemometrics, 17(3), 166-173.

## Examples

```
M = PLSDA(
    number_components = 2,
    factor_name = "V1",
    pred_method = "max_prob")
M = PLSDA('number_components'=2,factor_name='Species')
```

```
plsda_feature_importance_plot
    PLSDA feature importance summary plot
```


## Description

A plot of the selected feature significance metric for a PLSDA model for the top selected features.

## Usage

plsda_feature_importance_plot(n_features = 30, metric = "vip", ...)

## Arguments

$\mathrm{n}_{-}$features (numeric, integer) The number of features to include in the summary. The default is 30 .
metric (character) Metric to plot. Allowed values are limited to the following:

- "sr": Plot Selectivity Ratio.
- "sr_pvalue": Plot SR p-values.
- "vip": Plot Variable Importance in Projection scores.

The default is "vip".
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pls
- ggplot2
- reshape2
- cowplot


## Value

A plsda_feature_importance_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsda_feature_importance_plot object inherits the following struct classes:
[plsda_feature_importance_plot] » [chart] » [struct_class]

## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.
Wickham H (2016). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York. ISBN 978-3-319-24277-4, https://ggplot2.tidyverse.org.
Wickham H (2007). "Reshaping Data with the reshape Package." Journal of Statistical Software, 21(12), 1-20. http://www.jstatsoft.org/v21/i12/.

Wilke C (2023). cowplot: Streamlined Plot Theme and Plot Annotations for 'ggplot2'. R package version 1.1.2, https://CRAN.R-project.org/package=cowplot.

## Examples

```
M = plsda_feature_importance_plot(
            n_features = 50,
            metric = "vip")
D = iris_DatasetExperiment()
M = mean_centre()+PLSDA(factor_name='Species')
M = model_apply(M,D)
C = plsda_feature_importance_plot(n_features=30,metric='vip')
chart_plot(C,M[2])
```

```
plsda_predicted_plot PLSDA predicted plot
```


## Description

A plot of the regression coefficients from a PLSDA model.

## Usage

plsda_predicted_plot(factor_name, style = "boxplot", ycol = 1, ...)

## Arguments

| factor_name | (character) The name of a sample-meta column to use. |
| :--- | :--- |
| style | (character) Plot style. Allowed values are limited to the following: |

- "boxplot": A boxplot.
- "violin": A violin plot.
- "density": A density plot.

The default is "boxplot".
ycol (character, numeric, integer) The column of the Y block to be plotted. The default is 1 .
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pls
- ggplot2


## Value

A plsda_predicted_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsda_predicted_plot object inherits the following struct classes:
[plsda_predicted_plot]» [chart]» [struct_class]

## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.
Wickham H (2016). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York. ISBN 978-3-319-24277-4, https://ggplot2.tidyverse.org.

## Examples

```
M = plsda_predicted_plot(
        factor_name = "V1",
        style = "boxplot",
        ycol = 1)
D = iris_DatasetExperiment()
M = mean_centre()+PLSDA(factor_name='Species')
M = model_apply(M,D)
    C = plsda_predicted_plot(factor_name='Species')
    chart_plot(C,M[2])
```

    plsda_roc_plot PLSDA ROC plot
    
## Description

A Receiver Operator Characteristic (ROC) plot for PLSDA models computed by adjusting the threshold for assigning group labels from PLS predictions.

## Usage

plsda_roc_plot(factor_name, ycol = 1, ...)

## Arguments

factor_name (character) The name of a sample-meta column to use.
ycol (character, numeric, integer) The column of the Y block to be plotted. The default is 1 .
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pls
- ggplot2


## Value

A plsda_roc_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsda_roc_plot object inherits the following struct classes:
[plsda_roc_plot]» [chart]» [struct_class]

## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.
Wickham H (2016). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York. ISBN 978-3-319-24277-4, https://ggplot2. tidyverse.org.

## Examples

```
M = plsda_roc_plot(
        factor_name = "V1",
        ycol = 1)
D = iris_DatasetExperiment()
M = mean_centre()+PLSDA(factor_name='Species')
M = model_apply(M,D)
C = plsda_roc_plot(factor_name='Species')
chart_plot(C,M[2])
```

PLSR

Partial least squares regression

## Description

PLS is a multivariate regression technique that extracts latent variables maximising covariance between the input data and the response. For regression the response is a continuous variable.

## Usage

PLSR(number_components = 2, factor_name, ...)

## Arguments

number_components
(numeric, integer) The number of PLS components. The default is 2 .
factor_name (character) The name of sample meta column(s) to use.
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pls


## Value

A PLSR object with the following output slots:

```
scores (DatasetExperiment)
loadings (data.frame)
yhat (data.frame)
y (data.frame)
reg_coeff (data.frame)
vip (data.frame)
pls_model (list)
pred (data.frame)
sr (data.frame) Selectivity ratio for a variable represents a measure of a variable's importance in the PLS model. T
sr_pvalue (data.frame) A p-value computed from the Selectivity Ratio based on an F-distribution.
```


## Inheritance

A PLSR object inherits the following struct classes:
[PLSR]» [model]» [struct_class]

## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.

## Examples

```
M = PLSR(
    number_components = 2,
    factor_name = "V1")
M = PLSR(factor_name='run_order')
```

plsr_cook_dist Cook's distance barchart

## Description

A barchart of Cook's distance for each sample used to train a PLSR model. Cook's distance is used to estimate the influence of a sample on the model and can be used to identify potential outliers.

## Usage

plsr_cook_dist(ycol = 1, ...)

## Arguments

ycol (numeric, integer, character) The y-block column to plot. The default is 1.
... Additional slots and values passed to struct_class.

## Value

A plsr_cook_dist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsr_cook_dist object inherits the following struct classes:
[plsr_cook_dist]» [chart]» [struct_class]

## Examples

$$
\begin{gathered}
M=\text { plsr_cook_dist }\left(\begin{array}{c}
\text { ycol }=1) \\
C=\text { plsr_cook_dist }()
\end{array}\right.
\end{gathered}
$$

```
plsr_prediction_plot PLSR prediction plot
```


## Description

A scatter plot of the true response values against the predicted values for a PLSR model.

## Usage

plsr_prediction_plot(ycol = 1, ...)

## Arguments

ycol
(numeric, integer, character) The y-block column to plot. The default is 1 .
... Additional slots and values passed to struct_class.

## Value

A plsr_prediction_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsr_prediction_plot object inherits the following struct classes:
[plsr_prediction_plot]» [chart]» [struct_class]

## Examples

$$
\begin{gathered}
M=\text { plsr_prediction_plot( } \\
\text { ycol }=1) \\
C=\text { plsr_prediction_plot() }
\end{gathered}
$$

```
plsr_qq_plot PLSR QQ plot
```


## Description

A plot of the quantiles of the residuals from a PLSR model against the quantiles of a normal distribution.

## Usage

plsr_qq_plot(ycol = 1, ...)

## Arguments

ycol (numeric, integer, character) The y-block column to plot. The default is 1.

$$
\ldots \quad \text { Additional slots and values passed to struct_class. }
$$

## Value

A plsr_qq_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsr_qq_plot object inherits the following struct classes:
[plsr_qq_plot]» [chart]» [struct_class]

## Examples

$$
\begin{gathered}
M=\text { plsr_qq_plot }( \\
y c o l=1) \\
C=\text { plsr_qq_plot }()
\end{gathered}
$$

```
plsr_residual_hist PLSR residuals histogram
```


## Description

A histogram of the residuals for a PLSR model.

## Usage

plsr_residual_hist(ycol = 1, ...)

## Arguments

ycol (numeric, integer, character) The y-block column to plot. The default is 1.
... Additional slots and values passed to struct_class.

## Value

A plsr_residual_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A plsr_residual_hist object inherits the following struct classes:
[plsr_residual_hist]» [chart]» [struct_class]

## Examples

M = plsr_residual_hist( ycol = 1)

C = plsr_residual_hist()

```
pls_regcoeff_plot pls_regcoeff_plot class
```


## Description

Plots the regression coefficients of a PLSDA model.
Plots the regression coefficient scores of a PLSDA model

## Usage

pls_regcoeff_plot(ycol = 1, ...)

## Arguments

ycol (character, numeric, integer) The Y column to plot. The default is 1.
... additional slots and values passed to struct_class

## Details

This object makes use of functionality from the following packages:

- pls
- ggplot2


## Value

A pls_regcoeff_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.
struct object

## Inheritance

A pls_regcoeff_plot object inherits the following struct classes:

```
[pls_regcoeff_plot]»[chart]» [struct_class]
```


## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.
Wickham H (2016). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York. ISBN 978-3-319-24277-4, https://ggplot2. tidyverse. org.

## Examples

```
M = pls_regcoeff_plot(
    ycol = 1)
D = iris_DatasetExperiment()
M = mean_centre()+PLSDA(factor_name='Species')
M = model_apply(M,D)
C = pls_regcoeff_plot(ycol='setosa')
chart_plot(C,M[2])
```

```
pls_scores_plot PLSDA scores plot
```


## Description

A scatter plot of the selected PLSDA scores.

## Usage

```
pls_scores_plot(
    xcol = "LV1",
    ycol = "LV2",
    points_to_label = "none",
    factor_name,
    ellipse = "all",
    ellipse_type = "norm",
    ellipse_confidence = 0.95,
    label_filter = character(0),
    label_factor = "rownames",
    label_size = 3.88,
    components = NULL,
    ...
)
plsda_scores_plot(
    xcol = "LV1",
    ycol = "LV2",
    points_to_label = "none",
    factor_name,
    ellipse = "all",
    ellipse_type = "norm",
    ellipse_confidence = 0.95,
    label_filter = character(0),
    label_factor = "rownames",
    label_size = 3.88,
    components = NULL,
)
```


## Arguments

xcol (numeric, integer, character) The column name, or index, of data to plot on the x -axis. The default is "LV1".
ycol (numeric, integer, character) The column name, or index, of data to plot on the $y$-axis. The default is "LV2".
points_to_label
(character) Points to label. Allowed values are limited to the following:

- "none": No samples labels are displayed.
- "all": The labels for all samples are displayed.
- "outliers": Labels for for potential outlier samples are displayed.

The default is "none".
factor_name (character) The name of a sample-meta column to use
ellipse (character) Plot ellipses. Allowed values are limited to the following:

- "all": Ellipses are plotted for all groups and all samples.
- "group": Ellipses are plotted for all groups.
- "none": Ellipses are not included on the plot.
- "sample": An ellipse is plotted for all samples (ignoring group).

The default is "all".
ellipse_type (character) Type of ellipse. Allowed values are limited to the following:

- "norm": Multivariate normal ( $\mathrm{p}=0.95$ ).
- "t": Multivariate $\mathrm{t}(\mathrm{p}=0.95)$.

The default is "norm".
ellipse_confidence
(numeric) The confidence level for plotting ellipses. The default is 0.95 .
label_filter (character) Labels are only plotted for the named groups. If zero-length then all groups are included. The default is character (0).
label_factor (character) The column name of sample_meta to use for labelling samples on the plot. "rownames" will use the row names from sample_meta. The default is "rownames".
label_size (numeric) The text size of labels. Note this is not in Font Units. The default is 3.88 .
components (numeric, integer, NULL) The principal components used to generate the plot. If provided this parameter overrides xcol and ycol params. The default is NULL.
... Additional slots and values passed to struct_class.

## Value

A pls_scores_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pls_scores_plot object inherits the following struct classes:
[pls_scores_plot] » [scatter_chart] » [chart] » [struct_class]

## Examples

```
M = pls_scores_plot(
        components = NULL,
        xcol \(=1\),
        ycol = 2,
        points_to_label = "none",
    factor_name = "V1",
    ellipse = "all",
    label_filter = character(0),
    label_factor = "rownames",
    label_size = 3.88,
    ellipse_type = "norm",
    ellipse_confidence \(=0.95\) )
D = iris_DatasetExperiment()
\(M=\) mean_centre()+PLSDA(factor_name='Species')
M = model_apply (M,D)
C = pls_scores_plot(factor_name='Species')
chart_plot(C,M[2])
```

pls_vip_plot PLSDA VIP plot

## Description

A plot of the Variable Importance for Projection (VIP) scores for a PLSDA model.

## Usage

pls_vip_plot(threshold = 1, ycol = 1, ...)

## Arguments

threshold (numeric, integer) The threshold for indicating significant features. The default is 1 .
ycol (character, numeric, integer) The column of the Y block to be plotted. The default is 1 .
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pls
- ggplot2


## Value

A pls_vip_plot object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pls_vip_plot object inherits the following struct classes:
[pls_vip_plot]»[chart]»[stato]»[struct_class]

## References

Liland K, Mevik B, Wehrens R (2023). pls: Partial Least Squares and Principal Component Regression. R package version 2.8-3, https://CRAN.R-project.org/package=pls.
Wickham H (2016). ggplot2: Elegant Graphics for Data Analysis. Springer-Verlag New York. ISBN 978-3-319-24277-4, https://ggplot2.tidyverse.org.

## Examples

```
M = pls_vip_plot(
    threshold = 1,
    ycol = 1)
D = iris_DatasetExperiment()
M = mean_centre()+PLSDA(factor_name='Species')
M = model_apply(M,D)
C = pls_vip_plot(ycol='setosa')
chart_plot(C,M[2])
```

pqn_norm Probabilistic Quotient Normalisation (PQN)

## Description

PQN is used to normalise for differences in concentration between samples. It makes use of Quality Control (QC) samples as a reference. PQN scales by the median change relative to the reference in order to be more robust against changes caused by response to perturbation.

## Usage

pqn_norm(
qc_label = "QC",
factor_name,
qc_frac $=0$,
sample_frac = 0,
ref_method = "mean",
ref_mean = NULL,
)

## Arguments

| qc_label <br> factor_name | (character) The label used to identify QC samples. The default is "QC". (character) The name of a sample-meta column to use. |
| :---: | :---: |
| qc_frac | (numeric) A value between 0 and 1 to indicate the minimum proportion of QC samples a feature must be present in for it to be included when computing the reference. Default qc_frac $=0$. . The default is 0 . |
| sample_frac | (numeric) A value between 0 and 1 to indicate the minimum proportion of samples a feature must be present in for it to be considered when computing the normalisation coefficients. . The default is 0 . |
| ref_method | (character) Reference computation method. Allowed values are limited to the following: |
|  | - "mean": The reference is computed as the mean of the samples matching the qc_label input. <br> - "median": The reference is computed as the median of the samples matching the qc_label_input. |
|  | The default is "mean". |
| ref_mean | (numeric, NULL) A single sample to use as the reference for normalisation. If set to NULL then the reference will be computed based on the other input parameters (ref_mean, qc_label etc). . The default is NULL. |
|  | Additional slots and values passed to struct_class. |

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A pqn_norm object with the following output slots:
$\begin{array}{ll}\text { normalised } & \text { (DatasetExperiment) A DatasetExperiment object containing the normalised data. } \\ \text { coeff } & \text { (data.frame) The normalisation coefficients calculated by PQN. }\end{array}$

## Inheritance

A pqn_norm object inherits the following struct classes:
[pqn_norm] » [model] » [struct_class]

## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc. pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

```
M = pqn_norm(
        qc_label = "QC",
        factor_name = "V1",
        qc_frac = 0,
        sample_frac = 0,
        ref_mean = NULL,
        ref_method = "mean")
D = iris_DatasetExperiment()
M = pqn_norm(factor_name='Species',qc_label='all')
M = model_apply(M,D)
```

    pqn_norm_hist PQN coefficient histogram
    
## Description

A histogram of the PQN coefficients for all features

## Usage

pqn_norm_hist(...)

## Arguments

$$
\ldots \quad \text { Additional slots and values passed to struct_class. }
$$

## Value

A pqn_norm_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A pqn_norm_hist object inherits the following struct classes:
[pqn_norm_hist]» [chart]»[struct_class]

## Examples

M = pqn_norm_hist()
C = pqn_norm_hist()
prop_na Fisher's exact test for missing values

## Description

A Fisher's exact test is used to compare the number of missing values in each group. Multiple test corrected p-values are computed to indicate whether there is a significant difference in the number of missing values across groups for each feature.

## Usage

prop_na(alpha $=0.05$, mtc $=" f d r "$, factor_name, ...)

## Arguments

> alpha (numeric) The p-value cutoff for determining significance. The default is 0.05. mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
factor_name (character) The name of a sample-meta column to use.
... Additional slots and values passed to struct_class.

## Value

A prop_na object with the following output slots:

| p_value | (data.frame) The probability of observing the calculated statistic. |
| :--- | :--- |
| significant | (data.frame) TRUE if the calculated p-value is less than the supplied threshold (alpha). |
| na_count | (data.frame) The number of NA values per group of the chosen factor. |

struct object

## Inheritance

A prop_na object inherits the following struct classes:
[prop_na]» [model]» [struct_class]

## Examples

```
M = prop_na(
        alpha = 0.05,
        mtc = "fdr",
        factor_name = "V1")
M = prop_na(factor_name='Species')
```

    resample Data resampling
    
## Description

New training sets are generated from the original data by selecting samples at random. This can be based on levels in a factor or on the whole dataset.

## Usage

```
resample(
    number_of_iterations = 10,
    method = "split_data",
    factor_name,
    p_train = 0.8,
    collect = NULL,
    )
```


## Arguments

number_of_iterations
(numeric, integer) The number of training sets to generate. The default is 10.
method (character) Resampling method. Allowed values are limited to the following:

- "split_data": Samples for the training set are selected at random from the full dataset.
- "stratified_split": Samples for the training set are randomly selected from each level of the chosen factor.
- "equal_split": Samples for the training set are selected at random from each level of the main factor such that all group sizes are equal.

The default is "split_data".
factor_name
(character) The name of a sample-meta column to use.
p_train (numeric) The proportion of samples selected for the training set. The default is 0.8 .
collect
(NULL, character) The name of a model output to collect over all bootstrap repetitions, in addition to the input metric. The default is NULL.
... Additional slots and values passed to struct_class.

## Value

A resample object with the following output slots:

| results.training | (data.frame) |
| :--- | :--- |
| results.testing | (data.frame) |
| metric | (data.frame) |
| collected | (list) |
| metric.train | (numeric) |
| metric.test | (numeric) |

## Inheritance

A resample object inherits the following struct classes:
[resample]»[resampler]» [iterator]» [struct_class]

## Examples

```
M = resample(
            number_of_iterations = 100,
            method = "split_data",
            factor_name = "V1",
            p_train = 0.75,
            collect = NULL)
I = resample(
    number_of_iterations = 10,
    factor_name = 'Species',
    method = 'split_data',
    p_train = 0.8)
```

```
resample_chart resample_chart class
```


## Description

Plots the results of a resampling.

## Usage

resample_chart(style = "boxplot", binwidth = 0.05, ...)

## Arguments

style The plot style. One of 'boxplot', 'violin', 'histogram', 'density' or 'scatter'.
binwidth Binwidth for the "histogram" style. Ignored for all other styles.
additional slots and values passed to struct_class

## Value

struct object

## Examples

```
    C = resample_chart(style='boxplot')
```

    rsd_filter \(\quad\) RSD filter
    
## Description

An RSD filter calculates the relative standard deviation (the ratio of the standard deviation to the mean) for all features. Any feature with an RSD greater than a predefined threshold is excluded.

## Usage

rsd_filter(rsd_threshold = 20, qc_label = "QC", factor_name, ...)

## Arguments

rsd_threshold (numeric) The RSD threshold above which features are removed. The default is 20.
qc_label (character) The label used to identify QC samples. The default is "QC".
factor_name (character) The name of a sample-meta column to use.
Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A rsd_filter object with the following output slots:
filtered (DatasetExperiment) A DatasetExperiment object containing the filtered data.
flags (data.frame) RSD and a flag indicating whether the feature was rejected by the filter or not. rsd_qc (data.frame) The calculated RSD of the QC class.

## Inheritance

A rsd_filter object inherits the following struct classes:

```
[rsd_filter]» [model]»[struct_class]
```


## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc. pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

## Examples

```
M = rsd_filter(
    rsd_threshold = 20,
    qc_label = "QC",
    factor_name = "V1")
M = rsd_filter(factor_name='Class')
```

    rsd_filter_hist RSD histogram
    
## Description

A histogram of the calculated RSD values.

## Usage

rsd_filter_hist(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A rsd_filter_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A rsd_filter_hist object inherits the following struct classes:
[rsd_filter_hist] » [chart]» [struct_class]

## Examples

M = rsd_filter_hist()
C = rsd_filter_hist()
run, bootstrap, DatasetExperiment, metric-method
Runs an iterator, applying the chosen model multiple times.

## Description

Running an iterator will apply the iterator a number of times to a DatasetExperiment. For example, in cross-validation the same model is applied multiple times to the same data, splitting it into training and test sets. The input metric object can be calculated and collected for each iteration as an output.

## Usage

```
    ## S4 method for signature 'bootstrap,DatasetExperiment,metric'
    run(I, D, MET = NULL)
    ## S4 method for signature 'forward_selection_by_rank,DatasetExperiment,metric'
    run(I, D, MET)
    ## S4 method for signature 'grid_search_1d,DatasetExperiment,metric'
    run(I, D, MET)
    ## S4 method for signature 'kfold_xval,DatasetExperiment,metric'
    run(I, D, MET = NULL)
    ## S4 method for signature 'permutation_test,DatasetExperiment,metric'
```

```
run(I, D, MET = NULL)
## S4 method for signature 'permute_sample_order,DatasetExperiment,metric'
run(I, D, MET)
## S4 method for signature 'resample,DatasetExperiment,metric'
run(I, D, MET)
```


## Arguments

| I | an iterator object |
| :--- | :--- |
| D | a DatasetExperiment object |
| MET | a metric object |

## Value

Modified iterator object

## Examples

```
D = iris_DatasetExperiment() # get some data
MET = metric() # use a metric
I = example_iterator() # initialise iterator
models(I) = example_model() # set the model
I = run(I,D,MET) # run
```

$r$ _squared $\quad$ Coefficient of determination ( $R$-squared)

## Description

R -squared is a metric used to assess the goodness of fit for regression models. It measures how much variance of one variable can be explained by another variable.

## Usage

r_squared(...)

## Arguments

$$
\ldots \quad \text { Additional slots and values passed to struct_class. }
$$

## Value

A r_squared object. This object has no output slots.

## Inheritance

A r_squared object inherits the following struct classes:
[r_squared]» [metric]» [struct_class]

## Examples

M = r_squared()
MET $=$ r_squared()
sb_corr Signal/batch correction for mass spectrometry data

## Description

Applies Quality Control Robust Spline (QC-RSC) method to correct for signal drift and batch differences in mass spectrometry data.

## Usage

sb_corr
order_col, batch_col, qc_col, smooth $=0$, use_log = TRUE, min_qc $=4$, qc_label = "QC", spar_lim $=c(-1.5,1.5)$, ...
)

## Arguments

order_col (character) The column name of sample_meta indicating the run order of the samples.
batch_col (character) The column name of sample_meta indicating the batch each sample was measured in.
qc_col (character) The column name of sample_meta indicating the group each sample is a member of.
smooth (numeric) The amount of smoothing applied (0 to 1). If set to 0 the smoothing parameter will be estimated using leave-one-out cross-validation. The default is 0.
use_log (logical) Log tranformation. Allowed values are limited to the following:

- "TRUE": The data is log transformed prior to performing signal correction.
The default is TRUE.
min_qc
(numeric) The minimum number of QC samples required for signal correction.
The default is 4.

spar_lim | (character) The label used to identify QC samples. The default is "QC". |
| :--- |
| (numeric) A two element vector specifying the upper and lower limits when |
| spar = 0 . Allows the value of spar to be constrained within these limits to |
| prevent overfiting. The default is c $(-1.5,1.5)$. |

## Details

This object makes use of functionality from the following packages:

- pmp


## Value

A sb_corr object with the following output slots:
corrected (DatasetExperiment) The DatasetExperiment after signal/batch correction has been applied. fitted (data.frame) The fitted splines for each feature.
struct object

## Inheritance

A sb_corr object inherits the following struct classes:

```
[sb_corr]»[model]»[struct_class]
```


## References

Jankevics A, Lloyd GR, Weber RJM (2023). pmp: Peak Matrix Processing and signal batch correction for metabolomics datasets. doi:10.18129/B9.bioc.pmp https://doi.org/10.18129/B9. bioc.pmp, R package version 1.14.0, https://bioconductor.org/packages/pmp.

Kirwan JA, Broadhurst DI, Davidson RL, Viant MR (2013). "Characterising and correcting batch variation in an automated direct infusion mass spectrometry (DIMS) metabolomics workflow." Analytical and Bioanalytical Chemistry, 405(15), 5147-5157.

## Examples

M = sb_corr (
order_col = character(0),
batch_col = character(0),
qc_col = character(0),
smooth $=0$,
use_log = FALSE,
min_qc $=4$,
qc_label = "QC",
spar_lim = c(-1.5, 1.5))
M = sb_corr(order_col='run_order', batch_col='batch_no',qc_col='class')

```
scatter_chart Group scatter chart
```


## Description

Plots a 2d scatter plot of the input data.

## Usage

```
scatter_chart(
        xcol = 1,
        ycol = 2,
        points_to_label = "none",
        factor_name = "none",
        ellipse = "all",
        ellipse_type = "norm",
        ellipse_confidence = 0.95,
        label_filter = character(0),
        label_factor = "rownames",
        label_size = 3.88,
    )
```


## Arguments

xcol (numeric, integer, character) The column name, or index, of data to plot on the x -axis. The default is 1 .
ycol (numeric, integer, character) The column name, or index, of data to plot on the $y$-axis. The default is 2 .
points_to_label
(character) Points to label. Allowed values are limited to the following:

- "none": No samples labels are displayed.
- "all": The labels for all samples are displayed.
- "outliers": Labels for for potential outlier samples are displayed.

The default is "none".


## Value

A scatter_chart object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A scatter_chart object inherits the following struct classes:
[scatter_chart]» [chart]» [struct_class]

## Examples

```
M = scatter_chart(
    xcol = 1,
    ycol = 2,
    points_to_label = "none",
    factor_name = "V1",
```

```
        ellipse = "all",
        label_filter = character(0),
        label_factor = "rownames",
        label_size = 3.88,
        ellipse_type = "norm",
        ellipse_confidence = 0.95)
    D = iris_DatasetExperiment()
C = scatter_chart(
        xcol = 'Petal.Width',
        ycol = 'Sepal.Width',
        factor_name = 'Species'
        )
    chart_plot(C,D)
```

    split_data Split data
    
## Description

The data matrix is divided into two subsets.A predefined proportion of the samples are randomly selected for a training set, and the remaining samples are used for the test set.

## Usage

split_data(p_train, ...)

## Arguments

p_train (numeric) The proportion of samples selected for the training set.
... Additional slots and values passed to struct_class.

## Value

A split_data object with the following output slots:
training (DatasetExperiment) A DatasetExperiment object containing samples selected for the training set.
testing (DatasetExperiment) A DatasetExperiment object containing samples selected for the testing set.

## Inheritance

A split_data object inherits the following struct classes:
[split_data]» [model]» [struct_class]

## Examples

```
M = split_data(
        p_train = 0.75)
M = split_data(p_train=0.75)
```

stratified_split Stratified sampling

## Description

The dataset is divided into two subsets. A predefined proportion of samples from each level of a factor is selected for the training set, and the remaining samples are used for the test set. The stratification by factor level means that the relative number of samples per level is approximately equal to the original dataset.

## Usage

```
stratified_split(p_train, factor_name, ...)
```


## Arguments

| p_train | (numeric) The proportion of samples selected for the training set. |
| :--- | :--- |
| factor_name | (character) The name of a sample-meta column to use. |
| $\ldots$ | Additional slots and values passed to struct_class. |

## Value

A stratified_split object with the following output slots:
training (DatasetExperiment) A DatasetExperiment object containing samples selected for the training set.
testing (DatasetExperiment) A DatasetExperiment object containing samples selected for the testing set.

## Inheritance

A stratified_split object inherits the following struct classes:
[stratified_split]»[split_data]» [model]» [struct_class]

## Examples

```
M = stratified_split(
        factor_name = "V1",
        p_train = 0.75)
D = iris_DatasetExperiment()
M = stratified_split(p_train=0.75,factor_name='Species')
M = model_apply(M,D)
```

structToolbox
structToolbox: Examples of tools built using the Statistics in $R$ Using Class Templates (struct) package

## Description

This package extends the classes defined in the struct package

## SVM

Support Vector Machine Classifier

## Description

Support Vector Machines (SVM) are a machine learning algorithm for classification. They can make use of kernel functions to generate highly non-linear boundaries between groups.

## Usage

## SVM(

factor_name, kernel = "linear", degree $=3$,
gamma $=1$,
coef0 = 0,
cost = 1,
class_weights = NULL,
)

## Arguments

\(\left.$$
\begin{array}{ll}\text { factor_name } \\
\text { kernel } & \begin{array}{l}\text { (character) The name of a sample-meta column to use. } \\
\text { (character) Kernel type. Allowed values are limited to the following: } \\
\text { - "linear": }\end{array}
$$ <br>
- "polynomial": <br>
- "radial": <br>

- "sigmoid":\end{array}\right]\)| The default is "linear". |
| :--- |
| (numeric) The polynomial degree. The default is 3. |

## Details

This object makes use of functionality from the following packages:

- e1071


## Value

A SVM object with the following output slots:

| SV | (matrix) |
| :--- | :--- |
| index | (numeric) |
| coefs | (matrix) |
| pred | (data.frame) |
| decision_values | (data.frame) |

struct object

## Inheritance

A SVM object inherits the following struct classes:
[SVM]» [model]» [struct_class]

## References

Meyer D, Dimitriadou E, Hornik K, Weingessel A, Leisch F (2023). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-14, https://CRAN.R-project.org/package=e1071.
Brereton RG, Lloyd GR (2010). "Support Vector Machines for classification and regression." The Analyst, 135(2), 230-267.

## Examples

$$
M=S V M C
$$

factor_name = "V1",
kernel = "linear",

$$
\text { degree }=3 \text {, }
$$

gamma = 1,

$$
\operatorname{coef0}=0,
$$

$$
\cos t=1
$$

class_weights = 1)

M = SVM(factor_name='Species',gamma=1)

```
    svm_plot_2d SVM scatter plot
```


## Description

A scatter plot of the input data by group and the calculated boundary of a SVM model.

## Usage

svm_plot_2d(factor_name, npoints = 100, ...)

## Arguments

factor_name (character) The name of a sample-meta column to use.
npoints (numeric) The number of grid points used to plot the boundary. The default is 100.
... Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- e1071


## Value

A svm_plot_2d object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A svm_plot_2d object inherits the following struct classes:
[svm_plot_2d] » [chart]»[struct_class]

## References

Meyer D, Dimitriadou E, Hornik K, Weingessel A, Leisch F (2023). e1071: Misc Functions of the Department of Statistics, Probability Theory Group (Formerly: E1071), TU Wien. R package version 1.7-14, https://CRAN.R-project.org/package=e1071.

## Examples

```
M = svm_plot_2d(
            factor_name = "V1",
            npoints = 100)
D = iris_DatasetExperiment()
M = filter_smeta(mode='exclude',levels='setosa',factor_name='Species') +
    mean_centre()+PCA(number_components=2)+
    SVM(factor_name='Species',kernel='linear')
M = model_apply(M,D)
C = svm_plot_2d(factor_name='Species')
chart_plot(C,M[4],predicted(M[3]))
```


## tic_chart <br> Total Ion Count chart.

## Description

A scatter plot of Total Ion Count (sum of each sample) versus run order.

## Usage

tic_chart(run_order, factor_name, connected = FALSE, ...)

## Arguments

run_order (character) The column name of sample_meta indicating the run order of the samples.
factor_name (character) The name of a sample-meta column to use.
connected (logical) Plot samples connected by a grey line. The default is FALSE.
... Additional slots and values passed to struct_class.

## Value

A tic_chart object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A tic_chart object inherits the following struct classes:
[tic_chart]» [chart] » [struct_class]

## Examples

```
M = tic_chart(
        factor_name = "V1",
    run_order = character(0),
    connected = FALSE)
D = iris_DatasetExperiment()
D$sample_meta$run_order=1:nrow(D)
C = tic_chart(factor_name='Species',run_order='run_order')
chart_plot(C,D)
```

    tSNE
    \(t S N E\)
    
## Description

t-Distributed Stochastic Neighbor Embedding.

## Usage

tSNE (
dims $=2$,
perplexity $=30$,
max_iter = 100,
theta $=0.5$,
check_duplicates = FALSE,
init $=$ NULL,
eta $=200$,
...
)

## Arguments

dims (numeric) The number of tSNE dimensions computed. The default is 2.

```
perplexity (numeric) Perplexity parameter. The default is 30.
max_iter (numeric) The maximum number of tSNE iterations. The default is 100.
theta (numeric) Speed/accuracy trade-off. A value of 0 gives an exact tSNE. The de-
        fault is 0.5.
check_duplicates
    (logical) Check for duplicates. Allowed values are limited to the following:
    - "TRUE": Checks for the presence of exact duplicate samples.
                            - "FALSE": Does not check for exact duplicate samples.
        The default is FALSE.
    init (NULL, data.frame, DatasetExperiment) A set of coordinates for initialising the
        tSNE algorithm. NULL uses random initialisation. The default is NULL.
eta (numeric) The learning rate parameter. The default is 200.
    ... Additional slots and values passed to struct_class.
```


## Details

This object makes use of functionality from the following packages:

- Rtsne


## Value

A tSNE object with the following output slots:
Y (DatasetExperiment)

## Inheritance

A tSNE object inherits the following struct classes:
[tSNE]» [model]» [struct_class]

## References

Krijthe JH (2015). Rtsne: T-Distributed Stochastic Neighbor Embedding using Barnes-Hut Implementation. R package version 0.17 , https://github.com/jkrijthe/Rtsne.
van der Maaten L, Hinton G (2008). "Visualizing High-Dimensional Data Using t-SNE." Journal of Machine Learning Research, 9, 2579-2605.
van der Maaten L (2014). "Accelerating t-SNE using Tree-Based Algorithms." Journal of Machine Learning Research, 15, 3221-3245.

## Examples

```
    M = tSNE(
    dims = 2,
    perplexity = 30,
    max_iter = 1000,
    theta = 0.5,
    check_duplicates = FALSE,
    init = NULL,
    eta = 200)
M = tSNE()
```

tSNE_scatter Feature boxplot

## Description

plots the new representation of data after applying tSNE.

## Usage

tSNE_scatter(factor_name, ...)

## Arguments

factor_name (character) The name of a sample-meta column to use.
Additional slots and values passed to struct_class.

## Details

This object makes use of functionality from the following packages:

- Rtsne


## Value

A tSNE_scatter object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A tSNE_scatter object inherits the following struct classes:
[tSNE_scatter]» [chart]» [struct_class]

## References

Krijthe JH (2015). Rtsne: T-Distributed Stochastic Neighbor Embedding using Barnes-Hut Implementation. R package version 0.17, https://github.com/jkrijthe/Rtsne.
van der Maaten L, Hinton G (2008). "Visualizing High-Dimensional Data Using t-SNE." Journal of Machine Learning Research, 9, 2579-2605.
van der Maaten L (2014). "Accelerating t-SNE using Tree-Based Algorithms." Journal of Machine Learning Research, 15, 3221-3245.

## Examples

```
M = tSNE_scatter(
    factor_name = "V1")
M = tSNE_scatter(factor_name='Species')
```

ttest t-test

## Description

A t-test compares the means of two factor levels. Multiple-test corrected p-values are used to indicate the significance of the computed difference for all features.

## Usage

ttest
alpha $=0.05$,
mtc = "fdr",
factor_names,
paired = FALSE,
paired_factor = character(0),
equal_variance = FALSE,
conf_level = 0.95,
...
)

## Arguments

alpha (numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.

```
    - "none": No correction.
The default is "fdr".
factor_names (character) The name of sample meta column(s) to use.
paired (logical) Apply a paired t-test. The default is FALSE.
paired_factor (character) The factor name that encodes the sample id for pairing. The default
is character(0).
equal_variance (logical) Equal variance. Allowed values are limited to the following:
    - "TRUE": The variance of each group is treated as being equal using the
        pooled variance to estimate the variance.
    - "FALSE": The variance of each group is not assumed to be equal and the
    Welch (or Satterthwaite) approximation is used.
The default is FALSE.
conf_level (numeric) The confidence level of the interval. The default is 0.95.
... Additional slots and values passed to struct_class.
```


## Value

A ttest object with the following output slots:
t_statistic (data.frame) The value of the calculate statistics which is converted to a p-value when compared to at-distribu p_value (data.frame) The probability of observing the calculated $t$-statistic.
dof (numeric) The number of degrees of freedom used to calculate the test statistic.
significant (data.frame) TRUE if the calculated p-value is less than the supplied threhold (alpha).
conf_int (data.frame) Confidence interval for $t$ statistic.
estimates (data.frame) The group means estimated when computing the $t$-statistic.

## Inheritance

A ttest object inherits the following struct classes:
[ttest]» [model]» [struct_class]

## Examples

```
M = ttest(
        alpha = 0.05,
        mtc = "fdr",
        factor_names = "V1",
        paired = FALSE,
        paired_factor = "NA",
        equal_variance = FALSE,
        conf_level = 0.95)
M = ttest(factor_name='Class')
```

vec_norm Vector normalisation

## Description

The samples in the data matrix are normalised to account for differences in concentration by scaling each sample such that the sum of squares is equal to 1 .

## Usage

vec_norm(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A vec_norm object with the following output slots:
normalised (DatasetExperiment) A DatasetExperiment object containing the normalised data. coeff (data.frame) The normalisation coefficients calculated by PQN.
struct object

## Inheritance

A vec_norm object inherits the following struct classes:
[vec_norm] » [model] » [struct_class]

## Examples

M = vec_norm()
M = vec_norm()

```
wilcox_p_hist Histogram of p values
```


## Description

A histogram of $p$ values for the wilcoxon signed rank test

## Usage

wilcox_p_hist(...)

## Arguments

... Additional slots and values passed to struct_class.

## Value

A wilcox_p_hist object. This object has no output slots. See chart_plot in the struct package to plot this chart object.

## Inheritance

A wilcox_p_hist object inherits the following struct classes:
[wilcox_p_hist]» [chart]»[struct_class]

## Examples

M = wilcox_p_hist()
M = wilcox_p_hist()

```
wilcox_test wilcoxon signed rank test
```


## Description

A Mann-Whitney-Wilcoxon signed rank test compares ,the ranks of values in two groups. It is the non-parametric equivalent of a t-test. Multiple test corrected $p$-values are computed as indicators of significance for each variable/feature.

## Usage

```
wilcox_test(
    alpha = 0.05,
    mtc = "fdr",
    factor_names,
    paired = FALSE,
    paired_factor = character(0),
    conf_level = 0.95,
    ...
)
```


## Arguments

## alpha

(numeric) The p-value cutoff for determining significance. The default is 0.05 .
mtc (character) Multiple test correction method. Allowed values are limited to the following:

- "bonferroni": Bonferroni correction in which the p-values are multiplied by the number of comparisons.
- "fdr": Benjamini and Hochberg False Discovery Rate correction.
- "none": No correction.

The default is "fdr".
factor_names (character) The name of a sample-meta column to use.
paired (logical) Apply a paired test. The default is FALSE.
paired_factor (character) The factor name containing sample ids for paired data. The default is character (0).
conf_level (numeric) The confidence level of the interval. The default is 0.95.
... Additional slots and values passed to struct_class.

## Value

A wilcox_test object with the following output slots:

```
statistic (data.frame) The value of the calculated statistic which is converted to a p-value.
p_value (data.frame) The probability of observing the calculated t-statistic.
dof (numeric) The number of degrees of freedom used to calculate the test statistic.
significant (data.frame) TRUE if the calculated p-value is less than the supplied threhold (alpha).
conf_int (data.frame) Confidence interval for t statistic.
estimates (data.frame) The group estimates used when computing the statistic.
```

struct object
wilcox_test

Inheritance
A wilcox_test object inherits the following struct classes:
[wilcox_test]»[model]»[struct_class]

## Examples

M = wilcox_test alpha $=0.05$, mtc = "fdr", factor_names = "V1", paired = FALSE,
paired_factor = character(0), conf_level = 0.95)

M = wilcox_test(factor_name='Class')

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