

Package ‘oneway’

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Title One-Way Statistical Analyses

Version 0.0.2

Description Performs one-way tests of assumptions (normality and homoscedasticity), analysis of variance, robust and nonparametric alternatives, multiple comparison procedures, effect size estimators, confidence intervals, and descriptive summaries. Functions are designed with a consistent interface to support reproducible and user-friendly statistical workflows. For more details see Howell (2010, ISBN:978-0-495-59784-1), Zar (2014, ISBN:978-0-13-100846-5), Hollander et al. (2014, ISBN:978-0-470-38737-5), Montgomery (2017, ISBN:978-1-119-11347-8), Lakens (2013) <[doi:10.3389/fpsyg.2013.00863](https://doi.org/10.3389/fpsyg.2013.00863)>, and Piepho (2004) <[doi:10.1198/1061860043515](https://doi.org/10.1198/1061860043515)>.

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URL <https://github.com/P10911004-NPUST/oneway>

BugReports <https://github.com/P10911004-NPUST/oneway/issues>

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anorexia	<i>Weight Gain Following Therapy for Anorexia</i>
----------	---

Description

A dataset containing weight gain for young girls with anorexia following three different treatment conditions: a control group, cognitive-behavior therapy, and family therapy. The response variable is weight gain measured after treatment.

Usage

anorexia

Format

A data frame with 72 observations and 2 variables:

therapy A factor indicating the treatment group with three levels: "control", "cognitive", and "family".

weight_gain A numeric vector giving the observed weight gain after treatment.

Group sample sizes are:

- Control: 26

- Cognitive-behavior therapy: 29
- Family therapy: 17

References

Howell, D. C. (2013). Statistical methods for psychology (8th ed.). Cengage. Chapter 11, Table 11.5, pg. 342.

Cohen_d_s	<i>Effect size</i>
-----------	--------------------

Description

Calculate Cohen's d and Hedges' g introduced in Lakens (2013).

Usage

```
Cohen_d_s(
  y1,
  y2,
  alternative = "two.sided",
  alpha = 0.05,
  mu = 0,
  return_CI = FALSE
)

Hedges_g_s(
  y1,
  y2,
  alternative = "two.sided",
  alpha = 0.05,
  mu = 0,
  return_CI = FALSE
)
```

Arguments

y1	A numeric vector.
y2	A numeric vector.
alternative	Character (default: "two.sided"). Specifies the alternative hypothesis. Available options are c("two.sided", "less", "greater").
alpha	Numeric (default: 0.05). Significance level (0 - 1) for hypothesis testing.
mu	Numeric (default: 0).
return_CI	Logical (default: FALSE). Whether to return the confidence interval of the effect size.

Details

Refer to Lakens (2013), the Cohen's d is the formula 1, the Hedges' g is the formula 4. Their confidence interval (CI) were calculated based on formula 2, by transforming the d and g back to t to fit the general confidence interval calculation. For detailed and precise CI estimates, please use the `effectsize` package.

Value

A numeric scalar or a numeric vector of length 3, depends on `return_CI`.

References

Lakens, D. (2013). Calculating and reporting effect sizes to facilitate cumulative science: A practical primer for t-tests and ANOVAs. *Frontiers in Psychology*, 4. <https://doi.org/10.3389/fpsyg.2013.00863>

compact_letter_display

Compact Letter Display (CLD)

Description

Represent significance statements resulting from all-pairwise comparisons.

Usage

```
compact_letter_display(
  x1,
  x2,
  pvalues,
  grp_names,
  centers,
  alpha = 0.05,
  descending = TRUE,
  display_letters = base::letters,
  display_null_letter = "",
  misc = FALSE
)
```

Arguments

<code>x1</code>	Character vector. The names of the minuend.
<code>x2</code>	Character vector. The names of the subtrahend.
<code>pvalues</code>	Numeric vector. The p-values for the differences of each $x1 - x2$.
<code>grp_names</code>	Character vector. The group names (each factor levels).
<code>centers</code>	Numeric vector. Generally, the corresponding mean or median values of <code>grp_names</code> .

<code>alpha</code>	Numeric (default: 0.05). Significance level, range from 0 to 1.
<code>descending</code>	Logical (default: TRUE). If TRUE, sort the centers in decreasing order.
<code>display_letters</code>	Character vector (default: <code>base::letters</code>). Display symbols.
<code>display_null_letter</code>	Character (default: ""). Symbol for filling the letter's gap.
<code>misc</code>	Logical (default: FALSE). Return other unimportant variables, not for users.

Value

A character vector.

References

Piepho, H.-P. (2004). An algorithm for a letter-based representation of all-pairwise comparisons. *Journal of Computational and Graphical Statistics*, 13(2), 456–466. <https://doi.org/10.1198/1061860043515>

Piepho, H.-P. (2018). Letters in mean comparisons: What they do and don't mean. *Agronomy Journal*, 110(2), 431–434. <https://doi.org/10.2134/agronj2017.10.0580>

Examples

```
out <- Dunn_test(X_X_0, val ~ grp, verbose = FALSE)
tab <- row_arrange(out[["summary"]], "MED")
post <- out[["post_hoc"]]
print(post[, 1:7])
cld <- compact_letter_display(x1 = post$x1,
                             x2 = post$x2,
                             pvalues = post$`Padj (holm)`,
                             grp_names = tab$GROUP,
                             centers = tab$MED)

print(cld)
```

describe

Descriptive statistics

Description

Compute and summarize descriptive statistics for one or more groups.

Usage

```
describe(data, formula, rounding = 2)
```

Arguments

<code>data</code>	A data frame or a list.
<code>formula</code>	A formula with a dependent variable (DV) and an independent variable (IV). For example: DV ~ IV.
<code>rounding</code>	Integer (default: 2). Rounding digits.

Value

A data frame with 13 columns:

GROUP Group name.

CLD Compact letter display for multiple comparisons. This column is returned as an empty character vector and is intended to be filled by post hoc comparison functions. See `oneway::compact_letter_display`.

N Sample size.

AVG Arithmetic mean.

SD Sample standard deviation.

MED Median.

MIN Minimum observed value.

MAX Maximum observed value.

CI (95%) Two-sided 95% confidence interval for the population mean.

SKEW (= 0) Normal distribution has a skewness of 0. See `normality::skewness`.

KURT (= 3) Normal distribution has a kurtosis of 3. See `normality::kurtosis`.

normality Is the data normally distributed? See `normality::is_normal`.

n_outliers Number of possible outliers. See `outlying::Grubbs_test`.

Examples

```
y1 <- c(stats::rnorm(20), 7)
y2 <- c(stats::rnorm(22), -7, 9)
describe(list("apple" = y1, "banana" = y2))
```

<code>df_wide_to_long</code>	<i>Convert a data frame from wide to long format</i>
------------------------------	--

Description

Reshapes a data frame from wide format to long format by stacking one or more columns into a pair of key-value columns. This increases the number of rows while reducing the number of columns. For more advanced reshaping operations, consider using `tidyr::pivot_longer()`.

Usage

```
df_wide_to_long(df, columns, names_to = "grp", values_to = "val", keep = FALSE)
```

Arguments

<code>df</code>	A data frame.
<code>columns</code>	A numeric or character vector specifying the columns to pivot into long format.
<code>names_to</code>	A character string specifying the name of the new column containing the original column names. Default is "grp".
<code>values_to</code>	A character string specifying the name of the new column containing the values from the pivoted columns. Default is "val".
<code>keep</code>	Logical. If TRUE, columns not specified in <code>columns</code> are retained in the output. If FALSE (default), only the pivoted columns are returned.

Value

A data frame in long format. The output contains one column storing the original column names (`names_to`) and another storing the corresponding values (`values_to`). If `keep = TRUE`, non-pivoted columns are retained.

Examples

```
n <- 10
df0 <- data.frame(
  G1 = stats::rnorm(n, 6, 1),
  G2 = stats::rnorm(n, 6, 1),
  G3 = stats::rnorm(n, 3, 1)
)

df_wide_to_long(df0, c("G1", "G2"))
```

Dunn_test

*Dunn's multiple comparison test***Description**

Performs Dunn's multiple comparison procedure for all pairwise comparisons of groups following a significant Kruskal-Wallis test.

Usage

```
Dunn_test(
  data,
  formula = NULL,
  alpha = 0.05,
  p_adjust_method = "holm",
  rounding = 4,
  verbose = TRUE
)
```

Arguments

<code>data</code>	A data frame, or an object returned by <code>Kruskal_Wallis_test()</code> , from which the response and grouping variables are obtained.
<code>formula</code>	A two-sided formula specifying the response and grouping variables in the form <code>response ~ group</code> . Ignored if <code>data</code> is a "oneway_aov" object.
<code>alpha</code>	A numeric value between 0 and 1 specifying the significance level. The default is 0.05.
<code>p_adjust_method</code>	A character string specifying the method used to adjust p-values for multiple comparisons. Must be one of <code>stats::p.adjust.methods</code> . The default is "holm".
<code>rounding</code>	An integer specifying the number of decimal places to display in the output. The default is 4.
<code>verbose</code>	Logical (default: TRUE). Show warnings and messages.

Details

Dunn's test is a nonparametric post hoc procedure that compares group mean ranks after a significant Kruskal-Wallis test. The test statistic follows a normal approximation with an adjustment for tied ranks when ties are present.

Because multiple pairwise comparisons are performed, p-values should be adjusted to control the family-wise error rate or false discovery rate. The adjustment method is specified by `p_adjust_method` and is passed to `stats::p.adjust()`.

If the data satisfy the assumptions of one-way ANOVA, parametric procedures such as `Tukey_HSD_test()`, `REGWQ_test()`, or `Games_Howell_test()` may be more appropriate.

Value

A list containing the following components:

method The name of the statistical procedure.

data The input data used in the analysis.

pre_hoc The Kruskal-Wallis test results.

post_hoc Results of all pairwise Dunn comparisons, including differences in mean ranks, confidence intervals, Z statistics, unadjusted and adjusted p-values, and effect sizes.

summary Descriptive statistics for each group, including compact letter displays (CLD).

References

- Dinno, A. (2015). Nonparametric Pairwise Multiple Comparisons in Independent Groups using Dunn's Test. *The Stata Journal: Promoting Communications on Statistics and Stata*, 15(1), 292-300. <https://doi.org/10.1177/1536867X1501500117>
- Zar, J. H. (2014). *Biostatistical analysis* (5th edition). Pearson. Chapter 11: Multiple comparisons, pg. 255-256.

Examples

```
out <- Dunn_test(X_X_0, val ~ grp)
```

Games_Howell_test	<i>Games-Howell multiple comparison test</i>
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Description

Performs the Games-Howell multiple comparison procedure for all pairwise comparisons of group means. The Games-Howell test does not assume equal variances or equal sample sizes and is commonly used following a one-way analysis of variance when the homogeneity of variance assumption is violated.

Usage

```
Games_Howell_test(
  data,
  formula = NULL,
  alpha = 0.05,
  rounding = 4,
  verbose = TRUE
)
```

Arguments

data	A data frame, or an object returned by <code>oneway_anova()</code> , from which the response and grouping variables are obtained.
formula	A two-sided formula specifying the response and grouping variables in the form <code>response ~ group</code> . Ignored if data is a "oneway_aov" object.
alpha	A numeric value between 0 and 1 specifying the significance level. The default is 0.05.
rounding	An integer specifying the number of decimal places to display in the output. The default is 4.
verbose	Logical (default: TRUE). Show warnings and messages.

Details

The Games-Howell test is based on the studentized range distribution and uses Welch-Satterthwaite degrees of freedom for each pairwise comparison. Unlike Tukey's HSD test, it does not require equal variances or balanced sample sizes.

Value

A list containing the following components:

method The name of the statistical procedure.

data The input data used in the analysis.

pre_hoc The one-way analysis of variance results.

post_hoc Results of all pairwise Games-Howell comparisons, including mean differences, confidence intervals, test statistics, adjusted p-values, and effect sizes.

summary Descriptive statistics for each group, including compact letter displays (CLD).

References

Howell, D. C. (2013). Statistical Methods for Psychology (8th ed.). Cengage Learning. Chapter 12: Multiple comparisons among treatment means, Section 12.7: Tukey's test, pg. 395.

Zar, J. H. (2014). Biostatistical analysis (5th ed.). Pearson. Chapter 11: Multiple comparisons, pg. 246.

Examples

```
out <- Games_Howell_test(O_X_X, val ~ grp)
```

is_balance

Check Balance of Sample Sizes Among Groups

Description

Determines whether sample sizes are approximately balanced across groups. Sample sizes are considered balanced when the smallest and largest group sizes are within the specified tolerance range relative to the average sample size.

Usage

```
is_balance(data, formula, buffer_ratio = 0)
```

Arguments

data	A data frame or a list containing observations from each group.
formula	A formula specifying the dependent variable (DV) and the independent variable (IV) in the form DV ~ IV. This argument is required when data is a data frame and is ignored when data is already a list.
buffer_ratio	Numeric value between 0 and 1 (default: 0). The allowable proportional deviation from the mean sample size. For example, buffer_ratio = 0.2 allows group sizes to differ from the mean by up to 20%. When buffer_ratio = 0, the function requires exact equality of sample sizes among groups.

Details

The function compares each group's sample size with the mean sample size across groups. The sample sizes are considered balanced if:

$$(1 - r) \leq \frac{\min(n)}{\bar{n}} \leq \frac{\max(n)}{\bar{n}} \leq (1 + r)$$

where $\bar{n}$ is the mean sample size across groups and r is `buffer_ratio`.

Value

A logical value:

TRUE Sample sizes among groups are considered balanced.

FALSE At least one group has a sample size outside the specified tolerance range.

Examples

```
is_balance(list(rnorm(10), rnorm(13)), buffer_ratio = 0.2)
is_balance(list(rnorm(10), rnorm(13)), buffer_ratio = 0.1)
```

Kruskal_Wallis_test	<i>Kruskal–Wallis one-way analysis of variance by ranks</i>
---------------------	---

Description

Performs the Kruskal–Wallis rank-sum test for comparing the distributions of two or more independent groups. The test is a nonparametric alternative to one-way ANOVA and is based on the ranks of the observations rather than their original values.

Usage

```
Kruskal_Wallis_test(data, formula, alpha = 0.05, rounding = 4, verbose = TRUE)
```

Arguments

<code>data</code>	A data frame containing the response and grouping variables.
<code>formula</code>	A two-sided formula specifying the response and grouping variables.
<code>alpha</code>	Numeric (default: 0.05). Significance level (range from 0 to 1).
<code>rounding</code>	Integer (default: 4). Number of decimal places displayed in the output.
<code>verbose</code>	Logical (default: TRUE). Show warnings and messages.

Details

The Kruskal–Wallis test ranks all observations across groups and tests whether the mean ranks differ among groups. Under the null hypothesis, the test statistic approximately follows a chi-squared distribution with $k - 1$ degrees of freedom, where k is the number of groups. A correction for tied ranks is applied when ties are present.

Unlike classical one-way ANOVA, the Kruskal–Wallis test is not derived from a decomposition of variance into between-group and within-group sums of squares. Consequently, residual sums of squares and residual mean squares are not defined in the classical procedure.

Value

A data frame summarizing the Kruskal–Wallis test result in an ANOVA-like table:

DF Degrees of freedom.

SS Rank-based between-group sum of squares. Residual and total sums of squares are not defined for the classical Kruskal–Wallis test and are therefore returned as NA.

MS Rank-based mean square (SS / DF). Only the between-group value is defined.

H Observed Kruskal–Wallis H statistic.

Hcrit Critical chi-squared value at the specified significance level.

Pvalue P-value associated with the H statistic.

signif Significance code corresponding to the p-value.

p_omega2 Effect size (currently returned as NA).

method Statistical method ("Kruskal-Wallis").

The rows correspond to the between-group effect ("Group"), residuals ("Residuals"), and total ("Total"). Only the between-group row contributes to the Kruskal–Wallis test statistic.

References

Kruskal, W. H., & Wallis, W. A. (1952). Use of ranks in one-criterion variance analysis. *Journal of the American Statistical Association*, 47(260), 583–621.

Howell, D. C. (2013). *Statistical methods for psychology* (8th edition). Cengage. Chapter 18, Section 18.9, pg. 678-679.

Hollander, M., Wolfe, D. A., & Chicken, E. (2014). *Nonparametric Statistical Methods* (3rd ed.). Wiley. Chapter 6, pg. 204-206.

Examples

```
lst <- list(
  "depressant" = c(55, 0, 1, 0, 50, 60, 44),
  "stimulant" = c(73, 85, 51, 63, 85, 85, 66, 69),
  "placebo" = c(61, 54, 80, 47)
)
normality::is_normal(lst)
Kruskal_Wallis_test(lst)

Kruskal_Wallis_test(anorexia, weight_gain ~ therapy)
```

morphine*Morphine tolerance data*

Description

A dataset containing morphine tolerance measurements from five experimental treatment groups. The data consist of tolerance values recorded for eight experimental units in each group, resulting in a balanced one-way design with 40 observations.

Usage

`morphine`

Format

A data frame with 40 rows and 2 columns:

tolerance A numeric variable containing the morphine tolerance measurement.

grp A factor identifying the experimental treatment group with five levels: "MS", "MM", "SS", "SM", and "McM".

Details

The treatment groups are coded as:

MS Morphine followed by saline.

MM Morphine followed by morphine.

SS Saline followed by saline.

SM Saline followed by morphine.

McM Morphine followed by challenge morphine.

The dataset represents a balanced one-way experimental design with five independent groups and eight observations per group. It is suitable for demonstrating one-way ANOVA, nonparametric alternatives, homogeneity of variance tests, post hoc comparisons, and effect size estimation.

References

Howell, D. C. (2013). Statistical methods for psychology (8th ed.). Cengage. Chapter 11, Table 12.1, pg. 375.

oneway_anova	<i>One-Way Analysis of Variance</i>
--------------	-------------------------------------

Description

Performs a one-way analysis of variance (ANOVA) to compare the means of two or more independent groups. By default, the function automatically selects between the classical Fisher ANOVA and Welch's ANOVA according to whether the group variances are judged to be homogeneous.

Usage

```
oneway_anova(
  data,
  formula,
  alpha = 0.05,
  var_equal = NA,
  rounding = 4,
  verbose = TRUE
)
```

Arguments

data	A data frame containing the response and grouping variables.
formula	A two-sided formula specifying the response and grouping variables.
alpha	Numeric (default: 0.05). Significance level (range from 0 to 1).
var_equal	A logical value indicating whether equal variances should be assumed. If TRUE, Fisher's ANOVA is performed. If FALSE, Welch's ANOVA is performed. If NA (default), equality of variances is determined automatically using <code>varequal::is_var_equal()</code> .
rounding	Integer (default: 4). Number of decimal places displayed in the output.
verbose	Logical (default: TRUE). Show warnings and messages.

Details

If `var_equal = TRUE`, Fisher's ANOVA is performed. If `var_equal = FALSE`, Welch's ANOVA is performed. When `var_equal = NA` (the default), homogeneity of variances is assessed using `varequal::is_var_equal()`, and the appropriate test is selected automatically.

Fisher's ANOVA assumes independent observations, normally distributed populations, and equal population variances. When the equal-variance assumption is violated, Welch's ANOVA provides a more robust alternative by adjusting the test statistic and denominator degrees of freedom.

Value

A data frame representing the ANOVA table with the following columns:

DF Degrees of freedom.

SS Sum of squares.

MS Mean square.

Fvalue Observed F statistic.

Fcrit Critical F value at the specified significance level.

Pvalue P-value associated with the F statistic.

signif Significance code corresponding to the p-value.

p_omega2 Effect size. Partial omega squared.

method Show whether Fisher's or Welch's ANOVA was conducted.

The rows correspond to the treatment groups ("Group"), residual error ("Residuals"), and total variation ("Total").

References

Howell, D. C. (2013). Statistical methods for psychology (8th ed.). Cengage Learning. Chapter 11, pg. 325-345.

Montgomery, D. C. (2017). Design and analysis of experiments (Ninth edition). John Wiley & Sons, Inc.

Examples

```
# Automatically select the appropriate procedure
oneway_anova(O_X_X, val ~ grp)

# Classical one-way ANOVA
oneway_anova(O_O_X, val ~ grp, var_equal = TRUE)

# Welch's ANOVA
oneway_anova(O_X_X, val ~ grp, var_equal = FALSE)
```

oneway_art

One-Way Aligned Ranked Transformed Analysis of Variance (ART-ANOVA)

Description

Performs a one-way analysis of variance (ANOVA) to compare the means of two or more independent groups. By default, the function automatically selects between the classical Fisher ANOVA and Welch's ANOVA according to whether the group variances are judged to be homogeneous.

Usage

```
oneway_art(
  data,
  formula,
  alpha = 0.05,
  var_equal = NA,
  rounding = 4,
  verbose = TRUE
)
```

Arguments

<code>data</code>	A data frame containing the response and grouping variables.
<code>formula</code>	A two-sided formula specifying the response and grouping variables.
<code>alpha</code>	Numeric (default: 0.05). Significance level (range from 0 to 1).
<code>var_equal</code>	A logical value passes into the <code>oneway_anova()</code> to perform regular ANOVA on the ranked-response variable.
<code>rounding</code>	Integer (default: 4). Number of decimal places displayed in the output.
<code>verbose</code>	Logical (default: TRUE). Show warnings and messages.

Details

If `var_equal = TRUE`, Fisher's ANOVA is performed on the transformed response variable. If `var_equal = FALSE`, Welch's ANOVA is performed. When `var_equal = NA` (the default), homogeneity of variances is assessed using `varequal::is_var_equal()`, and the appropriate test is selected automatically.

Value

A data frame representing the ANOVA table with the following columns:

DF Degrees of freedom.

SS Sum of squares.

MS Mean square.

Fvalue Observed F statistic.

Fcrit Critical F value at the specified significance level.

Pvalue P-value associated with the F statistic.

signif Significance code corresponding to the p-value.

method A character specifying this is an ART-ANOVA procedure.

The rows correspond to the treatment groups ("Group"), residual error ("Residuals"), and total variation ("Total").

References

- Wobbrock, J. O., Findlater, L., Gergle, D., & Higgins, J. J. (2011). The aligned rank transform for nonparametric factorial analyses using only ANOVA procedures. *Proceedings of the SIGCHI Conference on Human Factors in Computing Systems*, 2011, 143–146. <https://doi.org/10.1145/1978942.1978963>
- Elkin, L. A., Kay, M., Higgins, J. J., & Wobbrock, J. O. (2021). An aligned rank transform procedure for multifactor contrast tests. *Proceedings of the 34th Annual ACM Symposium on User Interface Software and Technology*, 754–768. <https://doi.org/10.1145/3472749.3474784>

Examples

```
normality::is_normal(anorexia, weight_gain ~ therapy)
oneway_art(anorexia, weight_gain ~ therapy)
```

pairwise_comparison	<i>Multiple comparison procedure</i>
---------------------	--------------------------------------

Description

Performs a one-way analysis of variance followed by an appropriate post-hoc multiple pairwise comparison based on the distributional characteristics and design of the data. The procedure evaluates normality, homogeneity of variances, and sample-size balance to select an appropriate analysis protocol.

Usage

```
pairwise_comparison(
  data,
  formula,
  alpha = 0.05,
  rounding = 4,
  verbose = TRUE,
  p_adjust_method = "holm"
)
```

Arguments

data	A data frame containing the response and grouping variables.
formula	A two-sided formula specifying the response and grouping variables in the form of response ~ group.
alpha	A numeric significance level used for hypothesis testing. The default is 0.05.
rounding	An integer specifying the number of decimal places used when reporting numerical results. The default is 4.
verbose	A logical value indicating whether the results should be printed to the console. If TRUE, the default, the selected analysis protocol, post-hoc comparisons, and summary are printed. If FALSE, no output is printed.

p_adjust_method

A character string specifying the method used to adjust p-values for Dunn's multiple-comparison test when the Kruskal-Wallis/Dunn protocol is selected. The default is "holm". See `stats::p.adjust()` for available methods.

Details

If the response variable is normally distributed, the post-hoc procedure is selected according to variance homogeneity and sample-size balance:

- equal variances and balanced group sizes: REGWQ test;
- equal variances and unbalanced group sizes: Tukey-Kramer test;
- unequal variances: Games-Howell test.

If the response variable is not normally distributed, an aligned-rank-transform (ART) analysis is first attempted. Normality and variance homogeneity are then reassessed using the aligned-ranked response. If the aligned-ranked response is normally distributed, the same variance- and balance-based selection described above is applied. Otherwise, the procedure uses the Kruskal-Wallis test followed by Dunn's multiple-comparison test.

The resulting object contains the selected pre-hoc and post-hoc analysis results and identifies the complete analysis protocol in the method component.

The analysis proceeds according to the following decision sequence:

1. A preliminary one-way ANOVA is performed to assess normality and homoscedasticity.
2. If the response is normally distributed, the post-hoc procedure is selected according to variance homogeneity and sample-size balance.
3. If the response is not normally distributed, an ART analysis is performed.
4. Normality and variance homogeneity are reassessed using the aligned-ranked response from the ART analysis.
5. If the aligned-ranked response remains non-normal, the Kruskal-Wallis test followed by Dunn's test is used.
6. Otherwise, the REGWQ, Tukey-Kramer, or Games-Howell procedure is selected according to variance homogeneity and sample-size balance.

Thus, the possible analysis protocols are:

- one-way ANOVA + REGWQ;
- one-way ANOVA + Tukey-Kramer;
- one-way ANOVA + Games-Howell;
- ART-ANOVA + REGWQ;
- ART-ANOVA + Tukey-Kramer;
- ART-ANOVA + Games-Howell;
- Kruskal-Wallis + Dunn's test.

The function is intended as a general-purpose adaptive procedure for one-factor experimental designs. The selected method depends on the diagnostic results obtained from the supplied data and should therefore be interpreted together with the underlying assumptions and study design.

Value

A list containing the results of the selected analysis protocol. The method component identifies the complete protocol, combining the *a priori* and post-hoc procedures. The object also contains the post-hoc results and summary produced by the selected procedure.

See Also

[oneway_anova\(\)](#), [oneway_art\(\)](#), [REGWQ_test\(\)](#), [Tukey_Kramer_test\(\)](#), [Games_Howell_test\(\)](#), [Kruskal_Wallis_test\(\)](#), [Dunn_test\(\)](#)

Examples

```
# Normally distributed, variance equal, balanced groups
pairwise_comparison(O_O_O, val ~ grp)

# Normally distributed, variance equal, unbalanced groups
pairwise_comparison(O_O_X, val ~ grp)

# Normally distributed, variance unequal
pairwise_comparison(O_X_X, val ~ grp)

# Non-parametric
pairwise_comparison(X_X_X, val ~ grp)
```

plasma_etching	<i>Plasma Etching Experiment</i>
----------------	----------------------------------

Description

Etch Rate Data (in Å/min) from the Plasma Etching Experiment.

Usage

```
plasma_etching
```

Format

A data frame with 20 rows and 2 columns:

power A factor identifying the power (W) (160(20)220).

etch_rate A numeric response variable.

References

Montgomery, D. C. (2017). Design and analysis of experiments (Ninth edition). John Wiley & Sons, Inc.

pval2asterisk

Convert p-values to significance labels

Description

Converts numeric p-values into categorical significance labels according to user-defined thresholds. The function is commonly used to annotate statistical results in tables and figures.

Usage

```
pval2asterisk(
  x,
  break_points = c(0.055, 0.05, 0.01, 0.001, 0),
  symbols = c("ns", ".", "*", "**", "***")
)
```

Arguments

x	A numeric vector of p-values.
break_points	A numeric vector of significance thresholds in descending order. Each threshold defines the upper bound of a significance interval. The default values correspond to: • $p > 0.055$: "ns" • $0.05 < p \leq 0.055$: "." • $0.01 < p \leq 0.05$: "*" • $0.001 < p \leq 0.01$: "**" • $p \leq 0.001$: "***"
symbols	A character vector of significance labels corresponding to break_points. The lengths of break_points and symbols must be identical.

Details

Each p-value is assigned to exactly one interval defined by break_points. Values greater than the first threshold are assigned symbols[1], whereas values less than or equal to the last threshold are assigned the last element of symbols. Intermediate intervals are matched sequentially.

The function assumes that break_points are supplied in descending order.

Value

A character vector of the same length as x, where each element is the corresponding significance label.

Examples

```
p <- c(0.20, 0.04, 0.008, 0.0005, 1e-6)
pval2asterisk(p)

# Custom significance labels
pval2asterisk(
  p,
  break_points = c(0.05, 0.01, 0),
  symbols = c("Not significant", "Significant", "Highly significant")
)
```

REGWQ_test

*Ryan-Einot-Gabriel-Welsch Studentized Range (REGWQ) test***Description**

Performs the Ryan-Einot-Gabriel-Welsch Studentized Range (REGWQ) multiple comparison procedure for all pairwise comparisons of group means following a one-way analysis of variance. REGWQ is a stepwise procedure based on the studentized range distribution that generally provides greater statistical power than Tukey's HSD while maintaining strong control of the family-wise error rate under balanced designs.

Usage

```
REGWQ_test(data, formula = NULL, alpha = 0.05, rounding = 4, verbose = TRUE)
```

Arguments

data	A data frame, or an object returned by <code>oneway_anova()</code> , from which the response and grouping variables are obtained.
formula	A two-sided formula specifying the response and grouping variables in the form <code>response ~ group</code> . Ignored if data is a "oneway_aov" object.
alpha	A numeric value between 0 and 1 specifying the significance level. The default is 0.05.
rounding	An integer specifying the number of decimal places to display in the output. The default is 4.
verbose	Logical (default: TRUE). Show warnings and messages.

Details

The REGWQ procedure is a stepwise multiple comparison method based on the studentized range distribution. Group means are first ordered, after which pairwise comparisons are performed using significance levels that depend on the number of ordered means spanned by each comparison. This adaptive strategy generally yields greater statistical power than Tukey's HSD while maintaining control of the family-wise error rate.

The procedure assumes that observations are independent, residuals are approximately normally distributed, and population variances are equal. It is intended primarily for balanced one-way designs. When sample sizes are unequal but variances remain homogeneous, consider using `Tukey_Kramer_test()`. When variances are unequal, consider using `Games_Howell_test()`.

Value

A list containing the following components:

method The name of the statistical procedure.

data The input data used in the analysis.

pre_hoc The one-way analysis of variance results.

post_hoc Results of all pairwise REGWQ comparisons, including mean differences, confidence intervals, studentized range statistics, p-values, modified significance levels, and effect sizes.

summary Descriptive statistics for each group, including compact letter displays (CLD).

References

Howell, D. C. (2010). Statistical Methods for Psychology (7th ed.). Cengage Learning. Chapter 12: Multiple comparisons among treatment means, Section 12.6: Post hoc comparisons, pg. 393-394.

Examples

```
out <- REGWQ_test(morphine, tolerance ~ grp)
```

row_arrange

Rearrange rows by the order of one or more columns

Description

Reorders the rows of a data frame according to user-specified ordering of the values in one or more columns. If `by_order` is omitted, the unique values in each selected column are sorted and used as the ordering levels.

Usage

```
row_arrange(data, cols, by_order)
```

Arguments

data A data frame.

cols A character or numeric vector specifying the columns used to determine the row order. Character values are interpreted as column names, and numeric values as column indices.

by_order A list specifying the ordering of values for each column in `cols`. Each element should be a character vector containing the desired ordering of the corresponding column. If a single vector is supplied, it is treated as a list of length one. If omitted, the unique values in each column are sorted and used as the ordering.

Value

A data frame with the same columns as data, but with rows reordered according to the specified column orderings.

Examples

```
fct_lvl <- c("A", "B", "C", "D")

df0 <- data.frame(
  C1 = c("B", "D", "A", "A", "B", "C", "D"),
  C2 = c("D", "C", "D", "A", "C", "B", "A"),
  C3 = 1:7
)

row_arrange(df0, 1:2, list(fct_lvl, fct_lvl))

# Use the default ordering (sorted unique values)
row_arrange(df0, c("C2", "C3"))
```

simulated_data

Simulated datasets for statistical analysis

Description

A collection of six simulated datasets representing different combinations of distributional assumptions, variance homogeneity, and sample-size balance. These datasets are designed for demonstrating and evaluating statistical procedures under a range of common experimental conditions.

Usage

O_0_0

O_0_X

O_X_X

X_0_0

X_X_0

X_X_X

Format

Six data frames, each containing the following two variables:

grp Character vector identifying the experimental group.

val Numeric vector containing the simulated response values.

The number of observations differs according to the experimental design: balanced datasets contain 120 observations (20 per group), whereas unbalanced datasets contain 128 observations with group sizes of 27, 24, 16, 30, 11, and 20.

An object of class `data.frame` with 100 rows and 2 columns.

An object of class `data.frame` with 128 rows and 2 columns.

An object of class `data.frame` with 128 rows and 2 columns.

An object of class `data.frame` with 120 rows and 2 columns.

An object of class `data.frame` with 120 rows and 2 columns.

An object of class `data.frame` with 110 rows and 2 columns.

Details

All datasets contain six groups (G1 to G6) and two variables:

grp A character variable identifying the group.

val A numeric response variable.

The dataset names follow a three-character notation separated by underscores: D_V_B, where each position describes a different property of the data:

First position (D) Distribution. 0 indicates normally distributed data, whereas X indicates distribution-free or non-normally distributed data.

Second position (V) Variance. 0 indicates homoscedasticity, whereas X indicates heteroscedasticity.

Third position (B) Design balance. 0 indicates a balanced design, whereas X indicates an unbalanced design.

The six datasets are:

O_0_0 Normally distributed, homoscedastic, and balanced-designed data. Each group contains 20 observations.

O_0_X Normally distributed, homoscedastic, and unbalanced-designed data. The group sample sizes are 27, 24, 16, 30, 11, and 20, respectively.

O_X_X Normally distributed, heteroscedastic, and unbalanced-designed data. The group sample sizes are 27, 24, 16, 30, 11, and 20, respectively.

X_0_0 Distribution-free or non-normally distributed, homoscedastic, and balanced-designed data. Each group contains 20 observations.

X_X_0 Distribution-free or non-normally distributed, heteroscedastic, and balanced-designed data. Each group contains 20 observations.

X_X_X Distribution-free or non-normally distributed, heteroscedastic, and unbalanced-designed data. The group sample sizes are 27, 24, 16, 30, 11, and 20, respectively.

The normally distributed datasets were generated using `stats::rnorm()`. The distribution-free datasets were generated using combinations of `stats::rcauchy()`, `stats::runif()`, and `stats::rgamma()`, together with `stats::rnorm()`. Different location and scale parameters were used to produce the intended distributional and variance characteristics.

These datasets are intended for methodological examples, unit tests, demonstrations, and comparisons of statistical procedures under different combinations of assumptions. They should not be interpreted as empirical observations from a real population.

Dataset characteristics

Dataset	Distribution	Variance	Design
O_0_0	Normal	Homoscedastic	Balanced
O_0_X	Normal	Homoscedastic	Unbalanced
O_X_X	Normal	Heteroscedastic	Unbalanced
X_0_0	Non-normal	Homoscedastic	Balanced
X_X_0	Non-normal	Heteroscedastic	Balanced
X_X_X	Non-normal	Heteroscedastic	Unbalanced

Examples

```
data(O_0_0)

boxplot(val ~ grp, data = O_0_0)

data(X_X_X)

boxplot(val ~ grp, data = X_X_X)

aggregate(val ~ grp, data = O_0_0, FUN = mean)

aggregate(val ~ grp, data = X_X_X, FUN = mean)
```

Tukey_HSD_test	<i>Tukey's Honestly Significant Difference (Tukey-HSD) test</i>
----------------	---

Description

Performs Tukey's Honestly Significant Difference (HSD) multiple comparison procedure for all pairwise comparisons of group means following a one-way analysis of variance. The Tukey-HSD test controls the family-wise error rate and is appropriate when the assumptions of normality, homogeneity of variances, and balanced group sizes are reasonably satisfied.

Usage

```
Tukey_HSD_test(
  data,
  formula = NULL,
  alpha = 0.05,
  rounding = 4,
  verbose = TRUE
)
```

Arguments

<code>data</code>	A data frame, or an object returned by <code>oneway_anova()</code> , from which the response and grouping variables are obtained.
<code>formula</code>	A two-sided formula specifying the response and grouping variables in the form <code>response ~ group</code> . Ignored if <code>data</code> is a "oneway_aov" object.
<code>alpha</code>	A numeric value between 0 and 1 specifying the significance level. The default is 0.05.
<code>rounding</code>	An integer specifying the number of decimal places to display in the output. The default is 4.
<code>verbose</code>	Logical (default: TRUE). Show warnings and messages.

Details

Tukey's HSD test is based on the studentized range distribution and assumes equal variances across groups. It provides simultaneous confidence intervals and adjusted p -values that control the family-wise error rate for all pairwise comparisons.

This implementation is intended primarily for balanced one-way designs. When sample sizes are unequal but variances remain homogeneous, consider using `Tukey_Kramer_test()`. When variances are unequal, consider using `Games_Howell_test()`.

Value

A list containing the following components:

method The name of the statistical procedure.

data The input data used in the analysis.

pre_hoc The one-way analysis of variance results.

post_hoc Results of all pairwise Tukey-HSD comparisons, including mean differences, confidence intervals, studentized range statistics, adjusted p -values, and effect sizes.

summary Descriptive statistics for each group, including compact letter displays (CLD).

References

Howell, D. C. (2013). Statistical Methods for Psychology (8th ed.). Cengage Learning. Chapter 12: Multiple comparisons among treatment means, Section 12.7: Tukey's test, pg. 394.

Zar, J. H. (2014). Biostatistical analysis (5th edition). Pearson. Chapter 11: Multiple comparisons, pg. 241-243.

Examples

```
out <- Tukey_HSD_test(0_0_0, val ~ grp)
```

Tukey_Kramer_test	<i>Tukey-Kramer test</i>
-------------------	--------------------------

Description

Represent significance statements resulting from all-pairwise comparisons.

Usage

```
Tukey_Kramer_test(
  data,
  formula = NULL,
  alpha = 0.05,
  rounding = 4,
  verbose = TRUE
)
```

Arguments

data	A data frame in which the variables specified in the formula will be found.
formula	A formula specifying the model.
alpha	Numeric value range from 0 to 1 (default: 0.05). The error tolerance.
rounding	Integer (default: 4). Rounding digits.
verbose	Logical (default: TRUE). Show warnings and messages.

Value

A list with 4 elements:

method Statistical procedures that were conducted.

data The input data and possibly other transformed data.

pre_hoc *a priori* test result.

post_hoc Post-hoc test result.

summary Descriptive statistics.

References

Howell, D. C. (2013). Statistical Methods for Psychology (8th ed.). Cengage Learning. Chapter 12: Multiple comparisons among treatment means, Section 12.7: Tukey's test, pg. 394.

Zar, J. H. (2014). Biostatistical analysis (5th edition). Pearson. Chapter 11: Multiple comparisons, pg. 244-245.

Examples

```
out <- Tukey_Kramer_test(O_O_X, val ~ grp)
```

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