

Package ‘faux’

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Title Simulation for Factorial Designs

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Description Create datasets with factorial structure through simulation by specifying variable parameters. Extended documentation at <https://debruine.github.io/faux/>. Described in DeBruine (2020) [doi:10.5281/zenodo.2669586](https://doi.org/10.5281/zenodo.2669586).

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add_between	<i>Add between factors</i>
-------------	----------------------------

Description

Add between factors

Usage

```
add_between(.data, .by = NULL, ..., .shuffle = FALSE, .prob = NULL)
```

Arguments

.data	the data frame
.by	the grouping column (groups by row if NULL)
...	the names and levels of the new factors
.shuffle	whether to assign cells randomly or in "order"
.prob	probability of each level, equal if NULL

Value

data frame

Examples

```
add_random(subj = 4, item = 2) %>%
  add_between("subj", condition = c("cntl", "test")) %>%
  add_between("item", version = c("A", "B"))
```

add_contrast	<i>Add a contrast to a data frame</i>
--------------	---------------------------------------

Description

Add a contrast to a data frame

Usage

```
add_contrast(
  data,
  col,
  contrast = c("anova", "sum", "treatment", "helmert", "poly", "difference"),
  levels = NULL,
  ...,
  add_cols = TRUE,
  colnames = NULL
)
```

Arguments

data	the data frame
col	the column to recode
contrast	the contrast to recode to
levels	the levels of the factor in order
...	arguments to pass to the contrast function (base or omit)
add_cols	whether to just add the contrast to the existing column or also to create new explicit columns in the dataset (default)
colnames	optional list of column names for the added contrasts

Value

the data frame with the recoded column and added columns (if add_cols == TRUE)

Examples

```
df <- sim_design(between = list(time = 1:6), plot = FALSE) %>%
  add_contrast("time", "poly")

# test all polynomial contrasts
lm(y ~ time, df) %>% broom::tidy()

# test only the linear and quadratic contrasts
lm(y ~ `time^1` + `time^2`, df) %>% broom::tidy()
```

add_random

*Add random factors to a data structure***Description**

Add random factors to a data structure

Usage

```
add_random(.data = NULL, ..., .nested_in = NULL)
```

Arguments

<code>.data</code>	the data frame
<code>...</code>	the new random factor column name and the number of values of the random factor (if crossed) or the n per group (if nested); can be a vector of n per group if nested
<code>.nested_in</code>	the column(s) to nest in (if NULL, the factor is crossed with all columns)

Value

a data frame

Examples

```
# start a data frame
data1 <- add_random(school = 3)
# nest classes in schools (2 classes per school)
data2 <- add_random(data1, class = 2, .nested_in = "school")
# nest pupils in each class (different n per class)
data3 <- add_random(data2, pupil = c(20, 24, 23, 21, 25, 24), .nested_in = "class")
# cross each pupil with 10 questions
data4 <- add_random(data3, question = 10)

# compare nesting in 2 different factors
data <- add_random(A = 2, B = 2)
add_random(data, C = 2, .nested_in = "A")
add_random(data, C = 2, .nested_in = "B")
```

```
# specify item names
add_random(school = c("Hyndland Primary", "Hyndland Secondary")) %>%
  add_random(class = list(paste0("P", 1:7),
                          paste0("S", 1:6)),
             .nested_in = "school")
```

add_ranef	<i>Add random effects to a data frame</i>
-----------	---

Description

Add random effects to a data frame

Usage

```
add_ranef(.data, .by = NULL, ..., .cors = 0, .empirical = FALSE)
```

Arguments

.data	the data frame
.by	the grouping column (groups by row if NULL)
...	the name and standard deviation of each random effect
.cors	the correlations among multiple random effects, to be passed to rnorm_multi as r
.empirical	logical. To be passed to rnorm_multi as empirical

Value

data frame with new random effects columns

Examples

```
add_random(rater = 2, stimulus = 2, time = 2) %>%
  add_ranef("rater", u0r = 1.5) %>%
  add_ranef("stimulus", u0s = 2.2, u1s = 0.75, .cors = 0.5) %>%
  add_ranef(c("rater", "stimulus"), u0sr = 1.2)
```

add_recode	<i>Recode a categorical column</i>
------------	------------------------------------

Description

Recode a categorical column

Usage

```
add_recode(.data, .col, .newcol = paste0(col, ".c"), ...)
```

Arguments

.data	the data frame
.col	the column to recode
.newcol	the name of the recoded column (defaults to col.c)
...	coding for categorical column

Value

data frame with new fixed effects columns

Examples

```
add_random(subj = 4, item = 4) %>%  
  add_between("subj", cond = c("cntl", "test")) %>%  
  add_recode("cond", "cond.t", cntl = 0, test = 1)
```

add_within	<i>Add within factors</i>
------------	---------------------------

Description

Add within factors

Usage

```
add_within(.data, .by = NULL, ...)
```

Arguments

.data	the data frame
.by	the grouping column (groups by row if NULL)
...	the names and levels of the new factors

Value

data frame

Examples

```
add_random(subj = 2, item = 2) %>%
  add_within("subj", time = c("pre", "post"))
```

average_r2tau_0	<i>Average r to Random Intercept SD</i>
-----------------	---

Description

Average r to Random Intercept SD

Usage

```
average_r2tau_0(average_r, sigma)
```

Arguments

- average_r The average inter-item correlation
- sigma Total error variance

Value

The standard deviation of the random intercept

beta2norm	<i>Convert beta to normal</i>
-----------	-------------------------------

Description

Convert beta to normal

Usage

```
beta2norm(x, mu = 0, sd = 1, shape1 = NULL, shape2 = NULL, ...)
```

Arguments

- x the gamma distributed vector
- mu the mean of the normal distribution to convert to
- sd the SD of the normal distribution to convert to
- shape1, shape2 non-negative parameters of the beta distribution
- ... further arguments to pass to pbeta (e.g., ncp)

Value

a vector with a normal distribution

Examples

```
x <- rbeta(10000, 2, 3)
y <- beta2norm(x)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

binom2norm

Convert binomial to normal

Description

Convert a binomial distribution to a normal (gaussian) distribution with specified mu and sd

Usage

```
binom2norm(x, mu = 0, sd = 1, size = NULL, prob = NULL)
```

Arguments

x	the binomially distributed vector
mu	the mean of the normal distribution to return
sd	the SD of the normal distribution to return
size	number of trials (set to max value of x if not specified)
prob	the probability of success on each trial (set to mean probability if not specified)

Value

a vector with a gaussian distribution

Examples

```
x <- rbinom(10000, 20, 0.75)
y <- binom2norm(x, 0, 1, 20, 0.75)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

check_design	<i>Validates the specified design</i>
--------------	---------------------------------------

Description

Specify any number of within- and between-subject factors with any number of levels.

Usage

```
check_design(
  within = list(),
  between = list(),
  n = 100,
  mu = 0,
  sd = 1,
  r = 0,
  dv = list(y = "value"),
  id = list(id = "id"),
  vardesc = list(),
  plot = faux_options("plot"),
  design = NULL,
  fix_names = FALSE,
  sep = faux_options("sep")
)
```

Arguments

within	a list of the within-subject factors
between	a list of the between-subject factors
n	the number of samples required
mu	a vector giving the means of the variables
sd	the standard deviations of the variables
r	the correlations among the variables (can be a single number, full correlation matrix as a matrix or vector, or a vector of the upper right triangle of the correlation matrix)
dv	the name of the DV column list(y = "value")
id	the name of the ID column list(id = "id")
vardesc	a list of variable descriptions having the names of the within- and between-subject factors
plot	whether to show a plot of the design
design	a design list including within, between, n, mu, sd, r, dv, id
fix_names	deprecated
sep	separator for factor levels

Details

Specify `n` for each between-subject cell; `mu` and `sd` for each cell, and `r` for the within-subject cells for each between-subject cell.

This function returns a validated design list for use in `sim_data` to simulate a data table with this design, or to archive your design.

See `vignette("sim_design", package = "faux")` for details.

Value

list

Examples

```
within <- list(time = c("day", "night"))
between <- list(pet = c("dog", "cat"))
mu <- list(dog = 10, cat = 5)
vardesc <- list(time = "Time of Day", pet = "Type of Pet")
check_design(within, between, mu = mu, vardesc = vardesc)

between <- list(language = c("dutch", "thai"),
  pet = c("dog", "cat"))
mu <- list(dutch_dog = 12, dutch_cat = 7, thai_dog = 8, thai_cat = 3)
check_design(within, between, mu = mu)
```

check_mixed_design	<i>Get random intercepts for subjects and items</i>
--------------------	---

Description

Get error terms from an existing data table.

Usage

```
check_mixed_design(data, dv = 1, sub_id = 2, item_id = 3, formula = NULL)
```

Arguments

<code>data</code>	the existing tbl
<code>dv</code>	the column name or index containing the DV
<code>sub_id</code>	the column name or index for the subject IDs
<code>item_id</code>	the column name or index for the item IDs
<code>formula</code>	the formula to run in lmer (defaults to null model $dv \sim 1 + (1 sub_id) + (1 item_id)$)

Value

a list of parameters

Examples

```
des <- check_mixed_design(fr4, "rating", "rater_id", "face_id")
str(des[1:4])
```

codebook

Create PsychDS Codebook from Data

Description

See `vignette("codebook", package = "faux")` for details.

Usage

```
codebook(
  data,
  name = NULL,
  vardesc = list(),
  ...,
  schemaVersion = "Psych-DS 0.1.0",
  return = c("json", "list", "data"),
  interactive = FALSE
)
```

Arguments

<code>data</code>	The data frame to generate a codebook for
<code>name</code>	The name of this dataset (if NULL, will be the same as ‘data’, limited to 64 characters)
<code>vardesc</code>	Optional variable properties in the format of a named list of vectors (can be named or unnamed and in the same order as the data) from the options "description", "privacy", "dataType", "identifier", "minValue", "maxValue", "levels", "levelsOrdered", "na", "naValue", "alternateName", "privacy", "unitCode", "unitText"
<code>...</code>	Further dataset properties (e.g., description, license, author, citation, funder, url, identifier, keywords, privacyPolicy)
<code>schemaVersion</code>	defaults to "Psych-DS 0.1.0"
<code>return</code>	Whether the output should be in JSON format (json), a list (list) or the reformat- ted data with the codebook as an attribute (data)
<code>interactive</code>	Whether the function should prompt the user to describe columns and factor levels

Value

a list or json-formatted codebook, or reformatted data with the codebook as an attribute

Examples

```

vardesc = list(
  description = c("Length of the sepal",
                 "Width of the sepal",
                 "Length of the petal",
                 "Width of the petal",
                 "The flower species"),
  type = c("float", "float", "float", "float", "string")
)
codebook(iris, vardesc = vardesc)

```

contr_code_anova	<i>Anova code a factor</i>
------------------	----------------------------

Description

Anova coding (also called deviation or simple coding) sets the grand mean as the intercept. Each contrast compares one level with the reference level (base).

Usage

```
contr_code_anova(fct, levels = NULL, base = 1, colnames = NULL)
```

Arguments

fct	the factor to contrast code (or a vector)
levels	the levels of the factor in order
base	the index of the level to use as baseline
colnames	optional list of column names for the added contrasts

Value

the factor with contrasts set

Examples

```

df <- sim_design(between = list(pet = c("cat", "dog")),
                 mu = c(10, 20), plot = FALSE)
df$pet <- contr_code_anova(df$pet)
lm(y ~ pet, df) %>% broom::tidy()

df <- sim_design(between = list(pet = c("cat", "dog", "ferret")),
                 mu = c(2, 4, 9), empirical = TRUE, plot = FALSE)

df$pet <- contr_code_anova(df$pet, base = 1)
lm(y ~ pet, df) %>% broom::tidy()

```

```
df$pet <- contr_code_anova(df$pet, base = 2)
lm(y ~ pet, df) %>% broom::tidy()

df$pet <- contr_code_anova(df$pet, base = "ferret")
lm(y ~ pet, df) %>% broom::tidy()
```

contr_code_difference *Difference code a factor*

Description

Difference coding sets the grand mean as the intercept. Each contrast compares one level with the previous level.

Usage

```
contr_code_difference(fct, levels = NULL, colnames = NULL)
```

Arguments

fct	the factor to contrast code (or a vector)
levels	the levels of the factor in order
colnames	optional list of column names for the added contrasts

Value

the factor with contrasts set

Examples

```
df <- sim_design(between = list(pet = c("cat", "dog", "ferret")),
  mu = c(2, 4, 9), empirical = TRUE, plot = FALSE)

df$pet <- contr_code_difference(df$pet)
lm(y ~ pet, df) %>% broom::tidy()
```

contr_code_helmert	<i>Helmert code a factor</i>
--------------------	------------------------------

Description

Helmert coding sets the grand mean as the intercept. Each contrast compares one level with the mean of previous levels.

Usage

```
contr_code_helmert(fct, levels = NULL, colnames = NULL)
```

Arguments

fct	the factor to contrast code (or a vector)
levels	the levels of the factor in order
colnames	optional list of column names for the added contrasts

Value

the factor with contrasts set

Examples

```
df <- sim_design(between = list(pet = c("cat", "dog")),
                 mu = c(10, 20), plot = FALSE)
df$pet <- contr_code_helmert(df$pet)
lm(y ~ pet, df) %>% broom::tidy()

df <- sim_design(between = list(pet = c("cat", "dog", "ferret")),
                 mu = c(2, 4, 9), empirical = TRUE, plot = FALSE)

df$pet <- contr_code_helmert(df$pet)
lm(y ~ pet, df) %>% broom::tidy()

# reorder the levels to change the comparisons
df$pet <- contr_code_helmert(df$pet, levels = c("dog", "cat", "ferret"))
lm(y ~ pet, df) %>% broom::tidy()

df$pet <- contr_code_helmert(df$pet, levels = c("ferret", "dog", "cat"))
lm(y ~ pet, df) %>% broom::tidy()
```

contr_code_poly	<i>Polynomial code a factor</i>
-----------------	---------------------------------

Description

Polynomial coding sets the grand mean as the intercept. Each contrast tests a trend (linear, quadratic, cubic, etc.). This is only suitable for ordered factors.

Usage

```
contr_code_poly(fct, levels = NULL, colnames = NULL)
```

Arguments

fct	the factor to contrast code (or a vector)
levels	the levels of the factor in order
colnames	optional list of column names for the added contrasts

Value

the factor with contrasts set

Examples

```
df <- sim_design(within = list(time = 1:6),
                 mu = 1:6 + (1:6-3.5)^2,
                 long = TRUE, plot = FALSE)

df$time <- contr_code_poly(df$time)
lm(y ~ time, df) %>% broom::tidy()
```

contr_code_sum	<i>Sum code a factor</i>
----------------	--------------------------

Description

Sum coding sets the grand mean as the intercept. Each contrast compares one level with the grand mean.

Usage

```
contr_code_sum(fct, levels = NULL, omit = length(levels), colnames = NULL)
```


Arguments

fct	the factor to contrast code (or a vector)
levels	the levels of the factor in order
omit	the level to omit (defaults to the last level)
colnames	optional list of column names for the added contrasts

Value

the factor with contrasts set

Examples

```
df <- sim_design(between = list(pet = c("cat", "dog", "bird", "ferret")),
  mu = c(2, 4, 9, 13), empirical = TRUE, plot = FALSE)

df$pet <- contr_code_sum(df$pet)
lm(y ~ pet, df) %>% broom::tidy()

df$pet <- contr_code_sum(df$pet, omit = "cat")
lm(y ~ pet, df) %>% broom::tidy()

df$pet <- contr_code_sum(df$pet, omit = 1)
lm(y ~ pet, df) %>% broom::tidy()
```

contr_code_treatment *Treatment code a factor*

Description

Treatment coding sets the mean of the reference level (base) as the intercept. Each contrast compares one level with the reference level.

Usage

```
contr_code_treatment(fct, levels = NULL, base = 1, colnames = NULL)
```

Arguments

fct	the factor to contrast code (or a vector)
levels	the levels of the factor in order
base	the index of the level to use as baseline
colnames	optional list of column names for the added contrasts

Value

the factor with contrasts set

Examples

```
df <- sim_design(between = list(pet = c("cat", "dog")),
                 mu = c(10, 20), plot = FALSE)
df$pet <- contr_code_treatment(df$pet)
lm(y ~ pet, df) %>% broom::tidy()

df <- sim_design(between = list(pet = c("cat", "dog", "ferret")),
                 mu = c(2, 4, 9), empirical = TRUE, plot = FALSE)

df$pet <- contr_code_treatment(df$pet)
lm(y ~ pet, df) %>% broom::tidy()

df$pet <- contr_code_treatment(df$pet, base = 2)
lm(y ~ pet, df) %>% broom::tidy()

df$pet <- contr_code_treatment(df$pet, base = "ferret")
lm(y ~ pet, df) %>% broom::tidy()
```

convert_r

Convert r for NORTA

Description

Given a target r-value, returns the correlation you need to induce in a bivariate normal distribution to have the target correlation after converting distributions.

Usage

```
convert_r(
  target_r = 0,
  dist1 = "norm",
  dist2 = "norm",
  params1 = list(),
  params2 = list(),
  tol = 0.01
)
```

Arguments

target_r	The target correlation
dist1	The target distribution function for variable 1 (e.g., norm, binom, gamma, trunc-norm)
dist2	The target distribution function for variable 2
params1	Arguments to pass to the functions for distribution 1
params2	Arguments to pass to the functions for distribution 2
tol	Tolerance for optimise function

Details

See [Distributions](#) for distributions and their various arguments to specify in params1 and params2.

Value

r-value to induce in the bivariate normal variables

Examples

```
convert_r(target_r = 0.5,
  dist1 = "norm",
  dist2 = "binom",
  params1 = list(mean = 100, sd = 10),
  params2 = list(size = 1, prob = 0.5))
```

cormat	<i>Make a correlation matrix</i>
--------	----------------------------------

Description

cormat makes a correlation matrix from a single number, vars*vars matrix, vars*vars vector, or a vars*(vars-1)/2 vector.

Usage

```
cormat(cors = 0, vars = 3)
```

Arguments

- cors the correlations among the variables (can be a single number, vars*vars matrix, vars*vars vector, or a vars*(vars-1)/2 vector)
- vars the number of variables in the matrix

Value

matrix

Examples

```
cormat(.5, 3)
cormat(c( 1, .2, .3, .4,
  .2, 1, .5, .6,
  .3, .5, 1, .7,
  .4, .6, .7, 1), 4)
cormat(c(.2, .3, .4, .5, .6, .7), 4)
```

`cormat_from_triangle` *Make Correlation Matrix from Triangle*

Description

`cormat_from_triangle` makes a correlation matrix from a vector of the upper right triangle

Usage

```
cormat_from_triangle(cors)
```

Arguments

`cors` the correlations among the variables as a $\text{vars} \times (\text{vars}-1)/2$ vector

Value

matrix

Examples

```
cormat_from_triangle(c(.2, .3, .4,
                        .5, .6,
                        .7))
```

`distfuncs` *Get distribution functions*

Description

Get distribution functions

Usage

```
distfuncs(dist = "norm")
```

Arguments

`dist` The target distribution function (e.g., `norm`, `binom`, `gamma`, `truncnorm`, `likert`). If the distribution isn't defined in the packages `stats`, `truncnorm`, or `faux`, use the format `"package::dist"`.

Value

a list with the `r` and `q` functions

Examples

```

qfunc <- distfuncs("norm")$q # returns qnorm
p <- seq(0.1, 0.9, .1)
qfunc(p) == qnorm(p)

rfunc <- distfuncs("norm")$r # returns rnorm
rfunc(n = 10, mean = 100, sd = 10)

```

dlikert

Likert density function

Description

Likert density function

Usage

```
dlikert(x, prob, labels = names(prob))
```

Arguments

x	the likert distributed vector
prob	a vector of probabilities or counts; if named, the output is a factor
labels	a vector of values, defaults to names(prob) or 1:length(prob), if numeric, the output is numeric

Value

a vector of the densities

Examples

```

x <- 1:5
prob <- c(.1, .2, .4, .2, .1)
dlikert(x, prob)

x <- c("A", "C", "B", "B")
prob <- c(A = 10, B = 20, C = 30)
dlikert(x, prob)

# specify labels if prob not named and not 1:length(prob)
labels <- -2:2
x <- sample(labels, 10, replace = TRUE)
prob <- rep(1, length(labels)) # uniform probability
dlikert(x, prob, labels)

```

faceratings	<i>Attractiveness ratings of faces</i>
-------------	--

Description

A dataset containing attractiveness ratings (on a 1-7 scale from "much less attractiveness than average" to "much more attractive than average") for the neutral front faces from 2513 people (ages 17-90)

Usage

```
faceratings
```

Format

A data frame with 256326 rows and 9 variables:

rater_id rater's ID

rater_sex rater's sex (female, male, intersex, NA)

rater_age rater's age (17-90 years)

rater_sexpref rater's preferred sex for romantic relationships (either, men, neither, women, NA)

face_id face's ID

face_sex face's sex (female, male)

face_age face's age (in years)

face_eth face's ethnic group

rating attractiveness rating on a scale from 1 (much less attractive than average) to 7 (much more attractive than average)

Source

https://figshare.com/articles/dataset/Face_Research_Lab_London_Set/5047666

faux_options	<i>Set/get global faux options</i>
--------------	------------------------------------

Description

Global faux options are used, for example, to set the default separator for cell names.

Usage

```
faux_options(...)
```

Arguments

... One of four: (1) nothing, then returns all options as a list; (2) a name of an option element, then returns its value; (3) a name-value pair which sets the corresponding option to the new value (and returns nothing), (4) a list with option-value pairs which sets all the corresponding arguments.

Value

a list of options, values of an option, or nothing

Examples

```
faux_options() # see all options

faux_options("sep") # see value of faux.sep

# changes cell separator (e.g., A1.B2)
faux_options(sep = ".")

# changes cell separator back to default (e.g., A1_B2)
faux_options(sep = "_")
```

fh_bounds	<i>Get Fréchet-Hoeffding bounds</i>
-----------	-------------------------------------

Description

Fréchet-Hoeffding bounds are the limits to a correlation between different distributions.

Usage

```
fh_bounds(dist1 = "norm", dist2 = "norm", params1 = list(), params2 = list())
```

Arguments

dist1	The target distribution function for variable 1 (e.g., norm, binom, gamma, trunc-norm)
dist2	The target distribution function for variable 2
params1	Arguments to pass to the random generation function (e.g., rnorm) for distribution 1
params2	Arguments to pass to the random generation function (e.g., rnorm) for distribution 2

Value

a list of the min and max possible values

Examples

```
fh_bounds(dist1 = "pois",
          dist2 = "unif",
          params1 = list(lambda = 3),
          params2 = list(min = 0, max = 100))
```

fix_name_labels	<i>Fix name labels</i>
-----------------	------------------------

Description

Fixes if a factor list does not have named levels or has special characters in the names

Usage

```
fix_name_labels(x, pattern = NA, replacement = ".")
```

Arguments

- x the vector or list to fix
- pattern regex pattern to replace; defaults to non-word characters and the value of faux_options("sep") (default = _)
- replacement the character to replace; defaults to . (or _ if faux_options("sep") == ".")

Value

a named list with fixed names

Examples

```
source <- list("full.stop", " space ", "under_score", "plus+", "dash-", "tab\t", "line\nbreak")
fix_name_labels(source)
```

fr4	<i>Attractiveness rating subset</i>
-----	-------------------------------------

Description

The faceratings dataset cut down for demos to the first 4 raters of each sex and sexpref and the first 4 faces of each sex and ethnicity with non-NA ages

Usage

```
fr4
```


Format

A data frame with 768 rows and 9 variables:

rater_id rater's ID

rater_sex rater's sex (female, male)

rater_age rater's age (17.4-54.3 years)

rater_sexpref rater's preferred sex for romantic relationships (either, men, women)

face_id face's ID

face_sex face's sex (female, male)

face_age face's age (19-47 years)

face_eth face's ethnic group (black, east_asian, west_asian, white)

rating attractiveness rating on a scale from 1 (much less attractive than average) to 7 (much more attractive than average)

Source

https://figshare.com/articles/dataset/Face_Research_Lab_London_Set/5047666

gamma2norm	<i>Convert gamma to normal</i>
------------	--------------------------------

Description

Convert gamma to normal

Usage

```
gamma2norm(x, mu = 0, sd = 1, shape = NULL, rate = 1, scale = 1/rate)
```

Arguments

x	the gamma distributed vector
mu	the mean of the normal distribution to convert to
sd	the SD of the normal distribution to convert to
shape	gamma distribution parameter (must be positive)
rate	an alternative way to specify the scale
scale	gamma distribution parameter (must be positive)

Value

a vector with a normal distribution

Examples

```
x <- rgamma(10000, 2)
y <- gamma2norm(x)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

getcols	<i>Get data columns</i>
---------	-------------------------

Description

Get columns from a data table by specifying the index, column name as a string, or unquoted column name. Returns the column names or indices.

Usage

```
getcols(data, ..., as_index = FALSE)
```

Arguments

- data the existing tbl
- ... Columns to get
- as_index return the column indices (defaults to name)

Value

vector of column names or indices

Examples

```
getcols(mtcars, 1, cyl, "dis", 5:7)
```

get_coefs	<i>Get Coefficients from Data</i>
-----------	-----------------------------------

Description

You need model coefficients to simulate multilevel data, and can get them from data simulated from parameters using sim_design() or rmulti().

Usage

```
get_coefs(data, formula = NULL, fun = c("lm", "glm", "lmer", "glmer"), ...)
```

Arguments

data	A dataset in long format
formula	A formula (can be extracted from datasets created by sim_design)
fun	the model function (one of "lm", "glm", "lmer", or "glmer")
...	Further arguments to the model function

Value

a list of the model coefficients

Examples

```
# simulate some data
data <- sim_design(within = 2, between = 2,
                  mu = c(1, 0, 1, 1),
                  long = TRUE, empirical = TRUE)

# get coefs for the full factorial model
get_coefs(data)

# a reduced model
get_coefs(data, y ~ B1 + W1)

# specify a different model function
data$y <- norm2binom(data$y)
get_coefs(data, fun = "glm", family = binomial)
```

get_contrast_vals	<i>Get contrast values</i>
-------------------	----------------------------

Description

Get a data frame of contrast values from a factor vector

Usage

```
get_contrast_vals(v)
```

Arguments

v	a factor vector
---	-----------------

Value

a data frame

Examples

```
dat <- sim_design(  
  between = list(group = c("A", "B")),  
  n = 5, plot = FALSE)  
  
get_contrast_vals(dat$group)
```

get_design	<i>Get design</i>
------------	-------------------

Description

Get the design specification from a data table created in faux. This can be used to create more simulated data with the same design.

Usage

```
get_design(data)
```

Arguments

data The data table to check

Value

list with class design

Examples

```
data <- sim_design(2, 2, plot = FALSE)  
design <- get_design(data)  
data2 <- sim_design(design, plot = FALSE)
```

get_design_long	<i>Get design from long data</i>
-----------------	----------------------------------

Description

Makes a best guess at the design of a long-format data frame.

Usage

```
get_design_long(
  data,
  dv = c(y = "score"),
  id = c(id = "id"),
  plot = faux_options("plot")
)
```

Arguments

data	the data frame (in long format)
dv	the column name that identifies the DV
id	the column name(s) that identify a unit of analysis
plot	whether to show a plot of the design

Details

Finds all columns that contain a single value per unit of analysis (between factors), all columns that contain the same values per unit of analysis (within factors), and all columns that differ over units of analysis (dv, continuous factors)

Value

a design list

get_params	<i>Get parameters from a data table</i>
------------	---

Description

Generates a table of the correlations and means of numeric columns in a data frame. If data was generated by `sim_design` and has a "design" attribute, between, within, dv and id are retrieved from that, unless overridden (use `between = 0` to

Usage

```
get_params(
  data,
  between = NULL,
  within = NULL,
  dv = NULL,
  id = NULL,
  digits = 2
)

check_sim_stats(
```

```

    data,
    between = NULL,
    within = NULL,
    dv = NULL,
    id = NULL,
    digits = 2
  )

```

Arguments

data	the existing tbl
between	a vector of column names for between-subject factors
within	a vector of column names for within-subject factors (if data is long)
dv	the column name(s) of the dv, if NULL all numeric columns will be selected
id	the column name(s) of the subject ID, excluded from the table even if numeric
digits	how many digits to round to (default = 2)

Value

a tbl of correlations, means and sds

Examples

```
get_params(iris, "Species")
```

interactive_design	<i>Set design interactively</i>
--------------------	---------------------------------

Description

Set design interactively

Usage

```
interactive_design(output = c("faux"), plot = faux_options("plot"))
```

Arguments

output	what type of design to output (faux)
plot	whether to show a plot of the design

Value

list

Examples

```
if(interactive()){ des <- interactive_design() }
```

is_pos_def	<i>Check a Matrix is Positive Definite</i>
------------	--

Description

is_pos_def makes a correlation matrix from a vector

Usage

```
is_pos_def(cor_mat, tol = 1e-08)
```

Arguments

cor_mat	a correlation matrix
tol	the tolerance for comparing eigenvalues to 0

Value

logical value

Examples

```
is_pos_def(matrix(c(1, .5, .5, 1), 2)) # returns TRUE
is_pos_def(matrix(c(1, .9, .9,
                    .9, 1, -.2,
                    .9, -.2, 1), 3)) # returns FALSE
```

json_design	<i>Convert design to JSON</i>
-------------	-------------------------------

Description

Convert a design list to JSON notation for archiving (e.g. in scienceverse)

Usage

```
json_design(design, filename = NULL, digits = 8, pretty = FALSE, ...)
```

Arguments

design	a design list including within, between, n, mu, sd, r, dv, id
filename	option name of file to save the json to
digits	number of digits to save
pretty	whether to print condensed or readable
...	other options to send to jsonlite::toJSON

Value

a JSON string

Examples

```
des <- check_design(2,2)
json_design(des)
json_design(des, pretty = TRUE)
```

long2wide

Convert data from long to wide format

Description

Convert data from long to wide format

Usage

```
long2wide(
  data,
  within = c(),
  between = c(),
  dv = "y",
  id = "id",
  sep = faux_options("sep")
)
```

Arguments

data	the tbl in long format
within	the names of the within column(s)
between	the names of between column(s) (optional)
dv	the name of the DV (value) column
id	the names of the column(s) for grouping observations
sep	separator for factor levels

Value

a tbl in wide format

Examples

```
df_long <- sim_design(2, 2, long = TRUE)
long2wide(df_long, "A", "B")
```

make_id	<i>Make ID</i>
---------	----------------

Description

Make IDs with fixed length and a prefix (e.g., S001, S002, ..., S100).

Usage

```
make_id(n = 100, prefix = "S", digits = 0, suffix = "")
```

Arguments

n	the number of IDs to generate (or a vector of numbers)
prefix	the prefix to the number (default "S")
digits	the number of digits to use for the numeric part. Only used if this is larger than the largest number of digits in n.
suffix	the suffix to the number (default "")

Value

a vector of IDs

Examples

```
make_id(20, "SUBJECT_")  
make_id(10:30, digits = 3)
```

messy	<i>Simulate missing data</i>
-------	------------------------------

Description

Insert NA or another replacement value for some proportion of specified columns to simulate missing data.

Usage

```
messy(data, prop = 0, ..., replace = NA)
```

Arguments

data	the tbl
prop	the proportion of data to mess up
...	the columns to mess up (as a vector of column names or numbers)
replace	the replacement value (defaults to NA)

Value

the messed up table

Examples

```
messy(iris, 0.1, "Species", replace = "NO SPECIES")
messy(iris, 0.5, 1:4)
```

nbinom2norm	<i>Convert negative binomial to normal</i>
-------------	--

Description

Convert a negative binomial distribution to a normal (gaussian) distribution with specified mu and sd

Usage

```
nbinom2norm(x, mu = 0, sd = 1, size = NULL, prob = NULL)
```

Arguments

- x the negative binomially distributed vector
- mu the mean of the normal distribution to return
- sd the SD of the normal distribution to return
- size number of trials (set to max value of x if not specified)
- prob the probability of success on each trial (set to mean probability if not specified)

Value

a vector with a gaussian distribution

Examples

```
x <- rnbinom(10000, 20, 0.75)
y <- nbinom2norm(x, 0, 1, 20, 0.75)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

nested_list	<i>Output a nested list in RMarkdown list format</i>
-------------	--

Description

Output a nested list in RMarkdown list format

Usage

```
nested_list(x, pre = "", quote = "")
```

Arguments

x	The list
pre	Text to prefix to each line (e.g., if you want all lines indented 4 spaces to start, use " ")
quote	Text to quote values with (e.g., use "\"" to make sure values are not parsed as markdown)

Value

A character string

Examples

```
x <- list(
  a = list(a1 = "Named", a2 = "List"),
  b = list("Unnamed", "List"),
  c = c(c1 = "Named", c2 = "Vector"),
  d = c("Unnamed", "Vector"),
  e = list(e1 = list("A", "B", "C"),
    e2 = list(a = "A", b = "B"),
    e3 = c("A", "B", "C"),
    e4 = 100),
  f = "single item vector",
  g = list()
)
nested_list(x)
```

norm2beta	<i>Convert normal to beta</i>
-----------	-------------------------------

Description

Convert normal to beta

Usage

```
norm2beta(x, shape1, shape2, mu = mean(x), sd = stats::sd(x), ...)
```

Arguments

x	the normally distributed vector
shape1, shape2	non-negative parameters of the distribution to return
mu	the mean of x (calculated from x if not given)
sd	the SD of x (calculated from x if not given)
...	further arguments to pass to qbeta (e.g., ncp)

Value

a vector with a beta distribution

Examples

```
x <- rnorm(10000)
y <- norm2beta(x, 1, 3)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2binom	<i>Convert normal to binomial</i>
------------	-----------------------------------

Description

Convert normal to binomial

Usage

```
norm2binom(x, size = 1, prob = 0.5, mu = mean(x), sd = stats::sd(x))
```

Arguments

x	the normally distributed vector
size	number of trials (0 or more)
prob	the probability of success on each trial (0 to 1)
mu	the mean of x (calculated from x if not given)
sd	the SD of x (calculated from x if not given)

Value

a vector with a binomial distribution

Examples

```
x <- rnorm(10000)
y <- norm2binom(x)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2gamma

Convert normal to gamma

Description

Convert normal to gamma

Usage

```
norm2gamma(x, shape, rate = 1, scale = 1/rate, mu = mean(x), sd = stats::sd(x))
```

Arguments

x	the normally distributed vector
shape	gamma distribution parameter (must be positive)
rate	an alternative way to specify the scale
scale	gamma distribution parameter (must be positive)
mu	the mean of x (calculated from x if not given)
sd	the SD of x (calculated from x if not given)

Value

a vector with a gamma distribution

Examples

```
x <- rnorm(10000)
y <- norm2gamma(x, shape = 2)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2likert

*Convert normal to likert***Description**

Convert normal to likert

Usage

```
norm2likert(x, prob, labels = names(prob), mu = mean(x), sd = stats::sd(x))
```

Arguments

x	the normally distributed vector
prob	a vector of probabilities or counts; if named, the output is a factor
labels	a vector of values, defaults to names(prob) or 1:length(prob), if numeric, the output is numeric
mu	the mean of x (calculated from x if not given)
sd	the SD of x (calculated from x if not given)

Value

a vector with the specified distribution

Examples

```
x <- rnorm(10000)
y <- norm2likert(x, c(.1, .2, .35, .2, .1, .05))
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")

y <- norm2likert(x, c(40, 30, 20, 10))
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")

y <- norm2likert(x, c(lower = .5, upper = .5))
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2nbinom	<i>Convert normal to negative binomial</i>
-------------	--

Description

See the help for ‘qnbinom()’ for further info about prob versus mu parameter specification. Thanks for the suggested code, David Hugh-Jones!

Usage

```
norm2nbinom(
  x,
  size,
  prob,
  mu,
  lower.tail = TRUE,
  log.p = FALSE,
  x_mu = mean(x),
  x_sd = stats::sd(x)
)
```

Arguments

x	the normally distributed vector
size	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). (size > 0)
prob	the probability of success on each trial (0 to 1)
mu	alternative parametrization via mean (only specify one of prob or mu)
lower.tail	logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X > x]$
log.p	logical; if TRUE, probabilities p are given as $\log(p)$
x_mu	the mean of x (calculated from x if not given)
x_sd	the SD of x (calculated from x if not given)

Value

a vector with a negative binomial distribution

Examples

```
x <- rnorm(10000)
y <- norm2nbinom(x, 1, prob = 0.5)
z <- norm2nbinom(x, 1, mu = 1)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2norm	<i>Convert normal to normal</i>
-----------	---------------------------------

Description

Convert a normal distribution to a normal (gaussian) distribution with specified mu and sd

Usage

```
norm2norm(x, mu = 0, sd = 1, x_mu = mean(x), x_sd = stats::sd(x))
```

Arguments

x	the uniformly distributed vector
mu	the mean of the normal distribution to return
sd	the SD of the normal distribution to return
x_mu	the mean of x (calculated from x if not given)
x_sd	the SD of x (calculated from x if not given)

Value

a vector with a gaussian distribution

Examples

```
x <- rnorm(10000)
y <- norm2norm(x, 100, 10)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2pois	<i>Convert normal to poisson</i>
-----------	----------------------------------

Description

Convert normal to poisson

Usage

```
norm2pois(x, lambda, mu = mean(x), sd = stats::sd(x))
```


Arguments

x	the normally distributed vector
lambda	the mean of the distribution to return
mu	the mean of x (calculated from x if not given)
sd	the SD of x (calculated from x if not given)

Value

a vector with a poisson distribution

Examples

```
x <- rnorm(10000)
y <- norm2pois(x, 2)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2trunc

Convert normal to truncated normal

Description

Convert a normal (gaussian) distribution to a truncated normal distribution with specified minimum and maximum

Usage

```
norm2trunc(
  x,
  min = -Inf,
  max = Inf,
  mu = mean(x),
  sd = stats::sd(x),
  x_mu = mean(x),
  x_sd = stats::sd(x)
)
```

Arguments

x	the normally distributed vector
min	the minimum of the truncated distribution to return
max	the maximum of the truncated distribution to return
mu	the mean of the distribution to return (calculated from x if not given)
sd	the SD of the distribution to return (calculated from x if not given)
x_mu	the mean of x (calculated from x if not given)
x_sd	the SD of x (calculated from x if not given)

Value

a vector with a uniform distribution

Examples

```
x <- rnorm(10000)
y <- norm2trunc(x, 1, 7, 3.5, 2)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

norm2unif

Convert normal to uniform

Description

Convert a normal (gaussian) distribution to a uniform distribution with specified minimum and maximum

Usage

```
norm2unif(x, min = 0, max = 1, mu = mean(x), sd = stats::sd(x))
```

Arguments

x	the normally distributed vector
min	the minimum of the uniform distribution to return
max	the maximum of the uniform distribution to return
mu	the mean of x (calculated from x if not given)
sd	the SD of x (calculated from x if not given)

Value

a vector with a uniform distribution

Examples

```
x <- rnorm(10000)
y <- norm2unif(x)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

plikert	<i>Likert distribution function</i>
---------	-------------------------------------

Description

Likert distribution function

Usage

```
plikert(q, prob, labels = names(prob))
```

Arguments

q	the vector of quantiles
prob	a vector of probabilities or counts; if named, the output is a factor
labels	a vector of values, defaults to names(prob) or 1:length(prob), if numeric, the output is numeric

Value

a vector of the densities

Examples

```
q <- 1:5
prob <- c(.1, .2, .4, .2, .1)
plikert(q, prob)

q <- c("A", "C", "B", "B")
prob <- c(A = 10, B = 20, C = 30)
plikert(q, prob)

# specify labels if prob not named and not 1:length(prob)
labels <- -2:2
q <- labels
prob <- rep(1, length(labels)) # uniform probability
plikert(q, prob, labels)
```

plot_design	<i>Plot design</i>
-------------	--------------------

Description

Plots the specified within and between design. See `vignette("plots", package = "faux")` for examples and details.

Usage

```
plot_design(x, ..., geoms = NULL, palette = "Dark2", labeller = "label_value")

## S3 method for class 'design'
plot(x, ...)

## S3 method for class 'faux'
plot(x, ...)
```

Arguments

x	A list of design parameters created by <code>check_design()</code> or a data tbl (in long format)
...	A list of factor names to determine visualisation (see vignette) in the order color, x, facet row(s), facet col(s)
geoms	A list of ggplot2 geoms to display, defaults to "pointrangeSD" (mean $\pm$ 1SD) for designs and c("violin", "box") for data, options are: pointrangeSD, pointrangeSE, violin, box, jitter
palette	A brewer palette, defaults to "Dark2" (see <code>ggplot2::scale_colour_brewer</code>)
labeller	How to label the facets (see <code>ggplot2::facet_grid</code>). "label_value" is used by default.

Value

plot

Functions

- `plot(design)`: Plotting from a faux design list
- `plot(faux)`: Plotting from a faux data table

Examples

```
within <- list(time = c("day", "night"))
between <- list(pet = c("dog", "cat"))
des <- check_design(within, between, plot = FALSE)
plot_design(des)

data <- sim_design(within, between, plot = FALSE)
plot_design(data)
```

pos_def_limits	<i>Limits on Missing Value for Positive Definite Matrix</i>
----------------	---

Description

pos_def_limits returns min and max possible values for a positive definite matrix with a specified missing value

Usage

```
pos_def_limits(..., steps = 0.01, tol = 1e-08)
```

Arguments

...	the correlations among the variables as a $\text{vars} \times (\text{vars}-1)/2$ vector
steps	the tolerance for min and max values
tol	the tolerance for comparing eigenvalues to 0

Value

dataframe with min and max values

Examples

```
pos_def_limits(.8, .2, NA)
```

qlikert	<i>Likert quantile function</i>
---------	---------------------------------

Description

Likert quantile function

Usage

```
qlikert(p, prob, labels = names(prob))
```

Arguments

p	the vector of probabilities
prob	a vector of probabilities or counts; if named, the output is a factor
labels	a vector of values, defaults to names(prob) or 1:length(prob), if numeric, the output is numeric

Value

a vector of the quantiles

Examples

```
p <- seq(0, 1, .1)
prob <- c(.1, .2, .4, .2, .1)
qlikert(p, prob)

p <- seq(0, 1, .1)
prob <- c(A = 10, B = 20, C = 30)
qlikert(p, prob)

# specify labels if prob not named and not 1:length(prob)
labels <- -2:2
p <- seq(0, 1, .1)
prob <- rep(1, length(labels)) # uniform probability
qlikert(p, prob, labels)
```

readline_check	<i>Check readline input</i>
----------------	-----------------------------

Description

Check readline input

Usage

```
readline_check(
  prompt,
  type = c("numeric", "integer", "length", "grep"),
  min = -Inf,
  max = Inf,
  warning = NULL,
  default = NULL,
  ...
)
```

Arguments

prompt	the prompt for readline
type	what type of check to perform, one of c("numeric", "integer", "length", "grep")
min	the minimum value
max	the maximum value
warning	an optional custom warning message
default	the default option to return if the entry is blank, NULL allows no default, the default value will be displayed after the text as [default]
...	other arguments to pass to grep

Value

the validated result of readline

Examples

```
if(interactive()){
  readline_check("Type a number: ", "numeric")
  readline_check("Type two characters: ", "length", min = 2, max = 2)
  readline_check("Type at least 3 characters: ", "length", min = 3)
  readline_check("Type no more than 4 characters: ", "length", max = 4)
  readline_check("Type a letter and a number: ", "grep", pattern = "[a-zA-Z]\\d$")
}
```

rlikert

Random Likert distribution

Description

Random Likert distribution

Usage

```
rlikert(n, prob, labels = names(prob))
```

Arguments

n	the number of observations
prob	a vector of probabilities or counts; if named, the output is a factor
labels	a vector of values, defaults to names(prob) or 1:length(prob), if numeric, the output is numeric

Value

a vector sampled from a likert distribution with the specified parameters

Examples

```
# no names or labels returns integer vector of values 1:length(prob)
prob <- c(.1, .2, .4, .2, .1)
rlikert(10, prob)

# named prob returns factor
prob <- c(A = 10, B = 20, C = 30)
rlikert(10, prob)

# specify labels if prob not named and not 1:length(prob)
labels <- -2:2
prob <- rep(1, length(labels)) # uniform probability
rlikert(10, prob, labels)
```

rmulti	<i>Multiple correlated distributions</i>
--------	--

Description

Multiple correlated distributions

Usage

```
rmulti(
  n = 100,
  dist = c(A = "norm", B = "norm"),
  params = list(),
  r = 0,
  empirical = FALSE,
  as.matrix = FALSE
)
```

Arguments

n	the number of samples required
dist	A named vector of the distributions of each variable
params	A list of lists of the arguments to pass to each distribution function
r	the correlations among the variables (can be a single number, vars*vars matrix, vars*vars vector, or a vars*(vars-1)/2 vector)
empirical	logical. If true, params specify the sample parameters, not the population parameters
as.matrix	logical. If true, returns a matrix

Value

a tbl of vars vectors

Examples

```
dist <- c(A = "norm",
          B = "pois",
          C = "binom")
params <- list(A = list(mean = 100, sd = 10),
              B = list(lambda = 5),
              C = list(size = 10, prob = 0.5))
x <- rmulti(100, dist, params, c(0.2, 0.4, 0.6), empirical = TRUE)
get_params(x)
```

rnorm_multi*Multiple correlated normal distributions*

Description

Make normally distributed vectors with specified relationships. See `vignette("rnorm_multi", package = "faux")` for details.

Usage

```
rnorm_multi(  
  n = 100,  
  vars = NULL,  
  mu = 0,  
  sd = 1,  
  r = 0,  
  varnames = NULL,  
  empirical = FALSE,  
  as.matrix = FALSE,  
  seed = NULL  
)
```

Arguments

n	the number of samples required
vars	the number of variables to return
mu	a vector giving the means of the variables (numeric vector of length 1 or vars)
sd	the standard deviations of the variables (numeric vector of length 1 or vars)
r	the correlations among the variables (can be a single number, vars*vars matrix, vars*vars vector, or a vars*(vars-1)/2 vector)
varnames	optional names for the variables (string vector of length vars) defaults if r is a matrix with column names
empirical	logical. If true, mu, sd and r specify the empirical not population mean, sd and covariance
as.matrix	logical. If true, returns a matrix
seed	DEPRECATED use set.seed() instead before running this function

Value

a tbl of vars vectors

Examples

```
# 4 10-item vectors each correlated r = .5
rnorm_multi(10, 4, r = 0.5)

# set r with the upper right triangle
b <- rnorm_multi(100, 3, c(0, .5, 1), 1,
                  r = c(0.2, -0.5, 0.5),
                  varnames=c("A", "B", "C"))

cor(b)

# set r with a correlation matrix and column names from mu names
c <- rnorm_multi(
  n = 100,
  mu = c(A = 0, B = 0.5, C = 1),
  r = c( 1,    0.2, -0.5,
        0.2, 1,    0.5,
        -0.5, 0.5, 1)
)
cor(c)
```

rnorm_pre

*Make a normal vector correlated to existing vectors***Description**

rnorm_pre Produces a random normally distributed vector with the specified correlation to one or more existing vectors

Usage

```
rnorm_pre(x, mu = 0, sd = 1, r = 0, empirical = FALSE, threshold = 1e-12)
```

Arguments

x	the existing vector or data table of all vectors
mu	desired mean of returned vector
sd	desired SD of returned vector
r	desired correlation(s) between existing and returned vectors
empirical	logical. If true, mu, sd and r specify the empirical not population mean, sd and covariance
threshold	for checking correlation matrix

Value

vector

Examples

```
v1 <- rnorm(10)
v2 <- rnorm_pre(v1, 0, 1, 0.5)
cor(v1, v2)

x <- rnorm_multi(50, 2, .5)
x$y <- rnorm_pre(x, r = c(0.5, 0.25))
cor(x)
```

sample_from_pop	<i>Sample Parameters from Population Parameters</i>
-----------------	---

Description

Sample Parameters from Population Parameters

Usage

```
sample_from_pop(n = 100, mu = 0, sd = 1, r = 0)
```

Arguments

n	sample size
mu	population mean
sd	population SD
r	population r

Value

list of sample parameters (mu, sd, r)

Examples

```
sample_from_pop(10)
```

set_design	<i>Set design</i>
------------	-------------------

Description

Add a design specification to a data table

Usage

```
set_design(data, design)
```

Arguments

data	The data table
design	The design list

Value

A data frame with a design attribute

Examples

```
design <- check_design()
data <- data.frame(id = 1:100, y = rnorm(100)) %>%
  set_design(design)
```

sim_design	<i>Simulate data from design</i>
------------	----------------------------------

Description

Generates a data table with a specified within and between design. See `vignette("sim_design", package = "faux")` for examples and details.

Usage

```
sim_design(
  within = list(),
  between = list(),
  n = 100,
  mu = 0,
  sd = 1,
  r = 0,
  empirical = FALSE,
  long = faux_options("long"),
  dv = list(y = "value"),
```

```

    id = list(id = "id"),
    vardesc = list(),
    plot = faux_options("plot"),
    interactive = FALSE,
    design = NULL,
    rep = 1,
    nested = TRUE,
    seed = NULL,
    sep = faux_options("sep")
  )

```

Arguments

within	a list of the within-subject factors
between	a list of the between-subject factors
n	the number of samples required
mu	the means of the variables
sd	the standard deviations of the variables
r	the correlations among the variables (can be a single number, full correlation matrix as a matrix or vector, or a vector of the upper right triangle of the correlation matrix)
empirical	logical. If true, mu, sd and r specify the empirical not population mean, sd and covariance
long	Whether the returned tbl is in wide or long format (defaults to value of 'faux_options("long")')
dv	the name of the dv for long plots (defaults to y)
id	the name of the id column (defaults to id)
vardesc	a list of variable descriptions having the names of the within- and between-subject factors
plot	whether to show a plot of the design
interactive	whether to run the function interactively
design	a design list including within, between, n, mu, sd, r, dv, id, and vardesc
rep	the number of data frames to return (default 1); if greater than 1, the returned data frame is nested by rep (if nested = TRUE)
nested	Whether to nest data frames by rep if rep > 1
seed	DEPRECATED use set.seed() instead before running this function
sep	separator for factor levels

Value

a tbl

sim_df	<i>Simulate an existing dataframe</i>
--------	---------------------------------------

Description

Produces a data table with the same distributions and correlations as an existing data table Only returns numeric columns and simulates all numeric variables from a continuous normal distribution (for now).

Usage

```
sim_df(
  data,
  n = 100,
  within = c(),
  between = c(),
  id = "id",
  dv = "value",
  empirical = FALSE,
  long = faux_options("long"),
  seed = NULL,
  missing = FALSE,
  sep = faux_options("sep")
)
```

Arguments

data	the existing tbl
n	the number of samples to return per group
within	a list of the within-subject factor columns (if long format)
between	a list of the between-subject factor columns
id	the names of the column(s) for grouping observations
dv	the name of the DV (value) column
empirical	Should the returned data have these exact parameters? (versus be sampled from a population with these parameters)
long	whether to return the data table in long format
seed	DEPRECATED use set.seed() instead before running this function
missing	simulate missing data?
sep	separator for factor levels

Details

See `vignette("sim_df", package = "faux")` for details.

Value

a tbl

Examples

```
iris100 <- sim_df(iris, 100)
iris_species <- sim_df(iris, 100, between = "Species")

# set the names of within factors and (the separator character)
# if you want to return a long version
longdf <- sim_df(iris,
  between = "Species",
  within = c("type", "dim"),
  sep = ".",
  long = TRUE)

# or if you are simulating data from a table in long format
widedf <- sim_df(longdf,
  between = "Species",
  within = c("type", "dim"),
  sep = ".")
```

sim_joint_dist	<i>Simulate category joint distribution</i>
----------------	---

Description

This function is mainly used internally, such as for simulating missing data patterns, but is available in case anyone finds it useful.

Usage

```
sim_joint_dist(data, ..., n = 100, empirical = FALSE)
```

Arguments

data	the existing tbl
...	columns to calculate the joint distribution from, if none are chosen, all columns with 10 or fewer unique values will be chosen
n	the number of total observations to return
empirical	Should the returned data have the exact same distribution of conditions? (versus be sampled from a population with this distribution)

Value

data table

Examples

```
sim_joint_dist(ggplot2::diamonds, cut, color, n = 10)
```

```
sim_mixed_cc
```

Generate a cross-classified sample

Description

Makes a basic cross-classified design with random intercepts for subjects and items. See `vignette("sim_mixed", package = "faux")` for examples and details.

Usage

```
sim_mixed_cc(
  sub_n = 100,
  item_n = 20,
  grand_i = 0,
  sub_sd = 1,
  item_sd = 1,
  error_sd = 1,
  empirical = FALSE,
  seed = NULL
)
```

Arguments

<code>sub_n</code>	the number of subjects
<code>item_n</code>	the number of items
<code>grand_i</code>	the grand intercept (overall mean)
<code>sub_sd</code>	the SD of subject random intercepts (or a <code>sub_n</code> -length named vector of random intercepts for each subject)
<code>item_sd</code>	the SD of item random intercepts (or an <code>item_n</code> -length named vector of random intercepts for each item)
<code>error_sd</code>	the SD of the error term
<code>empirical</code>	Should the returned data have these exact parameters? (versus be sampled from a population with these parameters)
<code>seed</code>	DEPRECATED use <code>set.seed()</code> instead before running this function

Value

a `tbl`

Examples

```
sim_mixed_cc(10, 10)
```

sim_mixed_df	<i>Generate a mixed design from existing data</i>
--------------	---

Description

sim_mixed_df() produces a data table with the same distributions of by-subject and by-item random intercepts as an existing data table.

Usage

```
sim_mixed_df(  
  data,  
  sub_n = NULL,  
  item_n = NULL,  
  dv = "y",  
  sub_id = "sub_id",  
  item_id = "item_id"  
)
```

Arguments

data	the existing tbl
sub_n	the number of subjects to simulate (if NULL, returns data for the same subjects)
item_n	the number of items to simulate (if NULL, returns data for the same items)
dv	the column name or index containing the DV
sub_id	the column name or index for the subject IDs
item_id	the column name or index for the item IDs

Value

a tbl

Examples

```
sim_mixed_df(faceratings, 10, 10, "rating", "rater_id", "face_id")
```

std_alpha2average_r	<i>Standardized Alpha to Average R</i>
---------------------	--

Description

Standardized Alpha to Average R

Usage

```
std_alpha2average_r(std_alpha, n)
```

Arguments

std_alpha	The standarized alpha
n	The number of items

Value

The average inter-item correlation

Examples

```
std_alpha2average_r(.8, 10)
```

trunc2norm	<i>Convert truncated normal to normal</i>
------------	---

Description

Convert a truncated normal distribution to a normal (gaussian) distribution

Usage

```
trunc2norm(x, min = NULL, max = NULL, mu = mean(x), sd = stats::sd(x))
```

Arguments

x	the truncated normally distributed vector
min	the minimum of the truncated distribution (calculated from x if not given)
max	the maximum of the truncated distribution (calculated from x if not given)
mu	the mean of the distribution to return (calculated from x if not given)
sd	the SD of the distribution to return (calculated from x if not given)

Value

a vector with a uniform distribution

Examples

```
x <- truncnorm::rtruncnorm(10000, 1, 7, 3.5, 2)
y <- trunc2norm(x, 1, 7)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

unif2norm

Convert uniform to normal

Description

Convert a uniform distribution to a normal (gaussian) distribution with specified mu and sd

Usage

```
unif2norm(x, mu = 0, sd = 1, min = NULL, max = NULL)
```

Arguments

x	the uniformly distributed vector
mu	the mean of the normal distribution to return
sd	the SD of the normal distribution to return
min	the minimum possible value of x (calculated from x if not given)
max	the maximum possible value of x (calculated from x if not given)

Value

a vector with a gaussian distribution

Examples

```
x <- runif(10000)
y <- unif2norm(x)
g <- ggplot2::ggplot() + ggplot2::geom_point(ggplot2::aes(x, y))
ggExtra::ggMarginal(g, type = "histogram")
```

unique_pairs	<i>Make unique pairs of level names for correlations</i>
--------------	--

Description

Make unique pairs of level names for correlations

Usage

```
unique_pairs(v)
```

Arguments

`v` a vector of level names or a number of levels

Value

a vector of all unique pairs

Examples

```
unique_pairs(c("O", "C", "E", "A", "N"))
unique_pairs(3)
```

wide2long	<i>Convert data from wide to long format</i>
-----------	--

Description

Convert data from wide to long format

Usage

```
wide2long(
  data,
  within_factors = c(),
  within_cols = c(),
  dv = "y",
  id = "id",
  sep = faux_options("sep")
)
```

Arguments

<code>data</code>	the tbl in wide format
<code>within_factors</code>	the names of the within factors
<code>within_cols</code>	the names (or indices) of the within-subject (value) columns
<code>dv</code>	the name of the dv column (defaults to "y")
<code>id</code>	the name of the ID column(s) if they don't exist, a new column will be made (defaults to "id")
<code>sep</code>	separator for within-columns (to be used in <code>strsplit</code> , so can be regex), defaults to " "

Value

a tbl in long format

Examples

```
wide2long(iris, c("Feature", "Measure"), 1:4, sep = "\\.")
```

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