

Package ‘LikertMakeR’

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Type Package

Title Synthesise and Correlate Likert Scale and Rating-Scale Data
Based on Summary Statistics

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Description Generate and correlate synthetic Likert and rating-scale questionnaire responses with predefined means, standard deviations, Cronbach's Alpha, Factor Loading table, coefficients, and other summary statistics.

It can be used to simulate Likert data, construct multi-item scales, generate correlation matrices, and create example survey datasets for teaching statistics, psychometrics, and methodological research.

Worked examples and documentation are available in the package articles, accessible via the package website,

[<https://winzarh.github.io/LikertMakeR/>](https://winzarh.github.io/LikertMakeR/).

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URL <https://github.com/WinzarH/LikertMakeR/>,

<https://winzarh.github.io/LikertMakeR/>,

<https://github.com/WinzarH/LikertMakeR>

BugReports <https://github.com/WinzarH/LikertMakeR/issues>

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alpha

Calculate Cronbach's Alpha from a correlation matrix or dataframe

Description

alpha() calculates Cronbach's Alpha from a given correlation matrix or a given dataframe.

Usage

```
alpha(cormatrix = NULL, data = NULL)
```

Arguments

cormatrix (real) a square symmetrical matrix with values ranging from -1 to +1 and '1' in the diagonal
data (real) a dataframe or matrix

Value

a single value

See Also

[alpha_sensitivity](#), [reliability](#)

Examples

```
## Sample data frame
df <- data.frame(
  V1 = c(4, 2, 4, 3, 2, 2, 2, 1),
  V2 = c(4, 1, 3, 4, 4, 3, 2, 3),
  V3 = c(4, 1, 3, 5, 4, 1, 4, 2),
  V4 = c(4, 3, 4, 5, 3, 3, 3, 3)
)

## example correlation matrix
corMat <- matrix(
  c(
    1.00, 0.35, 0.45, 0.70,
    0.35, 1.00, 0.60, 0.55,
    0.45, 0.60, 1.00, 0.65,
    0.70, 0.55, 0.65, 1.00
  ),
  nrow = 4, ncol = 4
)

## apply function examples

alpha(cormatrix = corMat)

alpha(, df)

alpha(corMat, df)
```

alpha_sensitivity

Sensitivity of Cronbach's alpha to scale design parameters

Description

Computes how Cronbach's alpha changes as a function of either the average inter-item correlation (\bar{r}) or the number of items (k), holding the other parameter constant.

Usage

```
alpha_sensitivity(
  data = NULL,
  k = NULL,
  r_bar = NULL,
  vary = c("r_bar", "k"),
  k_range = NULL,
  r_bar_range = NULL,
  plot = TRUE,
  digits = 3
)
```

Arguments

<code>data</code>	A data frame or matrix of item responses. If provided, k and \bar{r} are computed from the data.
<code>k</code>	Number of items. Required if data is not supplied.
<code>r_bar</code>	Average inter-item correlation. Required if data is not supplied.
<code>vary</code>	Character string indicating which parameter to vary: "r_bar" (default) or "k".
<code>k_range</code>	Numeric vector of item counts to evaluate when vary = "k". Default is 2:20.
<code>r_bar_range</code>	Numeric vector of average inter-item correlations to evaluate when vary = "r_bar". Default is seq(0.05, 0.9, by = 0.05).
<code>plot</code>	Logical; if TRUE, a base R plot is produced. Default is TRUE.
<code>digits</code>	Number of decimal places for rounding output. Default is 3.

Details

The function supports two modes:

- Empirical: derive k and \bar{r} from a dataset
- Theoretical: specify k and \bar{r} directly

Value

A data frame with columns:

- `k`: number of items
- `r_bar`: average inter-item correlation
- `alpha`: Cronbach's alpha

The returned object includes an attribute "baseline" containing the reference k and \bar{r} values.

See Also

[alpha](#), [reliability](#)

Examples

```

# Theoretical example

## Not run:
alpha_sensitivity(k = 6) # produces plot

## End(Not run)

alpha_sensitivity(k = 6, r_bar = 0.4, plot = FALSE)

# Vary number of items
alpha_sensitivity(k = 6, r_bar = 0.4, vary = "k", plot = FALSE)

# Empirical example
df <- data.frame(
  V1 = c(1, 2, 3, 4, 5),
  V2 = c(3, 2, 4, 2, 5),
  V3 = c(2, 1, 5, 4, 3)
)

## Not run:
alpha_sensitivity(data = df) # produces plot

## End(Not run)

alpha_sensitivity(df, vary = "r_bar", plot = FALSE)

alpha_sensitivity(df, vary = "k", plot = FALSE)

```

correlateScales	<i>Dataframe of correlated scales from different dataframes of scale items</i>
-----------------	--

Description

correlateScales() creates a dataframe of scale items representing correlated constructs, as one might find in a completed questionnaire.

Usage

```
correlateScales(dataframes, scalecors)
```

Arguments

dataframes	a list of 'k' dataframes to be rearranged and combined
scalecors	target correlation matrix - should be a symmetric $k \times k$ positive-semi-definite matrix, where 'k' is the number of dataframes

Details

Correlated rating-scale items generally are summed or averaged to create a measure of an "unobservable", or "latent", construct. `correlateScales()` takes several such dataframes of rating-scale items and rearranges their rows so that the scales are correlated according to a predefined correlation matrix. Univariate statistics for each dataframe of rating-scale items do not change, but their correlations with rating-scale items in other dataframes do.

Value

Returns a dataframe whose columns are taken from the starter dataframes and whose summated values are correlated according to a user-specified correlation matrix

Examples

```
## three attitudes and a behavioural intention
n <- 32
lower <- 1
upper <- 5

### attitude #1
cor_1 <- makeCorrAlpha(items = 4, alpha = 0.90)
means_1 <- c(2.5, 2.5, 3.0, 3.5)
sds_1 <- c(0.9, 1.0, 0.9, 1.0)

Att_1 <- makeScales(
  n = n, means = means_1, sds = sds_1,
  lowerbound = rep(lower, 4), upperbound = rep(upper, 4),
  items = 4,
  cormatrix = cor_1
)

### attitude #2
cor_2 <- makeCorrAlpha(items = 5, alpha = 0.85)
means_2 <- c(2.5, 2.5, 3.0, 3.0, 3.5)
sds_2 <- c(1.0, 1.0, 0.9, 1.0, 1.5)

Att_2 <- makeScales(
  n = n, means = means_2, sds = sds_2,
  lowerbound = rep(lower, 5), upperbound = rep(upper, 5),
  items = 5,
  cormatrix = cor_2
)

### attitude #3
cor_3 <- makeCorrAlpha(items = 6, alpha = 0.75)
means_3 <- c(2.5, 2.5, 3.0, 3.0, 3.5, 3.5)
sds_3 <- c(1.0, 1.5, 1.0, 1.5, 1.0, 1.5)

Att_3 <- makeScales(
```

```
n = n, means = means_3, sds = sds_3,
lowerbound = rep(lower, 6), upperbound = rep(upper, 6),
items = 6,
cormatrix = cor_3
)

### behavioural intention
intent <- lfast(n, mean = 3.0, sd = 3, lowerbound = 0, upperbound = 10) |>
  data.frame()
names(intent) <- "int"

### target scale correlation matrix
scale_cors <- matrix(
  c(
    1.0, 0.6, 0.5, 0.3,
    0.6, 1.0, 0.4, 0.2,
    0.5, 0.4, 1.0, 0.1,
    0.3, 0.2, 0.1, 1.0
  ),
  nrow = 4
)

data_frames <- list("A1" = Att_1, "A2" = Att_2, "A3" = Att_3, "Int" = intent)

### apply the function
my_correlated_scales <- correlateScales(
  dataframes = data_frames,
  scalecors = scale_cors
)
head(my_correlated_scales)
```

eigenvalues

calculate eigenvalues of a correlation matrix with optional scree plot

Description

`eigenvalues()` calculates eigenvalues of a correlation matrix and optionally produces a scree plot.

Usage

```
eigenvalues(cormatrix, scree = FALSE)
```

Arguments

`cormatrix` (real, matrix) a correlation matrix
`scree` (logical) default = FALSE. If TRUE (or 1), then `eigenvalues()` produces a scree plot to illustrate the eigenvalues

Value

a vector of eigenvalues
 report on positive-definite status of cormatrix

Examples

```
## define parameters

correlationMatrix <- matrix(
  c(
    1.00, 0.25, 0.35, 0.40,
    0.25, 1.00, 0.70, 0.75,
    0.35, 0.70, 1.00, 0.80,
    0.40, 0.75, 0.80, 1.00
  ),
  nrow = 4, ncol = 4
)

## apply function

evals <- eigenvalues(cormatrix = correlationMatrix)
evals <- eigenvalues(correlationMatrix, 1)
```

lcor	<i>Rearrange elements in each column of a data-frame to fit a predefined correlation matrix</i>
------	---

Description

lcor() rearranges values in each column of a data-frame so that columns are correlated to match a predefined correlation matrix.

Usage

```
lcor(data, target, passes = 10)
```

Arguments

data	dataframe that is to be rearranged
target	target correlation matrix. Must have same dimensions as number of columns in data-frame.
passes	Number of optimization passes (default = 10). Increasing this value <i>MAY</i> improve results if n-columns (target correlation matrix dimensions) are many. Decreasing the value for 'passes' is faster but may decrease accuracy.

Details

Values in a column do not change, so univariate statistics remain the same.

Value

Returns a dataframe whose column-wise correlations approximate a user-specified correlation matrix

Examples

```
## parameters
n <- 32
lowerbound <- 1
upperbound <- 5
items <- 5

mydat3 <- data.frame(
  x1 = lfast(n, 2.5, 0.75, lowerbound, upperbound, items),
  x2 = lfast(n, 3.0, 1.50, lowerbound, upperbound, items),
  x3 = lfast(n, 3.5, 1.00, lowerbound, upperbound, items)
)

cor(mydat3) |> round(3)

tgt3 <- matrix(
  c(
    1.00, 0.50, 0.75,
    0.50, 1.00, 0.25,
    0.75, 0.25, 1.00
  ),
  nrow = 3, ncol = 3
)

## apply function
new3 <- lcor(mydat3, tgt3)

## test output
cor(new3) |> round(3)
```

lexact

Deprecated. Use lfast() instead

Description

lexact is DEPRECATED. Replaced in LikertMakeR Version 0.4.0 by new version of lfast.

lexact remains as a legacy for earlier package users. It is now just a wrapper for lfast

Previously, lexact used a Differential Evolution (DE) algorithm to find an optimum solution with desired mean and standard deviation, but we found that the updated lfast function is much faster and just as accurate.

Also the package is much less bulky.

Usage

```
lexact(n, mean, sd, lowerbound, upperbound, items = 1)
```

Arguments

n	(positive, int) number of observations to generate
mean	(real) target mean
sd	(real) target standard deviation
lowerbound	(positive, int) lower bound
upperbound	(positive, int) upper bound
items	(positive, int) number of items in the rating scale.

Value

a vector of simulated data approximating user-specified conditions.

Examples

```
x <- lexact(  
  n = 256,  
  mean = 4.0,  
  sd = 1.0,  
  lowerbound = 1,  
  upperbound = 7,  
  items = 6  
)  
  
x <- lexact(256, 2, 1.8, 0, 10)
```

lfast

Synthesise rating-scale data with predefined mean and standard deviation

Description

lfast() applies a simple Evolutionary Algorithm to find a vector that best fits the desired moments.

lfast() generates random discrete values from a scaled Beta distribution so the data replicate an ordinal rating scale - for example, a Likert scale made from multiple items (questions) or 0-10 likelihood-of-purchase scale. Data generated are generally consistent with real data, as shown in the *lfast() validation article*.

Usage

```
lfast(n, mean, sd, lowerbound, upperbound, items = 1, precision = 0)
```

Arguments

n	(positive, int) number of observations to generate
mean	(real) target mean, between upper and lower bounds
sd	(positive, real) target standard deviation
lowerbound	(int) lower bound (e.g. '1' for a 1-5 rating scale)
upperbound	(int) upper bound (e.g. '5' for a 1-5 rating scale)
items	(positive, int) number of items in the rating scale. Default = 1
precision	(positive, real) can relax the level of accuracy required. (e.g. '1' generally generates a vector with moments correct within '0.025', '2' generally within '0.05') Default = 0

Value

a vector approximating user-specified conditions.

See Also

- lfast validation article: https://winzarh.github.io/LikertMakeR/articles/lfast_validation.html
- Package website: <https://winzarh.github.io/LikertMakeR/>

Examples

```
## six-item 1-7 rating scale
x <- lfast(
  n = 256,
  mean = 4.0,
  sd = 1.25,
  lowerbound = 1,
  upperbound = 7,
  items = 6
)
```

```
## five-item -3 to +3 rating scale
x <- lfast(
  n = 64,
  mean = 0.025,
  sd = 1.25,
  lowerbound = -3,
  upperbound = 3,
  items = 5
)
```

```
## four-item 1-5 rating scale with medium variation
```

```
x <- lfast(
  n = 128,
  mean = 3.0,
  sd = 1.00,
  lowerbound = 1,
  upperbound = 5,
  items = 4,
  precision = 5
)

## eleven-point 'likelihood of purchase' scale
x <- lfast(256, 3, 3.0, 0, 10)
```

makeCorrAlpha

Correlation matrix from Cronbach's Alpha

Description

Generate a Positive-Definite Correlation Matrix for a target *Cronbach's alpha*.

Usage

```
makeCorrAlpha(
  items,
  alpha,
  variance = 0.1,
  alpha_noise = 0,
  diagnostics = FALSE
)
```

Arguments

items	Integer. Number of items (≥ 2).
alpha	Numeric. Target Cronbach's alpha ($0 < \alpha < 1$).
variance	Numeric. Controls heterogeneity of item loadings in the underlying one-factor model. Larger values produce greater dispersion among item loadings, which results in a wider spread of inter-item correlations while preserving the requested Cronbach's alpha. Typical guidance: <ul style="list-style-type: none"> • 0.05 — near-parallel items (very similar correlations) • 0.10 — modest heterogeneity (default) • 0.15 — strong heterogeneity • 0.20 — very strong heterogeneity • > 0.25 — extreme dispersion; internal shrinkage may occur

For most applied psychometric scales ($k < 20$), values between 0.05 and 0.15 produce realistic correlation structures. Values above 0.30 are automatically reduced to 0.30 to satisfy algorithm constraints.

alpha_noise	<p>Numeric. Controls random variation in the target Cronbach's alpha before the correlation matrix is constructed.</p> <p>When alpha_noise = 0 (default), the requested alpha is reproduced deterministically (subject to numerical tolerance).</p> <p>When alpha_noise > 0, a small amount of random variation is added to the requested alpha prior to constructing the matrix. This can be useful in teaching or simulation settings where slightly different reliability values are desired across repeated runs.</p> <p>Internally, noise is added on the Fisher z-transformed scale to ensure the resulting alpha remains within valid bounds (0, 1).</p> <p>Typical guidance:</p> <ul style="list-style-type: none"> • 0.00 — deterministic alpha (default) • 0.02 — very small variation • 0.05 — moderate variation • 0.10 — substantial variation (caution) <p>Larger values increase the spread of achieved alpha across runs.</p>
diagnostics	<p>Logical. If TRUE, returns a list containing the matrix and diagnostic information.</p>

Details

Constructs a correlation matrix with a specified number of items and target *Cronbach's alpha* using a constructive one-factor model.

Such a correlation matrix can be applied to the `makeScales()` function to generate synthetic data with the predefined alpha.

The algorithm directly builds a positive-definite correlation matrix by solving for item loadings that reproduce the desired average inter-item correlation implied by *alpha*. Unlike earlier versions of this function, this method guarantees positive definiteness by construction, without *post-hoc* repair.

The function computes the average inter-item correlation implied by the requested alpha and solves for a one-factor loading structure that reproduces this value. A small adaptive reduction in dispersion may be applied when necessary to ensure a valid positive-definite solution.

The constructive generator assumes a single common factor structure, consistent with typical psychometric scale construction.

Value

If `diagnostics = FALSE`, a positive-definite correlation matrix. If `diagnostics = TRUE`, a list containing:

- R: the correlation matrix
- diagnostics: list including achieved alpha, minimum eigenvalue, and internal variance used

Examples

```
# Example 1
# define parameters
items <- 4
alpha <- 0.85

# apply function
set.seed(42)
cor_matrix <- makeCorrAlpha(
  items = items,
  alpha = alpha
)

# test function output
print(cor_matrix) |> round(3)
alpha(cor_matrix)
eigenvalues(cor_matrix, 1)

# Example 2
# higher alpha, more items, more variability
cor_matrix2 <- makeCorrAlpha(
  items = 8,
  alpha = 0.95,
  variance = 0.10
)

# test output
cor_matrix2 |> round(2)
alpha(cor_matrix2) |> round(3)
eigenvalues(cor_matrix2, 1) |> round(3)

# Example 3
# large random variation around alpha
cor_matrix3 <- makeCorrAlpha(
  items = 6,
  alpha = 0.85,
  alpha_noise = 0.10
)

# test output
cor_matrix3 |> round(2)
alpha(cor_matrix3) |> round(3)
eigenvalues(cor_matrix3, 1) |> round(3)

# Example 4
# with diagnostics
cor_matrix4 <- makeCorrAlpha(
  items = 4,
  alpha = 0.80,
  diagnostics = TRUE
)
```

```
# test output
cor_matrix4
```

makeCorrLoadings	<i>Generate Inter-Item Correlation Matrix from Factor Loadings</i>
------------------	--

Description

Constructs an inter-item correlation matrix based on a user-supplied matrix of standardised factor loadings and (optionally) a factor correlation matrix. The `makeCorrLoadings()` function does a surprisingly good job of reproducing a target correlation matrix when all item-factor loadings are present, as shown in the *makeCorrLoadings() validation article*.

Usage

```
makeCorrLoadings(
  loadings,
  factorCor = NULL,
  uniquenesses = NULL,
  nearPD = FALSE,
  diagnostics = FALSE
)
```

Arguments

loadings	Numeric matrix. A $k \times f$ matrix of standardized factor loadings <i>items</i> \times <i>factors</i> . Row names and column names are used for diagnostics if present.
factorCor	Optional $f \times f$ matrix of factor correlations (Φ). If NULL, assumes orthogonal factors.
uniquenesses	Optional vector of length k . If NULL, calculated as $1 - \text{rowSums}(\text{loadings}^2)$.
nearPD	Logical. If TRUE, attempts to coerce non-positive-definite matrices using <code>Matrix::nearPD()</code> .
diagnostics	Logical. If TRUE, returns diagnostics including McDonald's Omega and item-level summaries.

Value

If `diagnostics = FALSE`, returns a correlation matrix (class: `matrix`). If `diagnostics = TRUE`, returns a list with: - R: correlation matrix - Omega: per-factor Omega or adjusted Omega - OmegaTotal: total Omega across all factors - Diagnostics: dataframe of communalities, uniquenesses, and primary factor

See Also

- `makeCorrLoadings()` validation article: https://winzarh.github.io/LikertMakeR/articles/makeCorrLoadings_validate.html
- Package website: <https://winzarh.github.io/LikertMakeR/>

Examples

```

# -----
# Example 1: Basic use without diagnostics
# -----

factorLoadings <- matrix(
  c(
    0.05, 0.20, 0.70,
    0.10, 0.05, 0.80,
    0.05, 0.15, 0.85,
    0.20, 0.85, 0.15,
    0.05, 0.85, 0.10,
    0.10, 0.90, 0.05,
    0.90, 0.15, 0.05,
    0.80, 0.10, 0.10
  ),
  nrow = 8, ncol = 3, byrow = TRUE
)

rownames(factorLoadings) <- paste0("Q", 1:8)
colnames(factorLoadings) <- c("Factor1", "Factor2", "Factor3")

factorCor <- matrix(
  c(
    1.0, 0.7, 0.6,
    0.7, 1.0, 0.4,
    0.6, 0.4, 1.0
  ),
  nrow = 3, byrow = TRUE
)

itemCor <- makeCorrLoadings(factorLoadings, factorCor)
round(itemCor, 3)

# -----
# Example 2: Diagnostics with factor correlations (Adjusted Omega)
# -----

result_adj <- makeCorrLoadings(
  loadings = factorLoadings,
  factorCor = factorCor,
  diagnostics = TRUE
)

# View outputs
round(result_adj$R, 3) # correlation matrix
round(result_adj$Omega, 3) # adjusted Omega
round(result_adj$OmegaTotal, 3) # total Omega
print(result_adj$Diagnostics) # communality and uniqueness per item

# -----
# Example 3: Diagnostics assuming orthogonal factors (Per-Factor Omega)

```

```
# -----
result_orth <- makeCorrLoadings(
  loadings = factorLoadings,
  diagnostics = TRUE
)

round(result_orth$Omega, 3) # per-factor Omega
round(result_orth$OmegaTotal, 3) # total Omega
print(result_orth$Diagnostics)
```

makeItemsScale	<i>Generate scale items from a summated scale, with desired Cronbach's Alpha</i>
----------------	--

Description

makeItemsScale() generates a random dataframe of scale items based on a predefined summated scale (such as created by the [lfast\(\)](#) function), and a desired *Cronbach's Alpha*.

Usage

```
makeItemsScale(
  scale,
  lowerbound,
  upperbound,
  items,
  alpha = 0.8,
  summated = TRUE,
  progress = TRUE
)
```

Arguments

scale	(int) a vector or dataframe of the summated rating scale. Should range from $lowerbound \times items$ to $upperbound \times items$
lowerbound	(int) lower bound of the scale item (example: '1' in a '1' to '5' rating)
upperbound	(int) upper bound of the scale item (example: '5' in a '1' to '5' rating)
items	(positive, int) k, or number of columns to generate
alpha	(positive, real) desired <i>Cronbach's Alpha</i> for the new dataframe of items. Default = '0.8'.
summated	logical. If TRUE (default), scale is assumed to be a summated scale. If FALSE, the input is treated as a mean scale and will be multiplied by items internally.
progress	logical. If TRUE (default is interactive()), a progress bar is displayed during optimisation. Set to FALSE for slightly faster execution.

Details

simulated annealing:

Simulated annealing is used in `makeItemsScale()` to search for item configurations that satisfy several competing objectives simultaneously. These include achieving a target Cronbach's alpha, maintaining similar item means, and preserving constant row sums. Because improving one objective can worsen another, a simple greedy search often becomes trapped in suboptimal solutions. Simulated annealing addresses this by allowing occasional acceptance of worse solutions early in the search, promoting exploration of the solution space. As the algorithm progresses, a cooling schedule reduces this randomness, increasingly favouring improvements across all criteria. This process enables the function to converge on a balanced solution that jointly satisfies the desired scale properties, rather than optimizing any single objective at the expense of others.

Value

a dataframe with 'items' columns and 'length(scale)' rows

Examples

```
summatedScale <- c(16, 11, 14, 7, 10, 13, 16, 13, 12, 14, 8, 11, 8, 18, 8, 13)

newItems <- makeItemsScale(
  scale = summatedScale,
  lowerbound = 1,
  upperbound = 5,
  items = 4,
  alpha = 0.8
)
```

makePaired

Synthesise a dataset from paired-sample t-test summary statistics

Description

Generates a dataset from paired-sample *t*-test summary statistics.

Usage

```
makePaired(
  n,
  means,
  sds,
  t_value,
  lowerbound,
  upperbound,
  items = 1,
  precision = 0
)
```

Arguments

n	(positive, integer) sample size
means	(real) 1:2 vector of target means for two before/after measures
sds	(real) 1:2 vector of target standard deviations
t_value	(real) desired paired <i>t</i> -statistic
lowerbound	(integer) lower bound (e.g. '1' for a 1-5 rating scale)
upperbound	(integer) upper bound (e.g. '5' for a 1-5 rating scale)
items	(positive, integer) number of items in the rating scale. Default = 1
precision	(positive, real) relaxes the level of accuracy required. Default = 0

Details

makePaired() generates correlated values so the data replicate rating scales taken, for example, in a before and after experimental design.

The function is effectively a wrapper function for lfast() and lcor() with the addition of a *t*-statistic from which the between-column correlation is inferred.

Paired *t*-tests apply to observations that are associated with each other. For example: the same people before and after a treatment; the same people rating two different objects; ratings by husband & wife; *etc.*

The paired-samples *t*-test is defined as:

$$t = \frac{\text{mean}(D)}{\text{sd}(D)/\sqrt{n}}$$

where:

- D = differences in values
- $\text{mean}(D)$ = mean of the differences
- $\text{sd}(D)$ = standard deviation of the differences, where

$$\text{sd}(D)^2 = \text{sd}(X_{\text{before}})^2 + \text{sd}(X_{\text{after}})^2 - 2 \text{cov}(X_{\text{before}}, X_{\text{after}})$$

A paired-sample *t*-test thus requires an estimate of the covariance between the two sets of observations. makePaired() rearranges these formulae so that the covariance is inferred from the *t*-statistic.

Value

a dataframe approximating user-specified conditions.

Note

Larger sample sizes usually result in higher t -statistics, and correspondingly small p -values.

Small sample sizes with relatively large standard deviations and relatively high t -statistics can result in impossible correlation values.

Similarly, large sample sizes with low t -statistics can result in impossible correlations. That is, a correlation outside of the $-1:1$ range.

If this happens, the function will fail with an *ERROR* message. The user should review the input parameters and insert more realistic values.

See Also

[lfast](#), [lcor](#)

Examples

```
n <- 20
pair_m <- c(2.5, 3.0)
pair_s <- c(1.0, 1.5)
lower <- 1
upper <- 5
k <- 6
t <- -2.5

pairedDat <- makePaired(
  n = n, means = pair_m, sds = pair_s,
  t_value = t,
  lowerbound = lower, upperbound = upper, items = k
)

str(pairedDat)
cor(pairedDat) |> round(2)

t.test(pairedDat$X1, pairedDat$X2, paired = TRUE)
```

makeRepeated

Reproduce Repeated-Measures Data from ANOVA Summary Statistics

Description

Constructs a synthetic dataset and inter-timepoint correlation matrix from a repeated-measures ANOVA result, based on reported means, standard deviations, and an F -statistic. This is useful when only summary statistics are available from published studies.

Usage

```

makeRepeated(
  n,
  k,
  means,
  sds,
  f_stat,
  df_between = k - 1,
  df_within = (n - 1) * (k - 1),
  structure = c("cs", "ar1", "toeplitz"),
  names = paste0("time_", 1:k),
  items = 1,
  lowerbound = 1,
  upperbound = 5,
  return_corr_only = FALSE,
  diagnostics = FALSE,
  ...
)

```

Arguments

n	Integer. Sample size used in the original study.
k	Integer. Number of repeated measures (timepoints).
means	Numeric vector of length k. Mean values reported for each timepoint.
sds	Numeric vector of length k. Standard deviations reported for each timepoint.
f_stat	Numeric. The reported repeated-measures ANOVA F-statistic for the within-subjects factor.
df_between	Degrees of freedom between conditions (default: k - 1).
df_within	Degrees of freedom within-subjects (default: (n - 1) * (k - 1)).
structure	Character. Correlation structure to assume: "cs", "ar1", or "toeplitz" (default = "cs").
names	Character vector of length k. Variable names for each timepoint (default: "time_1" to "time_k").
items	Integer. Number of items used to generate each scale score (passed to lfast).
lowerbound	Integer. Lower bounds for Likert-type response scales (default: 1).
upperbound	Integer. upper bounds for Likert-type response scales (default: 5).
return_corr_only	Logical. If TRUE, return only the estimated correlation matrix.
diagnostics	Logical. If TRUE, include diagnostic summaries such as feasible F-statistic range and effect sizes.
...	Reserved for future use.

Details

This function estimates the average correlation between repeated measures by matching the reported F-statistic, under one of three assumed correlation structures:

- "cs" (*Compound Symmetry*): The Default. Assumes all timepoints are equally correlated. Common in standard RM-ANOVA settings.
- "ar1" (*First-Order Autoregressive*): Assumes correlations decay exponentially with time lag.
- "toeplitz" (*Linearly Decreasing*): Assumes correlation declines linearly with time lag - a middle ground between "cs" and "ar1".

The function then generates a data frame of synthetic item-scale ratings using [lfast](#), and adjusts them to match the estimated correlation structure using [lcor](#).

Set `return_corr_only = TRUE` to extract only the estimated correlation matrix.

Value

A named list with components:

`data` A data frame of simulated repeated-measures responses (unless `return_corr_only = TRUE`).

`correlation_matrix` The estimated inter-timepoint correlation matrix.

`structure` The correlation structure assumed.

`achieved_f` The F-statistic produced by the estimated rho value (if `diagnostics = TRUE`).

`feasible_f_range` Minimum and maximum achievable F-values under the chosen structure (shown if diagnostics are requested).

`recommended_f` Conservative, moderate, and strong F-statistic suggestions for similar designs.

`effect_size_raw` Unstandardised effect size across timepoints.

`effect_size_standardised` Effect size standardised by average variance.

See Also

[lfast](#), [lcor](#)

Examples

```
set.seed(42)

out1 <- makeRepeated(
  n = 64,
  k = 3,
  means = c(3.1, 3.5, 3.9),
  sds = c(1.0, 1.1, 1.0),
  items = 4,
  f_stat = 4.87,
  structure = "cs",
  diagnostics = FALSE
)

head(out1$data)
```

```
out1$correlation_matrix

out2 <- makeRepeated(
  n = 32, k = 4,
  means = c(2.75, 3.5, 4.0, 4.4),
  sds = c(0.8, 1.0, 1.2, 1.0),
  f_stat = 16,
  structure = "ar1",
  items = 5,
  lowerbound = 1, upperbound = 7,
  return_corr_only = FALSE,
  diagnostics = TRUE
)

print(out2)

out3 <- makeRepeated(
  n = 64, k = 4,
  means = c(2.0, 2.25, 2.75, 3.0),
  sds = c(0.8, 0.9, 1.0, 0.9),
  items = 4,
  f_stat = 24,
  # structure = "toeplitz",
  diagnostics = TRUE
)

str(out3)
```

makeScales	<i>Synthesise rating-scale data with given first and second moments and a predefined correlation matrix</i>
------------	---

Description

Generates a dataframe of random discrete values so the data replicate a rating scale, and are correlated close to a predefined correlation matrix.

Usage

```
makeScales(n, means, sds, lowerbound = 1, upperbound = 5, items = 1, cormatrix)
```

Arguments

n	(positive, int) sample-size - number of observations
means	(real) target means: a vector of length k of mean values for each scale item
sds	(positive, real) target standard deviations: a vector of length k of standard deviation values for each scale item

lowerbound	(positive, int) a vector of length k (same as rows & columns of correlation matrix) of values for lower bound of each scale item (e.g. '1' for a 1-5 rating scale). Default = 1.
upperbound	(positive, int) a vector of length k (same as rows & columns of correlation matrix) of values for upper bound of each scale item (e.g. '5' for a 1-5 rating scale). Default = 5.
items	(positive, int) a vector of length k of number of items in each scale. Default = 1.
cormatrix	(real, matrix) the target correlation matrix: a square symmetric positive-semi-definite matrix of values ranging between -1 and +1, and '1' in the diagonal.

Details

makeScales() is wrapper function for:

- [lfast\(\)](#), generates a dataframe that best fits the desired moments, and
- [lcor\(\)](#), which rearranges values in each column of the dataframe so they closely match the desired correlation matrix.

Value

a dataframe of rating-scale values

See Also

[lfast](#), [lcor](#)

Examples

```
## Example 1: four correlated items (questions)

### define parameters

n <- 16
dfMeans <- c(2.5, 3.0, 3.0, 3.5)
dfSds <- c(1.0, 1.0, 1.5, 0.75)
lowerbound <- rep(1, 4)
upperbound <- rep(5, 4)

corMat <- matrix(
  c(
    1.00, 0.30, 0.40, 0.60,
    0.30, 1.00, 0.50, 0.70,
    0.40, 0.50, 1.00, 0.80,
    0.60, 0.70, 0.80, 1.00
  ),
  nrow = 4, ncol = 4
)

scale_names <- c("Q1", "Q2", "Q3", "Q4")
rownames(corMat) <- scale_names
```

```
colnames(corMat) <- scale_names

### apply function

df1 <- makeScales(
  n = n, means = dfMeans, sds = dfSds,
  lowerbound = lowerbound, upperbound = upperbound, cormatrix = corMat
)

### test function

str(df1)

#### means
apply(df1, 2, mean) |> round(3)

#### standard deviations
apply(df1, 2, sd) |> round(3)

#### correlations
cor(df1) |> round(3)

## Example 2: five correlated Likert scales

### a study on employee engagement and organizational climate:
# Job Satisfaction (JS)
# Organizational Commitment (OC)
# Perceived Supervisor Support (PSS)
# Work Engagement (WE)
# Turnover Intention (TI) (reverse-related to others).

### define parameters

n <- 128
dfMeans <- c(3.8, 3.6, 3.7, 3.9, 2.2)
dfSds <- c(0.7, 0.8, 0.7, 0.6, 0.9)
lowerbound <- rep(1, 5)
upperbound <- rep(5, 5)
items <- c(4, 4, 3, 3, 3)

corMat <- matrix(
  c(
    1.00, 0.72, 0.58, 0.65, -0.55,
    0.72, 1.00, 0.54, 0.60, -0.60,
    0.58, 0.54, 1.00, 0.57, -0.45,
    0.65, 0.60, 0.57, 1.00, -0.50,
    -0.55, -0.60, -0.45, -0.50, 1.00
  ),
  nrow = 5, ncol = 5
)

scale_names <- c("JS", "OC", "PSS", "WE", "TI")
```

```

rownames(corMat) <- scale_names
colnames(corMat) <- scale_names

### apply function

df2 <- makeScales(
  n = n, means = dfMeans, sds = dfSds,
  lowerbound = lowerbound, upperbound = upperbound,
  items = items, cormatrix = corMat
)

### test function

str(df2)

#### means
apply(df2, 2, mean) |> round(3)

#### standard deviations
apply(df2, 2, sd) |> round(3)

#### correlations
cor(df2) |> round(3)

```

makeScalesRegression *Generate Data from Multiple-Regression Summary Statistics*

Description

Generates synthetic rating-scale data that replicates reported regression results. This function is useful for reproducing analyses from published research where only summary statistics (standardised regression coefficients and R-squared) are reported.

Usage

```

makeScalesRegression(
  n,
  beta_std,
  r_squared,
  iv_cormatrix = NULL,
  iv_cor_mean = 0.3,
  iv_cor_variance = 0.01,
  iv_cor_range = c(-0.7, 0.7),
  iv_means,
  iv_sds,
  dv_mean,
  dv_sd,
  lowerbound_iv,

```

```

    upperbound_iv,
    lowerbound_dv,
    upperbound_dv,
    items_iv = 1,
    items_dv = 1,
    var_names = NULL,
    tolerance = 0.005
)

```

Arguments

n	Integer. Sample size
beta_std	Numeric vector of standardised regression coefficients (length k)
r_squared	Numeric. R-squared from regression (-1 to 1)
iv_cormatrix	k x k correlation matrix of independent variables. If missing (NULL), will be optimised.
iv_cor_mean	Numeric. Mean correlation among IVs when optimising (ignored if iv_cormatrix provided). Default = 0.3
iv_cor_variance	Numeric. Variance of correlations when optimising (ignored if iv_cormatrix provided). Default = 0.01
iv_cor_range	Numeric vector of length 2. Min and max constraints on correlations when optimising. Default = c(-0.7, 0.7)
iv_means	Numeric vector of means for IVs (length k)
iv_sds	Numeric vector of standard deviations for IVs (length k)
dv_mean	Numeric. Mean of dependent variable
dv_sd	Numeric. Standard deviation of dependent variable
lowerbound_iv	Numeric vector of lower bounds for each IV scale (or single value for all)
upperbound_iv	Numeric vector of upper bounds for each IV scale (or single value for all)
lowerbound_dv	Numeric. Lower bound for DV scale
upperbound_dv	Numeric. Upper bound for DV scale
items_iv	Integer vector of number of items per IV scale (or single value for all). Default = 1
items_dv	Integer. Number of items in DV scale. Default = 1
var_names	Character vector of variable names (length k+1: IVs then DV)
tolerance	Numeric. Acceptable deviation from target R-squared (default 0.005)

Details

Generate regression data from summary statistics

The function can operate in two modes:

Mode 1: With IV correlation matrix provided

When `iv_cormatrix` is provided, the function uses the given correlation structure among independent variables and calculates the implied IV-DV correlations from the regression coefficients.

Mode 2: With optimisation (IV correlation matrix not provided)

When `iv_cormatrix = NULL`, the function optimises to find a plausible correlation structure among independent variables that matches the reported regression statistics. Initial correlations are sampled using Fisher's z-transformation to ensure proper distribution, then iteratively adjusted to match the target R-squared.

The function generates Likert-scale data (not individual items) using `lfast()` for each variable with specified moments, then correlates them using `lcor()`. Generated data are verified by running a regression and comparing achieved statistics with targets.

Value

A list containing:

<code>data</code>	Generated dataframe with k IVs and 1 DV
<code>target_stats</code>	List of target statistics provided
<code>achieved_stats</code>	List of achieved statistics from generated data
<code>diagnostics</code>	Comparison of target vs achieved
<code>iv_dv_cors</code>	Calculated correlations between IVs and DV
<code>full_cormatrix</code>	The complete (k+1) x (k+1) correlation matrix used
<code>optimisation_info</code>	If IV correlations were optimised, details about the optimisation

See Also

[lfast](#) for generating individual rating-scale vectors with exact moments.

[lcor](#) for rearranging values to achieve target correlations.

[makeCorrAlpha](#) for generating correlation matrices from Cronbach's Alpha.

Examples

```
# Example 1: With provided IV correlation matrix
set.seed(123)
iv_corr <- matrix(c(1.0, 0.3, 0.3, 1.0), nrow = 2)

result1 <- makeScalesRegression(
  n = 64,
  beta_std = c(0.4, 0.3),
  r_squared = 0.35,
  iv_cormatrix = iv_corr,
  iv_means = c(3.0, 3.5),
  iv_sds = c(1.0, 0.9),
  dv_mean = 3.8,
  dv_sd = 1.1,
  lowerbound_iv = 1,
  upperbound_iv = 5,
```

```
    lowerbound_dv = 1,
    upperbound_dv = 5,
    items_iv = 4,
    items_dv = 4,
    var_names = c("Attitude", "Intention", "Behaviour")
  )

print(result1)
head(result1$data)

# Example 2: With optimisation (no IV correlation matrix)
set.seed(456)
result2 <- makeScalesRegression(
  n = 128,
  beta_std = c(0.3, 0.25, 0.2),
  r_squared = 0.40,
  iv_cormatrix = NULL, # Will be optimised
  iv_cor_mean = 0.3,
  iv_cor_variance = 0.02,
  iv_means = c(3.0, 3.2, 2.8),
  iv_sds = c(1.0, 0.9, 1.1),
  dv_mean = 3.5,
  dv_sd = 1.0,
  lowerbound_iv = 1,
  upperbound_iv = 5,
  lowerbound_dv = 1,
  upperbound_dv = 5,
  items_iv = 4,
  items_dv = 5
)

# View optimised correlation matrix
print(result2$target_stats$iv_cormatrix)
print(result2$optimisation_info)
```

ordinal_diagnostics *Extract ordinal diagnostics from a reliability() result*

Description

Extract ordinal diagnostics from a `reliability()` result

Usage

```
ordinal_diagnostics(x)
```

Arguments

x An object returned by `reliability()`.

Value

A data.frame describing observed response categories and sparsity checks, or NULL if no diagnostics are available.

reliability	<i>Estimate scale reliability for Likert and rating-scale data</i>
-------------	--

Description

Computes internal consistency reliability estimates for a single-factor scale, including Cronbach's alpha, McDonald's omega (total), and optional ordinal (polychoric-based) variants. Confidence intervals may be obtained via nonparametric bootstrap.

Usage

```
reliability(
  data,
  include = "none",
  ci = FALSE,
  ci_level = 0.95,
  n_boot = 1000,
  na_method = c("pairwise", "listwise"),
  min_count = 2,
  digits = 3,
  verbose = TRUE
)
```

Arguments

data	A data frame or matrix containing item responses. Each column represents one item; rows represent respondents.
include	Character vector specifying which additional estimates to compute. Possible values are: <ul style="list-style-type: none"> • "none" (default): Pearson-based alpha and omega only. • "lambda6": Include Guttman's lambda-6 (requires package psych). • "polychoric": Include ordinal (polychoric-based) alpha and omega. Multiple options may be supplied.
ci	Logical; if TRUE, confidence intervals are computed using nonparametric bootstrap. Default is FALSE.
ci_level	Confidence level for bootstrap intervals. Default is 0.95.
n_boot	Number of bootstrap resamples used when ci = TRUE. Default is 1000.
na_method	Method for handling missing values. Either "pairwise" (default) or "listwise".
min_count	Minimum observed frequency per response category required to attempt polychoric correlations. Ordinal reliability estimates are skipped if this condition is violated. Default is 2.

digits	Number of decimal places used when printing estimates. Default is 3.
verbose	Logical; if TRUE, warnings and progress indicators are displayed. Default is TRUE.

Details

The function is designed for Likert-type and rating-scale data and prioritises transparent diagnostics when ordinal reliability estimates are not feasible due to sparse response categories.

Cronbach's alpha and McDonald's omega are computed from Pearson correlations. When `include = "polychoric"`, ordinal reliability estimates are computed using polychoric correlations and correspond to Zumbo's alpha and ordinal omega.

Ordinal reliability estimates are skipped if response categories are sparse or if polychoric estimation fails. Diagnostics explaining these decisions are stored in the returned object and may be inspected using [ordinal_diagnostics](#).

This function assumes a single common factor and is not intended for multidimensional or structural equation modelling contexts.

Value

A tibble with one row per reliability coefficient and columns:

- `coef_name`: Name of the reliability coefficient.
- `estimate`: Point estimate.
- `ci_lower`, `ci_upper`: Confidence interval bounds (only present when `ci = TRUE`).
- `notes`: Methodological notes describing how the estimate was obtained.

The returned object has class `"likert_reliability"` and includes additional attributes containing diagnostics and bootstrap information.

See Also

[ordinal_diagnostics](#)
[alpha_sensitivity](#), [alpha](#)

Examples

```
## create dataset
my_cor <- LikertMakeR::makeCorrAlpha(
  items = 4,
  alpha = 0.80
)

my_data <- LikertMakeR::makeScales(
  n = 64,
  means = c(2.75, 3.00, 3.25, 3.50),
  sds = c(1.25, 1.50, 1.30, 1.25),
  lowerbound = rep(1, 4),
  upperbound = rep(5, 4),
  cormatrix = my_cor
)
```

```
)

## run function
reliability(my_data)

reliability(
  my_data,
  include = c("lambda6", "polychoric")
)

## slower (not run on CRAN checks)
reliability(
  my_data,
  include = "polychoric",
  ci = TRUE,
  n_boot = 200
)
```

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