# Package: pubh (via r-universe)

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Title A Toolbox for Public Health and Epidemiology

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Description A toolbox for making R functions and capabilities more accessible to students and professionals from Epidemiology and Public Health related disciplines. Includes a function to report coefficients and confidence intervals from models using robust standard errors (when available), functions that expand 'ggplot2' plots and functions relevant for introductory papers in Epidemiology or Public Health. Please note that use of the provided data sets is for educational purposes only.

**Depends** R (>= 4.1.0), emmeans, ggformula, gtsummary (>= 1.9.9), huxtable, stats

**Imports** car, dplyr, Epi, epiR, epitools, ggplot2, jtools, lmtest, moonBook, performance, sandwich, sjlabelled, sjmisc, survival, tibble, tidyselect

**Suggests** broom, broom.helpers (>= 1.15.0), cardx, easystats, effectsize, ggeffects, ggsci, ISwR, knitr, lme4, MASS, nlme, nnet, report, rmarkdown, rstatix, sjPlot, see

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**Encoding UTF-8** 

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axis\_labs

Apply labels from variables to axis-labels in plots.

# Description

axis\_labs takes labels from labelled data to use them as axis-labels for plots generated by gformula or ggplot2.

### Usage

```
axis_labs(object)
```

# Arguments

object

ggplot2 object (see examples).

### **Details**

This functions is helpful when data has been already labelled by sjlabelled. It retrives variable labels and use them for plotting.

### Value

A ggplot2 object.

bar\_error

#### **Examples**

```
data(kfm, package = "ISwR")
require(sjlabelled, quietly = TRUE)
kfm <- kfm |>
  var labels(
   dl.milk = "Breast-milk intake (dl/day)",
    sex = "Sex",
   weight = "Child weight (kg)",
   ml.suppl = "Milk substitute (ml/day)",
   mat.weight = "Maternal weight (kg)",
   mat.height = "Maternal height (cm)"
  )
kfm |>
  gf_point(weight ~ dl.milk) |>
  gf_lm(col = 2, interval = "confidence", col = 2) |>
  axis_labs()
kfm |>
  box_plot(dl.milk ~ sex, fill = "thistle", alpha = 0.8) |>
  axis_labs() |>
  gf_star(x1 = 1, y1 = 10.9, x2 = 2, y2 = 11, y3 = 11.2)
```

bar\_error

Bar charts with error bars.

#### **Description**

bar\_error constructs bar charts in with error bars showing 95 confidence intervals around mean values. High of bars represent mean values.

### Usage

```
bar_error(
  object = NULL,
  formula = NULL,
  data = NULL,
  fill = "indianred3",
  col = "black",
  alpha = 0.7,
  ...
)
```

#### **Arguments**

object When chaining, this holds an object produced in the earlier portions of the chain.

Most users can safely ignore this argument. See details and examples.

formula A formula with shape:  $y \sim x$  or  $y \sim x \mid z$  where y is a numerical variable and both

x and z are factors.

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data	A data frame where the variables in the formula can be found.
fill	Colour used to fill the bars.
col	Colour used for the borders of the bars.
alpha	Opacity of the colour fill $(0 = invisible, 1 = opaque)$ .
	Additional information passed to gf_summary.

### **Examples**

```
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)
data(birthwt, package = "MASS")
birthwt <- birthwt |>
  mutate(
   smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
   Race = factor(race > 1, labels = c("White", "Non-white"))
  ) |>
  var_labels(
   bwt = "Birth weight (g)",
   smoke = "Smoking status"
  )
birthwt |>
  bar_error(bwt ~ smoke, fill = "plum3")
  bar_error(bwt ~ smoke | Race, fill = "plum3")
birthwt |>
  bar_error(bwt ~ smoke, fill = ~Race)
```

Bernard

Survival of patients with sepsis.

### **Description**

A randomised, double-blind, placebo-controlled trial of intravenous ibuprofen in 455 patients who had sepsis, defined as fever, tachycardia, tachypnea, and acute failure of at least one organ system.

#### Usage

Bernard

### **Format**

A labelled tibble with 455 rows and 9 variables:

id Patient ID

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```
treat Treatment, factor with levels "Placebo" and "Ibuprofen".
race Race/ethnicity, factor with levels "White", "African American" and "Other".
fate Mortality status at 30 days, factor with levels "Alive" and "Dead".
apache Baseline APACHE score.
o2del Oxygen delivery at baseline.
followup Follow-up time in hours.
temp0 Baseline temperature in centigrades.
temp10 Temperature after 36 hr in centigrades.
```

#### **Source**

Bernard, GR, et al. (1997) The effects of ibuprofen on the physiology and survival of patients with sepsis, N Engl J Med 336: 912–918.

### **Examples**

```
require(dplyr, quietly = TRUE)
data(Bernard)

Bernard |>
    select(fate, treat) |>
    cross_tbl(by = "fate")

contingency(fate ~ treat, data = Bernard)
```

bland\_altman

Bland-Altman agreement plots.

### **Description**

Bland-Altman agreement plots.

```
bland_altman(
  object = NULL,
  formula = NULL,
  data = NULL,
  pch = 20,
  size = 1,
  col = "black",
  transform = FALSE,
  ...
)
```

box\_plot 7

### **Arguments**

object	When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples.
formula	A formula with shape: $y \sim x$ (see details).
data	A data frame where the variables in the formula can be found.
pch	Symbol for plotting data.
size	Size of the symbol using to plot data.
col	Colour used for the symbol to plot data.
transform	Logical, should ratios instead of difference be used to construct the plot?
	Further arguments passed to gf_point.

### **Details**

bland\_altman constructs Bland-Altman agreement plots.

Variables in formula are continuous paired observations. When the distribution of the outcome is not normal, but becomes normal with a log-transformation, bland\_altman can plot the ratio between outcomes (difference in the log scale) by using option transform = TRUE.

# **Examples**

```
data(wright, package = "ISwR")
wright |>
  bland_altman(mini.wright ~ std.wright,
    pch = 16,
    ylab = "Large-mini expiratory flow rate (l/min)",
    xlab = "Mean expiratory flow rate (l/min)"
) |>
  gf_labs(
    y = "Large-mini expiratory flow rate (l/min)",
    x = "Mean expiratory flow rate (l/min)"
)

data(Sharples)
Sharples |>
  bland_altman(srweight ~ weight, transform = TRUE) |>
  gf_labs(x = "Mean of weights (kg)", y = "Measured weight / Self-reported weight")
```

box\_plot

Construct box plots.

### **Description**

box\_plot is a wrap function that calls gf\_boxplot to construct more aesthetic box plots.

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#### Usage

```
box_plot(
  object = NULL,
  formula = NULL,
  data = NULL,
  fill = "indianred3",
  alpha = 0.7,
  outlier.shape = 20,
  outlier.size = 1,
  ...
)
```

#### **Arguments**

object When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples. formula A formula with shape:  $y \sim x$  where y is a numerical variable and x is a factor. data A data frame where the variables in the formula can be found. fill Colour used for the box passed to gf\_boxplot. alpha Opacity (0 = invisible, 1 = opaque). outlier.shape Shape (pch) used as symbol for the outliers. outlier.size Size of the outlier symbol. Further arguments passed to gf\_boxplot.

```
data(kfm, package = "ISwR")
require(sjlabelled, quietly = TRUE)
kfm <- kfm |>
 var_labels(
   dl.milk = "Breast-milk intake (dl/day)",
   sex = "Sex",
   weight = "Child weight (kg)",
   ml.suppl = "Milk substitute (ml/day)",
   mat.weight = "Maternal weight (kg)",
   mat.height = "Maternal height (cm)"
 )
kfm |>
 box_plot(dl.milk ~ sex, fill = "thistle", alpha = 0.8)
t.test(dl.milk ~ sex, data = kfm)
kfm |>
 box_plot(dl.milk ~ sex, fill = "thistle", alpha = 0.8) |>
 gf_star(1, 10.9, 2, 11, 11.4, legend = "p = 0.035", size = 2.5)
```

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Brenner

Prevalence of Helicobacter pylori infection in preschool children.

### **Description**

A data set containing the prevalence of Helicobacter pylori infection in preschool children according to parental history of duodenal or gastric ulcer.

# Usage

Brenner

#### **Format**

A labelled tibble with 863 rows and 2 variables:

```
ulcer History of duodenal or gastric ulcer, factor with levels "No" and "Yes". infected Infected with Helicobacter pylori, factor with levels "No" and "Yes".
```

#### **Source**

Brenner H, Rothenbacher D, Bode G, Adler G (1998) Parental history of gastric or duodenal ulcer and prevalence of Helicobacter pylori infection in preschool children: population based study. BMJ 316:665.

### **Examples**

```
require(dplyr, quietly = TRUE)
data(Brenner)

Brenner |>
    select(infected, ulcer) |>
    cross_tbl(by = "infected")

contingency(infected ~ ulcer, data = Brenner, method = "cross.sectional")
```

bst

Bootstrap Confidence Intervals.

#### **Description**

bst estimates confidence intervals around the mean, median or geo\_mean.

```
bst(x, stat = "mean", n = 1000, CI = 95, digits = 2)
```

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# Arguments

X	A numerical variable. Missing observations are removed by default.
stat	Statistic, either "mean" (default), "median" or "gmean" (geometric mean).
n	Number of replicates for the bootstrap (n=1000 by default).
CI	Confidence intervals (CI=95 by default).
digits	Number of digits for rounding (default = 2).

### Value

A data frame with the estimate and confidence intervals.

# **Examples**

```
data(IgM, package = "ISwR")
bst(IgM, "median")
bst(IgM, "gmean")
```

chisq.fisher Internal test for chi-squared assumption. Fisher  $(2 \ by \ 2)$ . If results = T, it fails

# Description

chisq.fisher is an internal function called by contingency and contingency2 that uses the Fisher exact test if results from the assumptions for the chi-squared test fail.

# Usage

```
chisq.fisher(tab)
```

### Arguments

tab A numeric two by two table.

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coef\_det

Coefficient of determination.

#### **Description**

coef\_det estimates the coefficient of determination (r-squared) from fitted (predicted) and observed values. Outcome from the model is assumed to be numerical.

### Usage

```
coef_det(obs, fit)
```

# Arguments

obs Vector with observed values (numerical outcome).

fit Vector with fitted (predicted) values.

### Value

A scalar, the coefficient of determination (r-squared).

```
## Linear regression:
Riboflavin <- seq(0, 80, 10)
OD <- 0.0125 * Riboflavin + rnorm(9, 0.6, 0.03)
titration <- data.frame(Riboflavin, OD)
model1 <- lm(OD ~ Riboflavin, data = titration)
summary(model1)
coef_det(titration$OD, fitted(model1))

## Non-linear regression:
library(nlme, quietly = TRUE)
data(Puromycin)
mm.tx <- gnls(rate ~ SSmicmen(conc, Vm, K),
    data = Puromycin,
    subset = state == "treated"
)
summary(mm.tx)
coef_det(Puromycin$rate[1:12], mm.tx$fitted)</pre>
```

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contingency

Measures of association from two by two contingency tables (formula).

### **Description**

contingency is a wrap that calls epi. 2by2 from package epiR.

### Usage

```
contingency(
  object = NULL,
  formula = NULL,
  data = NULL,
  method = "cohort.count",
   ...
)
```

#### **Arguments**

object	When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples.
formula	A formula with shape: outcome ~ exposure.
data	A data frame where the variables in the formula can be found.
method	A character string with options: "cohort.count", "cohort.time", "case.control", or "cross.sectional".
	Further arguments passed to epi. 2by2.

#### **Details**

contingency uses a formula as a way to input variables.

contingency displays the contingency table as a way for the user to check that the reference levels in the categorical variables (outcome and exposure) are correct. Then displays measures of association (table from epi.2by2). It also reports either chi-squared test or exact Fisher's test; contingency checks which one of the tests two is appropriate.

#### See Also

```
epi.2by2.
```

```
## A case-control study on the effect of alcohol on oesophageal cancer. Freq <- c(386, 29, 389, 171) status <- gl(2, 1, 4, labels = c("Control", "Case")) alcohol <- gl(2, 2, labels = c("0-39", "40+")) cancer <- data.frame(Freq, status, alcohol)
```

contingency2

```
cancer <- expand_df(cancer)
contingency(status ~ alcohol, data = cancer, method = "case.control")

data(Oncho)
require(dplyr, quietly = TRUE)

Oncho |>
    select(mf, area) |>
    cross_tbl(by = "mf") |>
    theme_pubh(2)

Oncho |>
    contingency(mf ~ area)
```

contingency2

Measures of association from two by two contingency tables (direct input).

### **Description**

contingency2 is a wrap that calls epi. 2by2 from package epiR.

#### Usage

```
contingency2(aa, bb, cc, dd, ...)
```

#### **Arguments**

aa	Number of cases where both exposure and outcome are present.
bb	Number of cases where exposure is present but outcome is absent.
сс	Number of cases where exposure is absent but outcome is present.
dd	Number of cases where both exposure and outcome are absent.
	Further arguments passed to epi. 2by2.

#### See Also

```
epi.2by2.
```

```
## A case-control study on the effect of alcohol on oesophageal cancer.
Freq <- c(386, 29, 389, 171)
status <- gl(2, 1, 4, labels = c("Control", "Case"))
alcohol <- gl(2, 2, labels = c("0-39", "40+"))
cancer <- data.frame(Freq, status, alcohol)
cancer <- expand_df(cancer)

contingency2(171, 389, 29, 386, method = "case.control")</pre>
```

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cosm_reg	Cosmetics for tables of regression coefficients. Converts tables gener-
	ated by tbl_regression to huxtable and adds some cosmetics.

### **Description**

Cosmetics for tables of regression coefficients. Converts tables generated by tbl\_regression to huxtable and adds some cosmetics.

### Usage

```
cosm_reg(gt_tbl, pad = 3, type = 3, bold = TRUE, head_label = "**Variable**")
```

### **Arguments**

gt\_tbl A table object generated by tbl\_regression.

pad Numerical, padding above and bellow rows.

type Anova's type to calculate global p-values.

bold Display labels in bold?

head\_label Character, label to be used as head for the variable's column.

#### Value

A huxtable.

```
require(sjlabelled, quietly = TRUE)
data(diet, package = "Epi")
diet <- diet |>
  var_labels(
    chd = "Coronary Heart Disease",
    fibre = "Fibre intake (g/day)"
model_binom <- glm(chd ~ fibre, data = diet, family = binomial)</pre>
model_binom |>
  tbl_regression(exponentiate = TRUE) |>
  cosm_reg(bold = TRUE) |>
  theme_pubh(1) |>
  add_footnote(get_r2(model_binom), font_size = 9)
data(birthwt, package = "MASS")
birthwt <- birthwt |>
  mutate(
   smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
```

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```
race = factor(race, labels = c("White", "African American", "Other"))
) |>
var_labels(
   bwt = "Birth weight (g)",
   smoke = "Smoking status",
   race = "Race"
)

model_norm <- lm(bwt ~ smoke + race, data = birthwt)

model_norm |>
   tbl_regression() |>
   cosm_reg(bold = TRUE) |>
   theme_pubh(1) |>
   add_footnote(get_r2(model_norm), font_size = 9)
```

cosm\_sum

Cosmetics for summary tables Adds some cosmetics to tables of descriptive statistics generated by tbl\_summary.

### Description

Cosmetics for summary tables Adds some cosmetics to tables of descriptive statistics generated by tbl\_summary.

#### Usage

```
cosm_sum(gt_tbl, pad = 3, bold = FALSE, head_label = "**Variable**")
```

#### **Arguments**

gt\_tbl A table object generated by tbl\_summary.

pad Numerical, padding above and bellow rows.

bold Display labels in bold?

head\_label Character, label to be used as head for the variable's column.

#### **Details**

Function cosm\_sum adds some cosmetics to tables generated by tbl\_summary, then converts the table as a huxtable and sets proper alignment.

#### Value

A huxtable.

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#### **Examples**

```
require(dplyr, quietly = TRUE)

data(Oncho)
Oncho |>
   select(-id) |>
   tbl_summary() |>
   cosm_sum(bold = TRUE) |>
   theme_pubh(1)
```

cross\_tbl

Table of descriptive statistics by categorical variable.

### **Description**

cross\_tbl is a wrapper to function from package tbl\_summary that constructs tables of descriptive statistics stratified by levels of a categorical outcome.

#### Usage

```
cross_tbl(
  data,
  by,
  head_label = " ",
  bold = TRUE,
  show_total = TRUE,
  p_val = FALSE,
  pad = 3,
  method = 2,
  ...
)
```

### **Arguments**

data A data frame where the variables in the formula can be found.

by The quoted name of the categorical variable (factor) used for the stratification.

head\_label Character, label to be used as head for the variable's column.

bold Display labels in bold?

show\_total Logical, show column with totals?

p\_val Logical, show p-values?

pad Numerical, padding above and bellow rows.

method An integer indicating methods for continuous variables. 1 Reports means and

standard deviations. 2 Reports medians and interquartile ranges.

... Additional arguments passed to tbl\_summary.

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#### **Details**

Function cross\_tbl is a relatively simple wrapper to function tbl\_summary. It constructs contingency tables and can also be used to report a table with descriptives for all variables stratified by one of the variables. Please see examples to see how to list variables. If data is labelled, the label of the stratifying variable is used as part of the header.

#### Value

A huxtable with descriptive statistics stratified by levels of the outcome.

#### See Also

```
tbl_summary
```

#### **Examples**

```
require(dplyr, quietly = TRUE)
#' data(Oncho)
## A two by two contingency table:
Oncho |>
  select(mf, area) |>
  cross_tbl(by = "mf", bold = TRUE) |>
  theme_pubh(2)
## Reporting prevalence:
Oncho |>
  select(mf, area) |>
  cross_tbl(by = "area", bold = TRUE) |>
  theme_pubh(2)
## Descriptive statistics for all variables in the \code{Oncho} data set except \code{id}.
Oncho |>
  select(-id) |>
  cross_tbl(by = "mf", bold = TRUE) |>
  theme_pubh(2)
```

diag\_test

Diagnostic tests from variables.

#### **Description**

diag\_test is a wrap function that calls epi.tests from package epiR. It computes sensitivity, specificity and other statistics related with screening tests.

```
diag_test(object = NULL, formula = NULL, data = NULL, ...)
```

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### **Arguments**

object	When chaining, this holds an object produced in the earlier portions of the chain.
	Most users can safely ignore this argument. See details and examples.
formula	A formula with shape: outcome ~ predictor (see details).
data	A data frame where the variables in the formula can be found.
	Further arguments passed to epi.tests.

#### **Details**

For the formula, the outcome is the gold standard and the explanatory variable is the new (screening) test. See examples.

#### See Also

```
epi.tests.
```

# **Examples**

```
## We compare the use of lung's X-rays on the screening of TB against the gold standard test.
Freq <- c(1739, 8, 51, 22)
BCG <- gl(2, 1, 4, labels = c("Negative", "Positive"))
Xray <- gl(2, 2, labels = c("Negative", "Positive"))
tb <- data.frame(Freq, BCG, Xray)
tb <- expand_df(tb)

tb |>
    diag_test(BCG ~ Xray)
```

diag\_test2

Diagnostic tests from direct input.

# Description

diag\_test2 is a wrap that calls epi.tests from package epiR. It computes sensitivity, specificity and other statistics related with screening tests.

#### Usage

```
diag_test2(aa, bb, cc, dd)
```

# Arguments

aa	Number of cases where both screening test and the gold standard are positive.
bb	Number of cases where screening test is positive but gold standard is negative.
сс	Number of cases where screening test is negative but gold standard is positive.
dd	Number of cases where both screening test and the gold standard are negative.

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### **Details**

diag.test uses direct input variables.

#### See Also

```
epi.tests.
```

### **Examples**

## We compare the use of lung's X-rays on the screening of TB against the gold standard test.  $diag_test2(22, 51, 8, 1739)$ 

estat

Descriptive statistics for continuous variables.

### **Description**

estat calculates descriptives of numerical variables.

# Usage

```
estat(object = NULL, formula = NULL, data = NULL, digits = 2, label = NULL)
```

# Arguments

object	When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples.
formula	A formula with shape: $\sim x$ or $\sim x   z$ (for groups).
data	A data frame where the variables in the formula can be found.
digits	Number of digits for rounding (default = 2).
label	Label used to display the name of the variable (see examples).

### Value

A data frame with descriptive statistics.

### See Also

```
summary, mytable.
```

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#### **Examples**

```
data(kfm, package = "ISwR")
require(sjlabelled, quietly = TRUE)
kfm <- kfm \mid >
  var_labels(
   dl.milk = "Breast-milk intake (dl/day)",
    sex = "Sex",
   weight = "Child weight (kg)",
   ml.suppl = "Milk substitute (ml/day)",
   mat.weight = "Maternal weight (kg)",
   mat.height = "Maternal height (cm)"
  )
kfm |>
  estat(~dl.milk)
estat(~ dl.milk | sex, data = kfm)
kfm |>
  estat(~ weight | sex)
```

expand\_df

Expand a data frame.

#### **Description**

expand\_df expands a data frame by a vector of frequencies.

#### Usage

```
expand_df(aggregate.data, index.var = "Freq", retain.freq = FALSE)
```

### **Arguments**

```
aggregate.data A data frame.

index.var A numerical variable with the frequencies (counts).

retain.freq Logical expression indicating if frequencies should be kept.
```

# **Details**

This is a generic function that resembles weighted frequencies in other statistical packages (for example, Stata). expand.df was adapted from a function developed by deprecated package epicalc (now package epiDisplay).

#### Value

An expanded data frame with replicates given by the frequencies.

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#### **Examples**

```
Freq <- c(5032, 5095, 41, 204)
Mortality <- gl(2, 2, labels = c("No", "Yes"))
Calcium <- gl(2, 1, 4, labels = c("No", "Yes"))
anyca <- data.frame(Freq, Mortality, Calcium)
anyca
anyca.exp <- expand_df(anyca)
with(anyca.exp, table(Calcium, Mortality))</pre>
```

**Fentress** 

Migraine pain reduction.

#### **Description**

Randomised control trial on children suffering from frequent and severe migraine. Control group represents untreated children. The active treatments were either relaxation alone or relaxation with biofeedback.

#### Usage

**Fentress** 

#### **Format**

A labelled tibble with 18 rows and 2 variables:

pain Reduction in weekly headache activity expressed as percentage of baseline data.

**group** Group, a factor with levels "Untreated", "Relaxation" (alone) and "Biofeedback" (relaxation and biofeedback).

#### Source

Fentress, DW, et al. (1986) Biofeedback and relaxation-response in the treatment of pediatric migraine. Dev Med Child Neurol 28:1 39-46.

Altman, DA (1991) Practical statistics for medical research. Chapman & Hall/CRC.

```
data(Fentress)
Fentress |>
  strip_error(pain ~ group)
```

gen\_bst\_df

frag	cont
ireq_	_COIIC

Relative and Cumulative Frequency.

### **Description**

freq\_cont tabulates a continuous variable by given classes.

### Usage

```
freq_cont(x, bks, dg = 2)
```

### **Arguments**

X	A numerical (continuous) variable. Ideally, relatively long (greater than 100 observations).
bks	Breaks defining the classes (see example).
dg	Number of digits for rounding (default = 2).

#### Value

A data frame with the classes, the mid-point, the frequencies, the relative and cumulative frequencies.

# **Examples**

```
data(IgM, package = "ISwR")
Ab <- data.frame(IgM)
estat(~IgM, data = Ab)
freq_cont(IgM, seq(0, 4.5, 0.5))</pre>
```

gen\_bst\_df

Generate a data frame with estimate and bootstrap CIs.

### **Description**

gen\_bst\_df is a function called that generates a data frame with confidence intervals of a continuous variable by levels of one or two categorical ones (factors).

```
gen_bst_df(object = NULL, formula = NULL, data = NULL, stat = "mean", ...)
```

gen\_bst\_df 23

### **Arguments**

object	When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples.
formula	A formula with shape: $y \sim x$ or $y \sim x \mid z$ where y is a numerical variable and both x and z are factors.
data	A data frame where the variables in the formula can be found.
stat	Statistic used for bst.
	Passes optional arguments to bst.

#### Value

A data frame with the confidence intervals by level.

```
data(kfm, package = "ISwR")
require(sjlabelled, quietly = TRUE)
kfm <- kfm |>
 var_labels(
   dl.milk = "Breast-milk intake (dl/day)",
   sex = "Sex",
   weight = "Child weight (kg)",
   ml.suppl = "Milk substitute (ml/day)",
   mat.weight = "Maternal weight (kg)",
   mat.height = "Maternal height (cm)"
  )
kfm |>
  gen_bst_df(dl.milk ~ sex)
data(birthwt, package = "MASS")
require(dplyr, quietly = TRUE)
birthwt <- mutate(birthwt,</pre>
  smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
  Race = factor(race > 1, labels = c("White", "Non-white"))
)
birthwt <- birthwt |>
  var_labels(
   bwt = "Birth weight (g)",
   smoke = "Smoking status"
gen_bst_df(bwt ~ smoke | Race, data = birthwt)
```

24 get\_r2

geo\_mean

Geometric mean.

# Description

Geometric mean.

### Usage

```
geo_mean(x)
```

### **Arguments**

Χ

A numeric variable with no negative values.

#### Value

A scalar, the calculated geometric mean.

### **Examples**

```
data(IgM, package = "ISwR")
Ab <- data.frame(IgM)
estat(~IgM, data = Ab)
geo_mean(IgM)</pre>
```

get\_r2

Estimate R2 or Pseudo-R2 from regression models

#### **Description**

get\_r2 is a is a wrap function that calls r2 from package performance. Calculates the R2 or pseudo-R2 value for different regression model objects, returning a character object for easy printing in tables of coefficients.

### Usage

```
get_r2(model, ...)
```

### **Arguments**

model A statistical regression model.
... Additional arguments passed to r2.

gf\_star 25

#### **Details**

The main purpose of get\_r2 is to allow easy printing of R2 value in tables of coefficients (see examples).

### See Also

r2.

### **Examples**

```
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)
data(birthwt, package = "MASS")
birthwt <- birthwt |>
  mutate(
    smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
   race = factor(race, labels = c("White", "African American", "Other"))
  var_labels(
   bwt = "Birth weight (g)",
   smoke = "Smoking status",
   race = "Race"
  )
model_norm <- lm(bwt ~ smoke + race, data = birthwt)</pre>
model_norm |>
  glm_coef(labels = model_labels(model_norm)) |>
  as_hux() |>
  set_align(everywhere, 2:3, "right") |>
  theme_pubh() |>
  add_footnote(get_r2(model_norm), font_size = 9)
```

gf\_star

Annotating a plot to display differences between groups.

### **Description**

gf\_star Is a function used to display differences between groups (see details).

```
gf_star(fig, x1, y1, x2, y2, y3, legend = "*", ...)
```

26 gf\_star

### **Arguments**

fig	A gformula object.
x1	Position in x for the start of the horizontal line.
y1	Position in y for the start of the vertical line, below to the horizontal line.
x2	Position in x for the end of the horizontal line.
y2	Position in y where the horizontal line is drawn.
у3	Position in y where the text is added.
legend	Character text used for annotating the plot.
	Additional information passed to gf_text.

#### **Details**

This function draws an horizontal line from coordinate (x1, y2) to coordinate (x2, y2). Draws vertical lines below the horizontal line, towards data, from (x1, y1) to (x1, y2) and from (x2, y1) to (x2, y2). Finally, adds text above the horizontal line, at the mid point between x1 and x2. See examples.

```
data(kfm, package = "ISwR")
require(sjlabelled, quietly = TRUE)
kfm <- kfm |>
 var_labels(
   dl.milk = "Breast-milk intake (dl/day)",
   sex = "Sex",
   weight = "Child weight (kg)",
   ml.suppl = "Milk substitute (ml/day)",
   mat.weight = "Maternal weight (kg)",
   mat.height = "Maternal height (cm)"
 )
kfm |>
 box_plot(dl.milk ~ sex, fill = "thistle", alpha = 0.8) |>
 gf_star(x1 = 1, y1 = 10.9, x2 = 2, y2 = 11, y3 = 11.2)
kfm |>
 box_plot(dl.milk ~ sex, fill = "thistle", alpha = 0.8) |>
 gf_star(1, 10.9, 2, 11, 11.4, legend = "p = 0.035", size = 2.5)
data(energy, package = "ISwR")
energy <- energy |>
 var_labels(
   expend = "Energy expenditure (MJ/day)",
   stature = "Stature"
 )
energy |>
 strip_error(expend ~ stature, col = "red") |>
 gf_star(1, 13, 2, 13.2, 13.4, "**")
```

glm\_coef 27

glm\_coef

Table of coefficients from generalised linear models.

### Description

glm\_coef displays estimates with confidence intervals and p-values from generalised linear models (see Details).

# Usage

```
glm_coef(
  model,
  digits = 2,
  alpha = 0.05,
  labels = NULL,
  se_rob = FALSE,
  type = "cond",
  exp_norm = FALSE
)
```

#### Arguments

model	A model from any of the classes listed in the details section.
digits	A scalar, number of digits for rounding the results (default = $2$ ).
alpha	Significant level (default = $0.05$ ) used to calculate confidence intervals.
labels	An optional character vector with the names of the coefficients (including intercept).
se_rob	Logical, should robust errors be used to calculate confidence intervals? (default = FALSE).
type	Character, either "cond" (condensed) or "ext" (extended). See details.
exp_norm	Logical, should estimates and confidence intervals should be exponentiated? (for family == "gaussian").

#### **Details**

glm\_coef recognises objects (models) from the following classes: clm, clogit, coxph, gee, glm, glmerMod, lm, lme, lmerMod, multinom, negbin, polr and surveg

For models from logistic regression (including conditional logistic, ordinal and multinomial), Poisson or survival analysis, coefficient estimates and corresponding confidence intervals are automatically exponentiated (back-transformed).

By default, glm\_coef uses naive standard errors for calculating confidence intervals but has the option of using robust standard errors instead.

glm\_coef can display two different data frames depending on the option of type, for type type = "cond" (the default), glm\_coef displays the standard table of coefficients with confidence intervals and p-values; for type = "ext", glm\_coef displays additional statistics including standard errors.

Please read the Vignette on Regression for more details.

28 harm\_mean

#### Value

A data frame with estimates, confidence intervals and p-values from glm objects.

#### **Examples**

```
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)
## Continuous outcome.
data(birthwt, package = "MASS")
birthwt <- birthwt |>
 mutate(
   smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
   race = factor(race, labels = c("White", "African American", "Other"))
 ) |>
 var_labels(
   bwt = "Birth weight (g)",
   smoke = "Smoking status",
   race = "Race"
 )
model_norm <- lm(bwt ~ smoke + race, data = birthwt)</pre>
glm_coef(model_norm, labels = model_labels(model_norm))
## Logistic regression.
data(diet, package = "Epi")
model_binom <- glm(chd ~ fibre, data = diet, family = binomial)</pre>
model_binom |>
 glm_coef(labels = c("Constant", "Fibre intake (g/day)"))
model_binom |>
 glm_coef(labels = c("Constant", "Fibre intake (g/day)"), type = "ext")
```

harm\_mean

Harmonic mean.

#### **Description**

Harmonic mean.

# Usage

harm\_mean(x)

### Arguments

Х

A numeric variable with no zero values.

hist\_norm 29

### Value

A scalar, the calculated harmonic mean.

# **Examples**

```
data(IgM, package = "ISwR")
Ab <- data.frame(IgM)
estat(~IgM, data = Ab)
harm_mean(IgM)</pre>
```

 $hist\_norm$ 

Histogram with Normal density curve.

# Description

hist\_norm constructs histograms and adds corresponding Normal density curve.

# Usage

```
hist_norm(
  object = NULL,
  formula = NULL,
  data = NULL,
  bins = 20,
  fill = "indianred3",
  color = "black",
  alpha = 0.4,
  ...
)
```

# Arguments

object	When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples.
formula	A formula with shape: $\sim y$ or $\sim y \mid x$ where y is a numerical variable and x is a factor.
data	A data frame where the variables in the formula can be found.
bins	Number of bins of the histogram.
fill	Colour to fill the bars of the histogram.
color	Colour used for the border of the bars.
alpha	Opacity $(0 = invisible, 1 = opaque)$ .
	Further arguments passed to gf_dhistogram.

30 Hodgkin

#### **Examples**

```
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)
data(birthwt, package = "MASS")
birthwt <- birthwt |>
  mutate(
    smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
    Race = factor(race > 1, labels = c("White", "Non-white"))
) |>
  var_labels(
    bwt = "Birth weight (g)",
    smoke = "Smoking status"
)
birthwt |>
  hist_norm(~bwt, alpha = 0.7, bins = 20, fill = "cadetblue")
birthwt |>
  hist_norm(~ bwt | smoke, alpha = 0.7, bins = 20, fill = "cadetblue")
```

Hodgkin

T-cell counts from Hodgkin's disease patients.

# Description

Number of CD4+ T-cells and CD8+ T-cells in blood samples from patients in remission from Hodgkin's disease or in remission from disseminated malignancies.

### Usage

Hodgkin

#### **Format**

A labelled tibble with 40 rows and 3 variables:

**CD4** Concentration of CD4+ T-cells (cells / mm<sup>3</sup>).

**CD8** Concentration of CD8+ T-cells (cells / mm<sup>3</sup>).

Group Group, factor with levels "Non-Hodgkin" and "Hodgkin".

#### **Source**

Shapiro, CM, et al (1986) Immunologic status of patients in remission from Hodgkin's disease and disseminated malignancies. Am J Med Sci 293:366-370.

Altman, DA (1991) Practical statistics for medical research. Chapman & Hall/CRC.

inv\_logit 31

#### **Examples**

```
data(Hodgkin)
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)

Hodgkin <- Hodgkin |>
    mutate(
      Ratio = CD4 / CD8
)    |>
    var_labels(
      Ratio = "CD4+ / CD8+ T-cells"
)

estat(~ Ratio | Group, data = Hodgkin)

Hodgkin |>
    qq_plot(~ Ratio | Group)

Hodgkin$Ratio <- Hodgkin$CD4 / Hodgkin$CD8
estat(~ Ratio | Group, data = Hodgkin)

qq_plot(~ Ratio | Group, data = Hodgkin)</pre>
```

inv\_logit

Inverse of the logit

#### **Description**

inv\_logit Calculates the inverse of the logit (probability in logistic regression)

### Usage

```
inv_logit(x)
```

#### **Arguments**

Х

Numerical value used to compute the inverse of the logit.

jack\_knife

Ranks leverage observations from Jackknife method.

#### **Description**

jack\_knife Ranks the squared differences between mean values from Jackknife analysis (arithmetic mean estimated by removing one observation at a time) and the original mean value.

32 Kirkwood

### Usage

```
jack_knife(x)
```

# Arguments

Х

A numeric variable. Missing values are removed by default.

#### Value

Data frame with the ranked squared differences.

### See Also

```
rank_leverage.
```

### **Examples**

```
x <- rnorm(10, 170, 8)
x
mean(x)
jack_knife(x)

x <- rnorm(100, 170, 8)
mean(x)
head(jack_knife(x))</pre>
```

Kirkwood

Body weight and plasma volume.

# Description

Body weight and plasma volume in eight healthy men.

### Usage

Kirkwood

### **Format**

A labelled data frame with 8 rows and 3 variables:

```
subject Subject ID.weight Body weight in kg.volume Plasma volume in litres.
```

### Source

Kirkwood, BR and Sterne, JAC (2003) Essential Medical Statistics. Second Edition. Blackwell.

knife\_mean 33

### **Examples**

```
data(Kirkwood)

Kirkwood |>
   gf_point(volume ~ weight) |>
   gf_lm(col = "indianred3", interval = "confidence", fill = "indianred3")
```

knife\_mean

Jackknife for means.

# Description

knife\_mean is an internal function. Calculates arithmetic means by removing one observation at a time.

# Usage

```
knife_mean(x)
```

### **Arguments**

Х

A numerical variable. Missing values are removed for the mean calculation.

### Value

A vector with the mean calculations.

#### **Examples**

```
x <- rnorm(10, 170, 8)
x
mean(x)
knife_mean(x)</pre>
```

leverage

Leverage.

# Description

leverage is an internal function called by rank\_leverage.

```
leverage(x)
```

34 logistic\_gof

#### **Arguments**

Х

A numeric variable. Missing values are removed by default.

#### **Details**

Estimates the leverage of each observation around the arithmetic mean.

#### Value

Variable with corresponding leverage estimations

#### **Examples**

```
x <- rnorm(10, 170, 8)
x
mean(x)
leverage(x)
rank_leverage(x)</pre>
```

logistic\_gof

Goodness of fit for Logistic Regression.

### **Description**

logistic\_gof performs the Hosmer and Lemeshow test to test the goodness of fit of a logistic regression model. This function is part of residuals.lrm from package rms.

### Usage

```
logistic_gof(model)
```

#### **Arguments**

model

A logistic regression model object.

### Author(s)

Frank Harell, Vanderbilt University <f.harrell@vanderbilt.edu>

#### References

Hosmer DW, Hosmer T, Lemeshow S, le Cessie S, Lemeshow S. A comparison of goodness-of-fit tests for the logistic regression model. Stat in Med 16:965–980, 1997.

```
data(diet, package = "Epi")
model <- glm(chd ~ fibre, data = diet, family = binomial)
glm_coef(model, labels = c("Constant", "Fibre intake (g/day)"))
logistic_gof(model)</pre>
```

Macmahon 35

Macmahon

Breast cancer and age of childbirth.

### Description

An international case-control study to test the hypothesis that breast cancer is related to the age that a woman gives childbirth.

#### Usage

Macmahon

#### **Format**

A labelled tibble with 185 rows and 2 variables:

```
cancer Diagnosed with breast cancer, a factor with levels "No" and "Yes". age Age mother gives childbirth, factor with levels "<20", "20-24", "25-29", "30-34" and ">34".
```

#### **Source**

Macmahon, B. et al. (1970). Age at first birth and breast cancer risk. Bull WHO 43, 209-221.

# **Examples**

```
require(dplyr, quietly = TRUE)
data(Macmahon)

Macmahon |>
  select(cancer, age) |>
  cross_tbl(by = "cancer")
```

mhor

Mantel-Haenszel odds ratio.

### **Description**

mhor computes odds ratios by levels of the stratum variable as well as the Mantel-Haenszel pooled odds ratio. The test for effect modification (test for interaction) is also displayed.

```
mhor(object = NULL, formula = NULL, data = NULL)
```

36 model\_labels

# Arguments

object When chaining, this holds an object produced in the earlier portions of the chain.

Most users can safely ignore this argument. See details and examples.

formula A formula with shape: outcome ~ stratum/exposure.

data A data frame containing the variables used in formula.

#### Value

Odds ratios with 95 outcome by levels of stratum. The Mantel-Haenszel pooled OR and the test for effect modification is also reported.

### **Examples**

```
data(oswego, package = "epitools")
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)
oswego <- oswego |>
 mutate(
   ill = factor(ill, labels = c("No", "Yes")),
   sex = factor(sex, labels = c("Female", "Male")),
   chocolate.ice.cream = factor(chocolate.ice.cream, labels = c("No", "Yes"))
 ) |>
 var_labels(
   ill = "Developed illness",
   sex = "Sex",
    chocolate.ice.cream = "Consumed chocolate ice cream"
 )
oswego |>
 select(ill, sex, chocolate.ice.cream) |>
 tbl_summary() |>
 cosm_sum() |>
 theme_pubh()
oswego |>
 mhor(ill ~ sex / chocolate.ice.cream)
```

model\_labels

Using labels as coefficient names in tables of coefficients.

### Description

model\_labels replaces row names in glm\_coef with labels from the original data frame.

```
model_labels(model, intercept = TRUE)
```

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# **Arguments**

model A generalised linear model.

intercept Logical, should the intercept be added to the list of coefficients?

#### **Details**

model\_labels does not handle yet interaction terms, see examples.

Please read the Vignette on Regression for more examples.

# **Examples**

```
require(dplyr, quietly = TRUE)
require(sjlabelled, quietly = TRUE)
data(birthwt, package = "MASS")
birthwt <- birthwt |>
  mutate(
    smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
    race = factor(race, labels = c("White", "African American", "Other"))
  ) |>
  var_labels(
    bwt = "Birth weight (g)",
    smoke = "Smoking status",
    race = "Race"
  )
model_norm <- lm(bwt ~ smoke + race, data = birthwt)</pre>
glm_coef(model_norm, labels = model_labels(model_norm))
model_int <- lm(formula = bwt ~ smoke * race, data = birthwt)</pre>
model_int |>
  glm\_coef(labels = c(
    model_labels(model_int),
    "Smoker: African American",
    "Smoker: Other"
  ))
```

multiple

Multiple comparisons with plot.

## **Description**

multiple displays results from post-doc analysis and constructs corresponding plot.

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## Usage

```
multiple(
  model,
  formula,
  adjust = "mvt",
  type = "response",
  reverse = TRUE,
  level = 0.95,
  digits = 2,
  ...
)
```

# **Arguments**

model	A fitted model supported by emmeans, such as the result of a call to aov, $lm$ , $glm$ , etc.
formula	A formula with shape: $\sim$ y or $\sim$ y x (for interactions). Where y is the term of the model on which comparisons are made and x is a term interacting with y.
adjust	Method to adjust CIs and p-values (see details).
type	Type of prediction (matching "linear.predictor", "link", or "response").
reverse	Logical argument. Determines the direction of comparisons.
level	Confidence interval significance level.
digits	Number of digits for rounding (default = 2).
	Further arguments passed to emmeans.

# **Details**

The default adjusting method is "mvt" which uses the multivariate t distribution. Other options are: "bonferroni", "holm", "hochberg", "tukey" and "none". The default option for argument reverse is to make reverse comparisons, i.e., against the reference level matching comparisons from 1m and g1m.

## Value

A list with objects: df A data frame with adjusted p-values, fig\_ci a plot with estimates and adjusted confidence intervals and fig\_pval a plot comparing adjusted p-values.

# See Also

```
emmeans, pwpp.
```

```
data(birthwt, package = "MASS")
birthwt$race <- factor(birthwt$race, labels = c("White", "African American", "Other"))
model_1 <- aov(bwt ~ race, data = birthwt)</pre>
```

odds\_trend 39

```
multiple(model_1, ~race)$df
multiple(model_1, ~race)$fig_ci |>
    gf_labs(y = "Race", x = "Difference in birth weights (g)")
multiple(model_1, ~race)$fig_pval |>
    gf_labs(y = "Race")
```

 $odds\_trend$ 

Function to calculate OR using Wald CI, and plot trend.

## **Description**

odds\_trend calculates the odds ratio with confidence intervals (Wald) for different levels (three or more) of the exposure variable, constructs the corresponding plot and calculates if the trend is significant or not.

# Usage

```
odds_trend(formula, data, angle = 45, hjust = 1, method = "wald", ...)
```

# **Arguments**

formula	A formula with shape: outcome ~ exposure.	
data	A data frame where the variables in the formula can be found.	
angle	Angle of for the x labels (default = $45$ ).	
hjust	Horizontal adjustment for x labels (default = $1$ ).	
method	Method for calculating confidence interval around odds ratio.	
	Passes optional arguments to oddsratio.	

## **Details**

```
odds_trend is a wrap function that calls oddsratio from package epitools.

Additional methods for confidence intervals include: "midp", "fisher", and "small".
```

#### Value

A list with components df a data frame with the results and fig corresponding plot.

#### See Also

oddsratio.

40 Oncho

## **Examples**

## A cross-sectional study looked at the association between obesity and a biopsy resulting ## from mammography screening.

```
Freq <- c(3441, 34, 39137, 519, 20509, 280, 12149, 196, 11882, 199)
Biopsy <- gl(2, 1, 10, labels = c("No", "Yes"))
Weight <- gl(5, 2, 10, labels = c(
    "Underweight", "Normal", "Over (11-24%)",
    "Over (25-39%)", "Over (> 39%)"
))
breast <- data.frame(Freq, Biopsy, Weight)
breast
breast <- expand_df(breast)
require(sjlabelled, quietly = TRUE)

breast <- var_labels(breast,
    Weight = "Weight group"
)

odds_trend(Biopsy ~ Weight, data = breast)$df
odds_trend(Biopsy ~ Weight, data = breast)$fig</pre>
```

0ncho

Onchocerciasis in Sierra Leone.

# **Description**

Study of onchocerciasis ("river blindness") in Sierra Leone, in which subjects were classified according to whether they lived in villages in savannah or rainforest area.

#### Usage

**Oncho** 

## **Format**

A labelled tibble with 1302 rows and 7 variables:

```
id Subject ID.
```

mf Infected with Onchocerciasis volvulus, factor with levels "Not-infected" and "Infected".

area Area of residence, factor with levels "Savannah" and "Rainforest".

agegrp Age group in years, factor with levels "5-9", "10-19", "20-39" and "40+".

sex Subject sex, factor with levels "Male" and "Female".

mfload Microfiliariae load.

lesions Severe eye lesions, factor with levels "No" and "Yes".

predict\_inv 41

## **Source**

McMahon, JE, Sowa, SIC, Maude, GH and Kirkwood BR (1988) Onchocerciasis in Sierra Leone 2: a comparison of forest and savannah villages. Trans Roy Soc Trop Med Hyg 82: 595-600.

Kirkwood, BR and Sterne, JAC (2003) Essential Medical Statistics. Second Edition. Blackwell.

# **Examples**

```
require(dplyr, quietly = TRUE)
data(Oncho)

Oncho |>
  select(mf, agegrp) |>
  cross_tbl(by = "mf")
```

predict\_inv

Given y solve for x in a simple linear model.

## **Description**

predict\_inv Calculates the value the predictor x that generates value y with a simple linear model.

## Usage

```
predict_inv(model, y)
```

# Arguments

model A simple linear model object (class lm).

y A numerical scalar, the value of the outcome for which we want to calculate the

predictor x.

#### Value

The estimated value of the predictor.

# **Examples**

## Spectrophotometry example. Titration curve for riboflavin (nmol/ml). The sample has an absorbance ## of 1.15. Aim is to estimate the concentration of riboflavin in the sample.

```
Riboflavin <- seq(0, 80, 10)
OD <- 0.0125 * Riboflavin + rnorm(9, 0.6, 0.03)
titration <- data.frame(Riboflavin, OD)

require(sjlabelled, quietly = TRUE)
titration <- titration |>
    var_labels(
    Riboflavin = "Riboflavin (nmol/ml)",
```

prop\_or

```
OD = "Optical density"
)

titration |>
    gf_point(OD ~ Riboflavin) |>
    gf_smooth(col = "indianred3", se = TRUE, lwd = 0.5, method = "loess")

## Model with intercept different from zero:
model <- lm(OD ~ Riboflavin, data = titration)
glm_coef(model)

predict_inv(model, 1.15)</pre>
```

prop\_or

Proportion, p1 from proportion p2 and OR.

# Description

prop\_or is a simple function to calculate a proportion, from another proportion and the odds ratio between them.

## Usage

```
prop_or(p2, or)
```

## **Arguments**

p2 The value of a proportion in the unexposed group (p2).

or The odds ratio of p1/p2.

# Value

p1, the proportion in the exposed group (p1).

```
flu <- matrix(c(20, 80, 220, 140), nrow = 2)
colnames(flu) <- c("Yes", "No")
rownames(flu) <- c("Vaccine", "Placebo")
flu

or <- (20 * 140) / (80 * 220)
p2 <- 80 / 220
prop_or(p2 = p2, or = or)
20 / 240</pre>
```

pseudo\_r2 43

pseudo\_r2

Pseudo R2 (logistic regression) pseudo\_r2 Calculates R2 analogues (pseudo R2) of logistic regression.

## **Description**

Pseudo R2 (logistic regression) pseudo\_r2 Calculates R2 analogues (pseudo R2) of logistic regression.

# Usage

```
pseudo_r2(model)
```

## **Arguments**

model

A logistic regression model.

# **Details**

pseudo\_r2 calculates three pseudo R2 of logistic regression models: 1) Nagelkerke, @0 Cox and Snell, 3) Hosmer and Lemeshow.

#### Value

A data frame with the calculated pseudo R2 values.

# **Examples**

```
\label{lem:data_oncho} $$ \mbox{model_oncho} <- glm(mf ~ area, data = Oncho, binomial) $$ glm_coef(model_oncho, labels = c("Constant", "Area (rainforest/savannah)")) $$ pseudo_r2(model_oncho) $$
```

qq\_plot

Quantile-quantile plots against the standard Normal distribution.

# **Description**

qq\_plot constructs quantile-quantile plots against the standard normal distribution (also known as quantile-normal plots).

44 qq\_plot

## Usage

```
qq_plot(
  object = NULL,
  formula = NULL,
  data = NULL,
  pch = 20,
  col = "indianred3",
  ylab = NULL,
   ...
)
```

# Arguments

object When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples. formula A formula with shape:  $\sim x$  or  $\sim x \mid z$  where x is a numerical variable and z is a factor. A data frame where the variables in the formula can be found. data pch Point character passed to gf\_qq. Colour of the reference line, passed to gf\_line. col Optional character passed as label for the y-axis. ylab Further arguments passed to gf\_qq. . . .

```
data(kfm, package = "ISwR")
require(sjlabelled, quietly = TRUE)
kfm <- kfm |>
   var_labels(
      dl.milk = "Breast-milk intake (dl/day)",
      sex = "Sex",
      weight = "Child weight (kg)",
      ml.suppl = "Milk substitute (ml/day)",
      mat.weight = "Maternal weight (kg)",
      mat.height = "Maternal height (cm)"
)
kfm |>
   qq_plot(~dl.milk)
```

rank\_influence 45

rank\_influence

Ranks observations based upon influence measures on models.

# Description

rank\_influence calculates influence measures of each data observation on models and then ranks them.

# Usage

```
rank_influence(model)
```

## **Arguments**

model

A generalised linear model object.

## **Details**

rank\_influence is a wrap function that calls influence.measures, ranks observations on their significance influence on the model and displays the 10 most influential observations (if they are significant).

## See Also

```
influence.measures.
```

# **Examples**

```
data(diet, package = "Epi")
model <- glm(chd ~ fibre, data = diet, family = binomial)
rank_influence(model)</pre>
```

rank\_leverage

Ranks observations by leverage.

## **Description**

rank\_leverage ranks observations by their leverage (influence) on the arithmetic mean.

## Usage

```
rank_leverage(x)
```

## **Arguments**

Χ

A numeric variable. Missing values are removed by default.

46 reference\_range

## Value

A data frame ranking observations by their leverage around the mean.

## See Also

```
jack_knife.
```

## **Examples**

```
x <- rnorm(10, 170, 8)
x
mean(x)
rank_leverage(x)

x <- rnorm(100, 170, 8)
mean(x)
head(rank_leverage(x))</pre>
```

reference\_range

Reference range (reference interval).

# Description

reference\_range estimates the reference range (reference interval) of a numerical variable.

# Usage

```
reference_range(avg, std)
```

## **Arguments**

avg The arithmetic mean (a scalar numerical value).

std The standard deviation (a scalar numerical value).

## **Details**

The reference range assumes normality and represents the limits that would include 95 observations.

# Value

A data frame with the reference range limits.

```
x <- rnorm(100, 170, 8)
round(mean(x), 2)
round(sd(x), 2)
round(reference_range(mean(x), sd(x)), 2)</pre>
```

rel\_dis 47

rel\_dis

Relative Dispersion.

# **Description**

Calculates the coefficient of variation (relative dispersion) of a variable. The relative dispersion is defined as the standard deviation over the arithmetic mean.

# Usage

```
rel_dis(x)
```

# **Arguments**

Х

A numerical variable. NA's observations are removed by default.

#### Value

The coefficient of variation (relative dispersion).

# **Examples**

```
height <- rnorm(100, 170, 8)
rel_dis(height)</pre>
```

Roberts

Extracorporeal membrane oxygenation in neonates.

# Description

A clinical trial on the value of extracorporeal membrane oxygenation for term neonates with severe respiratory failure. RCT compares active treatment against conventional management.

# Usage

Roberts

#### **Format**

A labelled tibble with 185 rows and 2 variables:

**emo** Extracorporeal membrane oxygenation treatment, factor with levels "No" and "Yes". **survived** One year survival, factor with levels "No" and "Yes".

48 Rothman

#### **Source**

Roberts, TE (1998) Extracorporeal Membrane Oxygenation Economics Working Group. Economic evaluation and randomised controlled trial of extracorporeal membrane oxygenation: UK collaborative trial. Brit Med J 317:911-16.

## **Examples**

```
require(dplyr, quietly = TRUE)
data(Roberts)

Roberts |>
  select(survived, emo) |>
  cross_tbl(by = "survived")
```

Rothman

Oral contraceptives and stroke.

## **Description**

A case-control study of oral contraceptives and stroke in young women with presence or absence of hypertension. Cases represent thrombotic stroke and controls are hospital controls. The group of no hypertension includes normal blood pressure (<140/90 mm Hg) and borderline hypertension (140-159/90-94 mm Hg). Hypertension group includes moderate hypertension (160-179/95-109 mm Hg) and severe hypertension (180+/110+ mm Hg). This data has been used as an example of join exposure by Rothman for measuring interactions (see examples).

## Usage

Rothman

#### **Format**

A labelled tibble with 477 rows and 3 variables:

stroke Thrombotic stroke, factor with levels "No" and "Yes".

oc Current user of oral contraceptives, factor with levels "Non-user" and "User".

ht Hypertension, factor with levels "No" (<160/95 mm Hg) and "Yes".

#### Source

Collaborative Group for the Study of Stroke in Young Women (1975) Oral contraceptives and stroke in young women. JAMA 231:718-722.

Rothman, KJ (2002) Epidemiology. An Introduction. Oxford University Press.

round\_pval 49

## **Examples**

```
require(dplyr, quietly = TRUE)
data(Rothman)
Rothman |>
  select(stroke, oc, ht) |>
  tbl_summary() |>
  cosm_sum()
mhor(stroke ~ ht / oc, data = Rothman)
## Model with standard interaction term:
model1 <- glm(stroke ~ ht * oc, data = Rothman, family = binomial)</pre>
glm_coef(model1)
## Model considering join exposure:
Rothman$join <- 0
Rothman$join[Rothman$oc == "Non-user" & Rothman$ht == "Yes"] <- 1</pre>
Rothman$join[Rothman$oc == "User" & Rothman$ht == "No"] <- 2</pre>
Rothman$join[Rothman$oc == "User" & Rothman$ht == "Yes"] <- 3</pre>
Rothman$join <- factor(Rothman$join, labels = c(</pre>
  "Unexposed", "Hypertension", "OC user",
  "OC and hypertension"
))
require(sjlabelled, quietly = TRUE)
Rothman$join <- set_label(Rothman$join, label = "Exposure")</pre>
Rothman |>
  select(stroke, join) |>
  cross_tbl(by = "stroke")
model2 <- glm(stroke ~ join, data = Rothman, family = binomial)</pre>
glm_coef(model2)
```

round\_pval

Rounding p-values.

## **Description**

round\_pval is an internal function called by glm\_coef to round p-values from model coefficients.

## Usage

```
round_pval(pval)
```

# Arguments

pval

vector of p-values, numeric.

Sharples

Sandler

Passive smoking in adulthood and cancer risk.

# Description

A case-control study to investigate the effects of passive smoking on cancer. Passive smoking was defined as exposure to the cigarette smoke of a spouse who smoked at least one cigarette per day for at least 6 months.

# Usage

Sandler

## **Format**

A labelled tibble with 998 rows and 3 variables:

```
passive Passive smoker, factor with levels "No" and "Yes".cancer Diagnosed with cancer, factor with levels "No" and "Yes".smoke Active smoker, factor with levels "No" and "Yes".
```

## **Source**

Sandler, DP, Everson, RB, Wilcox, AJ (1985). Passive smoking in adulthood and cancer risk. Amer J Epidem, 121: 37-48.

# **Examples**

```
require(dplyr, quietly = TRUE)
data(Sandler)

Sandler |>
   select(cancer, passive) |>
   cross_tbl(by = "cancer")

mhor(cancer ~ smoke / passive, data = Sandler)
```

Sharples

Measured and self-reported weight in New Zealand.

## **Description**

Data on measured and self-reported weight from 40–50 year old participants in the 1989/1990 Life In New Zealand Survey.

ss\_jk 51

## **Usage**

Sharples

#### **Format**

```
A tibble with 343 rows and 4 variables:
```

```
srweight Self-reported weight in kg.
```

weight Measured weight in kg.

srbmi Body mass index calculated from self-reported weight and self-reported height in kg/m^2.

**mbmi** Body mass index calculated from measured weight and measured height in kg/m^2.

## **Source**

Sharples, H, et al. (2012) Agreement between measured and self-reported height, weight and BMI in predominantly European middle-aged New Zealanders: findings from a nationwide 1989 survey. New Zealand Med J 125: 60-69.

## **Examples**

```
Sharples |>
  bland_altman(srweight ~ weight, transform = TRUE) |>
  gf_labs(x = "Mean of weights (kg)", y = "Measured weight / Self-reported weight")
```

ss\_jk

Sum of squares for Jackknife.

# **Description**

ss\_jk is an internal function called by jack\_knife. It calculates the squared difference of a numerical variable around a given value (for example, the mean).

# Usage

```
ss_jk(obs, stat)
```

# Arguments

obs A numerical vector with no missing values (NA's).

Stat The value of the statistic that is used as a reference.

#### Value

The squared difference between a variable and a given value.

strip\_error

## **Examples**

```
x <- rnorm(10, 170, 8)
x
mean(x)
ss_jk(x, mean(x))
jack_knife(x)</pre>
```

stats\_quotes

Internal function to calculate descriptive statistics.

# Description

stats\_quotes is an internal function called by estat.

# Usage

```
stats_quotes(x, data2, digits = 2)
```

# Arguments

x a numeric variabledata2 A data frame where x can be found.digits Number of digits for rounding.

strip\_error

Strip plots with error bars.

# **Description**

strip\_error constructs strip plots with error bars showing 95 confidence intervals around mean values.

# Usage

```
strip_error(
  object = NULL,
  formula = NULL,
  data = NULL,
  pch = 20,
  size = 1,
  alpha = 0.7,
  col = "indianred3",
  ...
)
```

strip\_error 53

## Arguments

object When chaining, this holds an object produced in the earlier portions of the chain. Most users can safely ignore this argument. See details and examples. formula A formula with shape:  $y \sim x$  or  $y \sim x \mid z$  where y is a numerical variable and both x and z are factors. data A data frame where the variables in the formula can be found. Point character passed to gf\_point or gf\_jitter. pch Size of the symbol (pch) for representing data values. size Opacity of the symbol (0 = invisible, 1 = opaque). alpha A colour or formula used for mapping colour. col Additional information passed to gf\_jitter or gf\_point.

```
data(energy, package = "ISwR")
require(sjlabelled, quietly = TRUE)
energy <- energy |>
 var_labels(
   expend = "Energy expenditure (MJ/day)",
   stature = "Stature"
 )
energy |>
 strip_error(expend ~ stature, col = "red")
t.test(expend ~ stature, data = energy)
## Adding an horizontal line to show significant difference:
energy |>
 strip_error(expend ~ stature, col = "red") |>
 gf_star(1, 13, 2, 13.2, 13.4, "**")
data(birthwt, package = "MASS")
require(dplyr, quietly = TRUE)
birthwt <- birthwt |>
 mutate(
   smoke = factor(smoke, labels = c("Non-smoker", "Smoker")),
   Race = factor(race > 1, labels = c("White", "Non-white"))
 ) |>
 var_labels(
   bwt = "Birth weight (g)",
   smoke = "Smoking status"
 )
 strip_error(bwt ~ smoke | Race, col = "darksalmon")
birthwt |>
 strip_error(bwt ~ smoke, col = ~Race)
```

54 Thall

```
birthwt |>
   strip_error(bwt ~ smoke, pch = ~Race, col = ~Race)
birthwt |>
   strip_error(bwt ~ smoke | Race)
```

Thall

RCT on the treatment of epilepsy.

# **Description**

Randomised control trial of an antiepilectic drug (prograbide), in which the number of seizures of 59 patients at baseline and other four follow-up visits were recorded.

# Usage

Thall

## **Format**

A tibble with 59 rows and 8 variables:

id Subject ID.

treat Treatment, factor with levels "Control" and "Prograbide".

base Number of seizures at baseline.

age Age in years at baseline.

- y1 Number of seizures at year one follow-up.
- y2 Number of seizures at year two follow-up.
- y3 Number of seizures at year three follow-up.
- y4 Number of seizures at year four follow-up.

#### Source

Thall, PF and Vail, SC (1990) Some covariance models for longitudinal count data with over-dispersion. Biometrics, 46: 657-671.

Stukel, TA (1993) Comparison of methods for the analysis of longitudinal data. Statistics Med 12: 1339-1351.

Shoukri, MM and Chaudhary, MA (2007) Analysis of correlated data with SAS and R. Third Edition. Chapman & Hall/CRC.

theme\_pubh 55

## **Examples**

```
data(Thall)

c1 <- cbind(Thall[, c(1:5)], count = Thall$y1)[, c(1:4, 6)]

c2 <- cbind(Thall[, c(1:4, 6)], count = Thall$y2)[, c(1:4, 6)]

c3 <- cbind(Thall[, c(1:4, 7)], count = Thall$y3)[, c(1:4, 6)]

c4 <- cbind(Thall[, c(1:4, 8)], count = Thall$y3)[, c(1:4, 6)]

epilepsy <- rbind(c1, c2, c3, c4)

require(lme4, quietly = TRUE)

model_glmer <- glmer(count ~ treat + base + I(age - mean(age, na.rm = TRUE)) +

(1 | id), data = epilepsy, family = poisson)

glm_coef(model_glmer, labels = c(
    "Treatment (Prograbide/Control)",
    "Baseline count", "Age (years)"

))</pre>
```

theme\_pubh

A theme for huxtables This function quickly set a default style for a huxtable.

## **Description**

A theme for huxtables This function quickly set a default style for a huxtable.

#### Usage

```
theme_pubh(ht, rw = 1)
```

#### **Arguments**

ht A huxtable object.

rw A numeric vector with the rows on which a bottom border is desired.

## **Details**

theme\_pubh is a variation of theme\_article with the added flexibility of adding a bottom border line at desired row numbers.

```
require(dplyr, quietly = TRUE)
data(Oncho)

Oncho |>
   select(area, mf) |>
   cross_tbl(by = "area") |>
   theme_pubh(2)
```

56 Tuzson

```
data(Bernard)

t1 <- estat(~ apache | fate, data = Bernard)
t2 <- estat(~ o2del | fate, data = Bernard)
rbind(t1, t2) |>
  as_hux() |>
  theme_pubh(c(1, 3))
```

Tuzson

Peak knee velocity in walking at flexion and extension.

# **Description**

Data of peak knee velocity in walking at flexion and extension in studies about functional performance in cerebral palsy.

# Usage

Tuzson

#### **Format**

A labelled tibble with 18 rows and 2 variables:

```
flexion Peak knee velocity in gait: flexion (degree/s). extension Peak knee velocity in gait: extension (degree/s).
```

# Source

Tuzson, AE, Granata, KP, and Abel, MF (2003) Spastic velocity threshold constrains functional performance in cerebral palsy. Arch Phys Med Rehabil 84: 1363-1368.

```
data(Tuzson)

Tuzson |>
   gf_point(flexion ~ extension)

cor.test(~ flexion + extension, data = Tuzson)
```

Vanderpump 57

Vanderpump

Smoking and mortality in Whickham, England.

# **Description**

Data represents women participating in a health survey in Whickham, England in 1972-1974.

# Usage

Vanderpump

#### **Format**

A labelled tibble with 1314 rows and 3 variables:

```
vstatus Vitality status, factor with levels "Alive" and "Death".

smoker Smoking status, factor with levels "Non-smoker" and "Smoker".

agegrp Age group, factor with levels "18-44", "45-64" and "64+".
```

#### **Source**

Vanderpump, MP, et al (1996) Thyroid, 6:155-160.

Appleton, DR, French, JM and Vanderpump, PJ (1996) Ignoring a covariate: An example of Simpson's paradox. The American Statistician 50:340-341.

Vittinghoff, E, Glidden, DV, Shiboski, SC and McCulloh, CE (2005) Regression methods in Biostatistics. Springer.

```
require(dplyr, quietly = TRUE)
data(Vanderpump)

Vanderpump |>
    select(vstatus, agegrp, smoker) |>
    tbl_strata(
        strata = smoker,
        .tbl_fun = ~ .x |>
            tbl_summary(by = agegrp)
    ) |>
    cosm_sum(bold = TRUE, head_label = " ")

mhor(vstatus ~ agegrp / smoker, data = Vanderpump)
```

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