

Package ‘psychonetrics’

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Type Package

Title Structural Equation Modeling and Confirmatory Network Analysis

Version 0.15

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Description

Multi-group (dynamical) structural equation models in combination with confirmatory network models from cross-sectional, time-series and panel data <doi:10.31234/osf.io/8ha93>. Allows for confirmatory testing and fit as well as exploratory model search.

License GPL-2

LinkingTo Rcpp (>= 0.11.3), RcppArmadillo, RcppEigen, pbv, roptim

Depends R (>= 4.3.0)

Imports methods, qgraph, numDeriv, dplyr, abind, Matrix (>= 1.6-5),
lavaan, corpcor, glasso, mgcv, optimx, nloptr, VCA, pbapply,
parallel, magrittr, IsingSampler, tidy, psych, GA, combinat,
rlang

Suggests psychTools, semPlot, graphicalVAR, metaSEM, mvtnorm, ggplot2

ByteCompile true

URL <https://psychonetrics.org/>

BugReports <https://github.com/SachaEpskamp/psychonetrics/issues>

StagedInstall true

NeedsCompilation yes

RoxygenNote 7.3.3

Author Sacha Epskamp [aut, cre]

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psychometrics-package *Structural Equation Modeling and Confirmatory Network Analysis*

Description

Multi-group (dynamical) structural equation models in combination with confirmatory network models from cross-sectional, time-series and panel data <doi:10.31234/osf.io/8ha93>. Allows for confirmatory testing and fit as well as exploratory model search.

Details

The DESCRIPTION file:

```

Package:      psychometrics
Type:        Package
Title:       Structural Equation Modeling and Confirmatory Network Analysis
Version:     0.15
Authors@R:   person(given = "Sacha",family = "Epskamp", role = c("aut", "cre"), email = "mail@sachaepskamp.com")
Maintainer:  Sacha Epskamp <mail@sachaepskamp.com>
Description: Multi-group (dynamical) structural equation models in combination with confirmatory network models
License:     GPL-2
LinkingTo:   Rcpp (>= 0.11.3), RcppArmadillo, RcppEigen, pbv, roptim
Depends:     R (>= 4.3.0)
Imports:     methods, qgraph, numDeriv, dplyr, abind, Matrix (>= 1.6-5), lavaan, corpcor, glasso, mgcv, optimx, nloptr
Suggests:   psychTools, semPlot, graphicalVAR, metaSEM, mvtnorm, ggplot2
ByteCompile: true
URL:        https://psychometrics.org/
BugReports:  https://github.com/SachaEpskamp/psychometrics/issues
StagedInstall: true
NeedsCompilation: yes
RoxygenNote: 7.3.3
Author:     Sacha Epskamp [aut, cre]

```

Index of help topics:

CIplot	Plot Analytic Confidence Intervals
Ising	Ising model
Jonas	Jonas dataset
MIs	Print modification indices
NA2020	Network Analysis 2020 Self-Report Data
StarWars	Star Wars dataset
addMIs	Model updating functions
aggregate_bootstraps	Aggregate Bootstrapped Models
bifactor	Bi-factor models
bootstrap	Bootstrap a psychonetrics model
changedata	Change the data of a psychonetrics object
checkJacobian	Diagnostic functions
compare	Model comparison
covML	Maximum likelihood covariance estimate
dlvm1	Lag-1 dynamic latent variable model family of psychonetrics models for panel data
duplicationMatrix	Model matrices used in derivatives
emergencystart	Reset starting values to simple defaults
esa	Ergodic Subspace Analysis
factorscores	Compute factor scores
find_penalized_lambda	Automated Lambda Selection for Penalized Estimation
fit	Print fit indices
fixpar	Parameters modification
fixstart	Attempt to Fix Starting Values
generate	Generate data from a fitted psychonetrics object
getVCOV	Obtain the asymptotic covariance matrix
getmatrix	Extract an estimated matrix
groupequal	Group equality constrains
latentgrowth	Latnet growth curve model
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meta_var1	Meta-analytic VAR(1) model
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ml_tsdlvm1	Multi-level Lag-1 dynamic latent variable model family of psychonetrics models for time-series data
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parequal	Set equality constrains across parameters
partialprune	Partial pruning of multi-group models
penalize	Penalized Maximum Likelihood Functions

prune	Stepdown model search by pruning non-significant parameters.
psychonetrics-class	Class '"psychonetrics"'
psychonetrics-package	Structural Equation Modeling and Confirmatory Network Analysis
psychonetrics_bootstrap-class	Class '"psychonetrics_bootstrap"'
psychonetrics_log-class	Class '"psychonetrics"'
ri_clpm	Random intercept cross-lagged panel models
runmodel	Run a psychonetrics model
setestimator	Convenience functions
setverbose	Should messages of computation progress be printed?
simplestructure	Generate factor loadings matrix with simple structure
stepup	Stepup model search along modification indices
transmod	Transform between model types
tsdlvm1	Lag-1 dynamic latent variable model family of psychonetrics models for time-series data
unionmodel	Unify models across groups
var1	Lag-1 vector autoregression family of psychonetrics models
varcov	Variance-covariance family of psychonetrics models
write_psychonetrics	Write comprehensive model output to a text file

This package can be used to perform Structural Equation Modeling and confirmatory network modeling. Current implemented families of models are (1) the variance-covariance matrix (`varcov`), (2) the latent variable model (`lvm`), (3) the lag-1 vector autoregression model (`var1`), and (4) the dynamical lag-1 latent variable model for panel data (`dlvm1`) and for time-series data (`tsdlvm1`).

Author(s)

Sacha Epskamp [aut, cre]

Maintainer: Sacha Epskamp <mail@sachaepskamp.com>

References

More information: psychonetrics.org

aggregate_bootstraps *Aggregate Bootstrapped Models*

Description

Aggregates bootstrap results into a `psychonetrics_bootstrap` object

Usage

```
aggregate_bootstraps(sample, bootstraps, remove_problematic = TRUE)
```

Arguments

`sample` The original psychonetrics object (not bootstrapped)

`bootstraps` A list of bootstrapped psychonetrics objects (i.e., using `bootstrap = TRUE`)

`remove_problematic`
 Remove bootstraps that did not converge (sum of absolute gradient > 1)

Details

After running this function, the helper functions `parameters`, `fit`, and `CIplot` can be used to investigate bootstrap results.

Value

An object of the class `psychonetrics_bootstrap`

Author(s)

Sacha Epskamp

bifactor

Bi-factor models

Description

Wrapper to `lvm` to specify a bi-factor model.

Usage

```
bifactor(data, lambda, latents, bifactor = "g", ...)
```

Arguments

`data` The data as used by `lvm`

`lambda` The factor loadings matrix **without** the bifactor, as used by `lvm`

`latents` A vector of names of the latent variables, as used by `lvm`

`bifactor` Name of the bifactor

`...` Arguments sent to `lvm`

Value

An object of the class `psychonetrics` ([psychonetrics-class](#))

Author(s)

Sacha Epskamp

`bootstrap`*Bootstrap a psychometrics model*

Description

This function will bootstrap the data (once) and return a new unevaluated psychometrics object. It requires `storedata = TRUE` to be used when forming a model.

Usage

```
bootstrap(x, replacement = TRUE, proportion = 1, verbose = TRUE, storedata = FALSE,
          baseline_saturated = TRUE)
```

Arguments

<code>x</code>	A psychometrics model.
<code>replacement</code>	Logical, should new samples be drawn with replacement?
<code>proportion</code>	Proportion of sample to be drawn. Set to lower than 1 for subsampling.
<code>verbose</code>	Logical, should messages be printed?
<code>storedata</code>	Logical, should the bootstrapped data also be stored?
<code>baseline_saturated</code>	Logical, should the baseline and saturated models be included?

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

changedata	<i>Change the data of a psychonetrics object</i>
------------	--

Description

This function can be used to change the data in a psychonetrics object.

Usage

```
changedata(x, data, covs, nobs, means, groups, missing = "listwise")
```

Arguments

x	A psychonetrics model.
data	A data frame encoding the data used in the analysis. Can be missing if covs and nobs are supplied.
covs	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. IMPORTANT NOTE: psychonetrics expects the maximum likelihood (ML) covariance matrix, which is NOT obtained from <code>cov</code> directly. Manually rescale the result of <code>cov</code> with $(nobs - 1)/nobs$ to obtain the ML covariance matrix.
nobs	The number of observations used in covs and means, or a vector of such numbers of observations for multiple groups.
means	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
groups	An optional string indicating the name of the group variable in data.
missing	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be "listwise" for listwise deletion, or "pairwise" for pairwise deletion.

Value

An object of the class psychonetrics ([psychonetrics-class](#))

Author(s)

Sacha Epskamp

CIplot

*Plot Analytic Confidence Intervals***Description**

Function to plot analytic confidence intervals (CI) of matrix elements estimated in psychometrics.

Usage

```
CIplot(x, matrices, alpha_ci = 0.05, alpha_color = c(0.05,
  0.01, 0.001, 1e-04), labels, labels2, labelstart,
  print = TRUE, major_break = 0.2, minor_break = 0.1,
  split0, prop0, prop0_cex = 1, prop0_alpha = 0.95,
  prop0_minAlpha = 0.25)
```

Arguments

x	A psychometrics model.
matrices	Vector of strings indicating the matrices to plot CIs for
alpha_ci	The alpha level used for the CIs
alpha_color	A vector of alphas used for coloring the CIs
labels	The labels for the variables associated with the rows of a matrix.
labels2	The labels for the variables associated with the columns of a matrix. Defaults to the value of labels for square matrices.
labelstart	The value to determine if labels are printed to the right or to the left of the CI
print	Logical, should the plots also be printed? Only works when one matrix is used in 'matrices'
major_break	Numeric indicating the step size between major breaks
minor_break	Numeric indicating the step size between minor breaks
split0	Logical only used for results of aggregate_bootstraps. When set to TRUE, the displayed intervals are based on occasions when the parameter was not estimated to be zero, and an extra box is added indicating the number of times a parameter is estimated to be zero. Defaults to TRUE when model selection is used and FALSE otherwise.
prop0	Logical only used for results of aggregate_bootstraps, should boxes indicating the proportion of times parameters were estimated to be zero be added to the plot? Defaults to the value of split0.
prop0_cex	Only used for results of aggregate_bootstraps. Size of the boxes indicating number of times a parameter was set to zero.
prop0_alpha	Only used for results of aggregate_bootstraps. Transparency of the boxes indicating number of times a parameter was set to zero.
prop0_minAlpha	Only used for results of aggregate_bootstraps. Minimal transparency of the *lines* of plotted intervals as the proportion of times an edge was not included goes to 0.

Value

A single ggplot2 object, or a list of ggplot2 objects for each matrix requested.

Author(s)

Sacha Epskamp

Examples

```
### Example from ?ggm ###
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit an empty GGM:
mod0 <- ggm(ConsData, vars = vars)

# Run the model:
mod0 <- mod0 %>% runmodel

# Labels:
labels <- c(
  "indifferent to the feelings of others",
  "inquire about others' well-being",
  "comfort others",
  "love children",
  "make people feel at ease")

# Plot the CIs:
CIplot(mod0, "omega", labels = labels, labelstart = 0.2)

### Example from ?gvar ###
library("dplyr")
library("graphicalVAR")

beta <- matrix(c(
  0,0.5,
  0.5,0
),2,2,byrow=TRUE)
kappa <- diag(2)
simData <- graphicalVARsim(50, beta, kappa)
```

```
# Form model:
model <- gvar(simData)

# Evaluate model:
model <- model %>% runmodel

# Plot the CIs:
CIplot(model, "beta")
```

compare

Model comparison

Description

This function will print a table comparing multiple models on chi-square, AIC and BIC.

Usage

```
compare(...)
```

```
## S3 method for class 'psychometrics_compare'
print(x, ...)
```

Arguments

...	Any number of psychometrics models. Can be named to change the rownames of the output.
x	Output of the compare function.

Value

A data frame with chi-square values, degrees of freedoms, RMSEAs, AICs, and BICs.

Author(s)

Sacha Epskamp

`covML`*Maximum likelihood covariance estimate*

Description

These functions complement the base R `cov` function by simplifying obtaining maximum likelihood (ML) covariance estimates (denominator n) instead of unbiased (UB) covariance estimates (denominator $n-1$). The function `covML` can be used to obtain ML estimates, the function `covUBtoML` transforms from UB to ML estimates, and the function `covMLtoUB` transforms from ML to UB estimates.

Usage

```
covML(x, ...)  
covUBtoML(x, n, ...)  
covMLtoUB(x, n, ...)
```

Arguments

<code>x</code>	A dataset
<code>n</code>	The sample size
<code>...</code>	Arguments sent to the <code>cov</code> function.

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

Examples

```
data("StarWars")  
Y <- StarWars[,1:10]  
  
# Unbiased estimate:  
UB <- cov(Y)  
  
# ML Estimate:  
ML <- covML(Y)  
  
# Check:  
all(abs(UB - covMLtoUB(ML, nrow(Y))) < sqrt(.Machine$double.eps))  
all(abs(ML - covUBtoML(UB, nrow(Y))) < sqrt(.Machine$double.eps))
```

diagnostics

Diagnostic functions

Description

The 'checkJacobian' function can be used to check if the analytic gradient / Jacobian is aligned with the numerically approximated gradient / Jacobian, and the 'checkFisher' function can be used to check if the analytic Hessian is aligned with the numerically approximated Hessian.

Usage

```
checkJacobian(x, f = "default", jac = "default", transpose = FALSE,  
             plot = TRUE, perturbStart = FALSE, method = "Richardson")
```

```
checkFisher(x, f = "default", fis = "default", transpose = FALSE,  
           plot = TRUE, perturbStart = FALSE)
```

Arguments

x	A 'psychometrics' object
f	A custom fit function or the psychometrics default fit function (default).
jac	A custom Jacobian function or the psychometrics default Jacobian function (default).
fis	A custom Fischer information function or the psychometrics default Fischer information function (default).
transpose	Should the numeric Jacobian be transposed?
plot	Should a diagnostic plot be produced?
perturbStart	Should start values be perturbed (only used in development)
method	Numeric derivative method (default: Richardson)

Author(s)

Sacha Epskamp

dlvm1	<i>Lag-1 dynamic latent variable model family of psychometrics models for panel data</i>
-------	--

Description

This is the family of models that models a dynamic factor model on panel data. The function accepts both wide-format data (with a design matrix for vars) and long-format data (with a character vector for vars plus idvar and optionally beepvar). The format is auto-detected from the type of vars. There are four covariance structures that can be modeled in different ways: within_latent, between_latent for the within-person and between-person latent (contemporaneous) models respectively, and within_residual, between_residual for the within-person and between-person residual models respectively. The panelgvar wrapper function sets the lambda to an identity matrix, all residual variances to zero, and models within-person and between-person latent (contemporaneous) models as GGMs. The panelvar wrapper does the same but models contemporaneous relations as a variance-covariance matrix. Finally, the panellvvar wrapper automatically models all latent networks as GGMs. The old name panel_lvvar is deprecated in favor of panellvvar.

Usage

```
dlvm1(data, vars, datatype = c("auto", "wide", "long"),
      idvar, beepvar,
      standardize = c("none", "z", "quantile", "z_per_wave"),
      lambda, within_latent = c("cov", "chol",
                                "prec", "ggm"), within_residual = c("cov", "chol",
                                                                    "prec", "ggm"), between_latent = c("cov", "chol",
                                                                                                        "prec", "ggm"), between_residual = c("cov", "chol",
                                                                                                                            "prec", "ggm"), beta = "full", omega_zeta_within =
      "full", delta_zeta_within = "diag", kappa_zeta_within
      = "full", sigma_zeta_within = "full",
      lowertri_zeta_within = "full", omega_epsilon_within =
      "zero", delta_epsilon_within = "diag",
      kappa_epsilon_within = "diag", sigma_epsilon_within =
      "diag", lowertri_epsilon_within = "diag",
      omega_zeta_between = "full", delta_zeta_between =
      "diag", kappa_zeta_between = "full",
      sigma_zeta_between = "full", lowertri_zeta_between =
      "full", omega_epsilon_between = "zero",
      delta_epsilon_between = "diag", kappa_epsilon_between
      = "diag", sigma_epsilon_between = "diag",
      lowertri_epsilon_between = "diag", nu, mu_eta,
      identify = TRUE, identification = c("loadings",
                                          "variance"), latents, groups, groupvar, covs,
      cors, means, nobs, corinput, start
      = "version2", covtype = c("choose", "ML", "UB"),
      missing = "auto", equal = "none",
      baseline_saturated = TRUE, estimator = "ML",
```

```

optimizer, storedata = FALSE, verbose = FALSE,
sampleStats, baseline =
c("stationary_random_intercept", "stationary",
"independence", "none"), bootstrap = FALSE, boot_sub,
boot_resample, within, between,
penalty_lambda = NA, penalty_alpha = 1,
penalize_matrices)

panelgvar(data, vars, within_latent = c("ggm","chol","cov","prec"),
between_latent = c("ggm","chol","cov","prec"), ...)

panelvar(data, vars, within_latent = c("cov","chol","prec","ggm"),
between_latent = c("cov","chol","prec","ggm"), ...)

panellvgvar(...)

panel_lvgvar(...)

```

Arguments

data	A data frame encoding the data used in the analysis. Can be missing if covs and nobs are supplied.
vars	Required argument. For wide-format data, this must be a <i>*matrix*</i> with each row indicating a variable and each column indicating a measurement. The matrix must be filled with names of the variables in the dataset corresponding to variable <i>i</i> at wave <i>j</i> . NAs can be used to indicate missing waves. The rownames of this matrix will be used as variable names. For long-format data, this should be a character vector of variable names in the dataset.
datatype	Data format: "auto" (default) auto-detects based on whether vars is a matrix (wide) or character vector (long). "wide" forces wide-format interpretation. "long" forces long-format interpretation with automatic reshape to wide.
idvar	Optional string indicating the subject/cluster ID variable in data. Required when datatype = "long" or when vars is a character vector.
beepvar	Optional string indicating the time point / measurement occasion variable. If missing when datatype = "long", a sequential measurement variable is created per subject.
standardize	Standardization method: "none" (default), "z" for global z-scores per variable across all waves (uses the overall mean and standard deviation for each variable computed across all waves of data), "quantile" for quantile normalization across all waves, "z_per_wave" for independent z-scores per wave column (note: this imposes stationarity as means become 0 and variances become 1 at each wave).
lambda	Required argument. A model matrix encoding the factor loading structure. Each row indicates an indicator and each column a latent. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

<code>within_latent</code>	The type of within-person latent contemporaneous model to be used.
<code>within_residual</code>	The type of within-person residual model to be used.
<code>between_latent</code>	The type of between-person latent model to be used.
<code>between_residual</code>	The type of between-person residual model to be used.
<code>beta</code>	A model matrix encoding the temporal relationships (transpose of temporal network). A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix. Can also be "full" for a full temporal network or "zero" for an empty temporal network.
<code>omega_zeta_within</code>	Only used when <code>within_latent = "ggm"</code> . Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>delta_zeta_within</code>	Only used when <code>within_latent = "ggm"</code> . Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>kappa_zeta_within</code>	Only used when <code>within_latent = "prec"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>sigma_zeta_within</code>	Only used when <code>within_latent = "cov"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>lowertri_zeta_within</code>	Only used when <code>within_latent = "chol"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>omega_epsilon_within</code>	Only used when <code>within_residual = "ggm"</code> . Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`delta_epsilon_within`

Only used when `within_residual = "ggm"`. Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`kappa_epsilon_within`

Only used when `within_residual = "prec"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`sigma_epsilon_within`

Only used when `within_residual = "cov"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`lowertri_epsilon_within`

Only used when `within_residual = "chol"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`omega_zeta_between`

Only used when `between_latent = "ggm"`. Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`delta_zeta_between`

Only used when `between_latent = "ggm"`. Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`kappa_zeta_between`

Only used when `between_latent = "prec"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`sigma_zeta_between`

Only used when `between_latent = "cov"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

lowertri_zeta_between	Only used when <code>between_latent = "chol"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega_epsilon_between	Only used when <code>between_residual = "ggm"</code> . Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_epsilon_between	Only used when <code>between_residual = "ggm"</code> . Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_epsilon_between	Only used when <code>between_residual = "prec"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
sigma_epsilon_between	Only used when <code>between_residual = "cov"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_epsilon_between	Only used when <code>between_residual = "chol"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
nu	Optional vector encoding the intercepts of the observed variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
mu_eta	Optional vector encoding the means of the latent variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
identify	Logical, should the model be automatically identified?
identification	Type of identification used. "loadings" to fix the first factor loadings to 1, and "variance" to fix the diagonal of the latent variable model matrix (<code>sigma_zeta</code> , <code>lowertri_zeta</code> , <code>delta_zeta</code> or <code>kappa_zeta</code>) to 1.

latents	An optional character vector with names of the latent variables.
groups	Deprecated. Use groupvar instead. An optional string indicating the name of the group variable in data.
groupvar	An optional string indicating the name of the group variable in data. Replaces the deprecated groups argument; if both are supplied, groupvar takes precedence with a warning.
covs	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. IMPORTANT NOTE: psychometrics expects the maximum likelihood (ML) covariance matrix, which is NOT obtained from cov directly. Manually rescale the result of cov with $(nobs - 1)/nobs$ to obtain the ML covariance matrix.
cors	A sample correlation matrix, or a list/array of such matrices for multiple groups. When supplied, the matrix is treated as a covariance matrix with a warning (appropriate for standardized data). Requires nobs. Note: corinput = TRUE is not supported in d1vm1.
means	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
nobs	The number of observations used in covs and means, or a vector of such numbers of observations for multiple groups.
corinput	Logical. Not supported in d1vm1() and will produce an error if set to TRUE.
start	Start value specification. Can be either a string or a psychometrics model. If it is a string, "version2" indicates the latest version of start value computation, "version1" indicates start values as they were computed up to version 0.11, and "simple" indicate simple starting values. If this is a psychometrics model the starting values will be based on the output. This can be useful, for example, if you first estimate a model with matrices set to a Cholesky decomposition, then use those values as start values for estimating Gaussian graphical models.
missing	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be "auto" (default) for automatic detection, "listwise" for listwise deletion, or "pairwise" for pairwise deletion. When "auto", the function checks for missing data and switches ML to FIML, PML to PFIML, or defaults to listwise for LS estimators.
equal	A character vector indicating which matrices should be constrained equal across groups.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
estimator	The estimator to be used. Currently implemented are "ML" for maximum likelihood estimation, "FIML" for full-information maximum likelihood estimation, "PML" for penalized maximum likelihood estimation, "PFIML" for penalized full-information maximum likelihood estimation, "ULS" for unweighted least squares estimation, "WLS" for weighted least squares estimation, and "DWLS" for diagonally weighted least squares estimation. When missing = "auto" (default), "ML" is automatically switched to "FIML" and "PML" to "PFIML" if missing data is detected.

optimizer	The optimizer to be used. Can be one of "nlminb" (the default R nlminb function), "ucminf" (from the optimr package), "nloptr_TNEWTON" (preconditioned truncated Newton via nloptr), and "LBFGS++" (pure C++ L-BFGS-B). Defaults to "nlminb".
storedata	Logical, should the raw data be stored? Needed for bootstrapping (see bootstrap).
verbose	Logical, should progress be printed to the console?
sampleStats	An optional sample statistics object. Mostly used internally.
covtype	If 'covs' is used, this is the type of covariance (maximum likelihood or unbiased) the input covariance matrix represents. Set to "ML" for maximum likelihood estimates (denominator n) and "UB" to unbiased estimates (denominator n-1). The default will try to find the type used, by investigating which is most likely to result from integer valued datasets.
baseline	What baseline model should be used? "stationary_random_intercept" includes both within- and between person variances constrained equal across time (default), "stationary" only includes within-person variances constrained equal across time, "independence" (default up to version 0.11) includes a variance for every variable at every time point (not constrained equal across time), and "none" includes no baseline model.
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets boot_sub = 1 and boot_resample = TRUE) or "case" (which sets boot_sub = 0.75 and boot_resample = FALSE).
boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
within	Optional alias for within_latent. If both within and within_latent are explicitly specified, within_latent takes precedence and a warning is issued.
between	Optional alias for between_latent. If both between and between_latent are explicitly specified, between_latent takes precedence and a warning is issued.
penalty_lambda	Numeric penalty strength for penalized ML estimation (PML/PFIML). NA (default) triggers automatic selection via EBIC-based grid search when a penalized estimator is used; set to a specific numeric value to use a fixed penalty strength (0 = no penalty). See find_penalized_lambda and penalize .
penalty_alpha	Elastic net mixing parameter: 1 = LASSO (default), 0 = ridge.
penalize_matrices	Character vector of matrix names to penalize. If missing, defaults are selected based on the model type.
...	Arguments sent to dlvm1.

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

Examples

```
library("dplyr")

# Smoke data cov matrix, based on LISS data panel https://www.dataarchive.lisssdata.nl
smoke <- structure(c(47.2361758611759, 43.5366809116809, 41.0057465682466,
                    43.5366809116809, 57.9789886039886, 47.6992521367521,
                    41.0057465682466,
                    47.6992521367521, 53.0669434731935), .Dim = c(3L, 3L),
                  .Dimnames = list(
                    c("smoke2008", "smoke2009", "smoke2010"), c("smoke2008",
                    "smoke2009", "smoke2010")))

# Design matrix:
design <- matrix(rownames(smoke),1,3)

# Form model:
mod <- panelvar(vars = design,
               covs = smoke, nobs = 352
               )

# Run model:
mod <- mod %>% runmodel

# Evaluate fit:
mod %>% fit
```

duplicationMatrix *Model matrices used in derivatives*

Description

These matrices are used in the analytic gradients

Usage

```
duplicationMatrix(n, diag = TRUE)
```

```
eliminationMatrix(n, diag = TRUE)
```

```
diagonalizationMatrix(n)
```

Arguments

n Number of rows and columns in the original matrix
diag Logical indicating if the diagonal should be included (set to FALSE for derivative of `vech(x)`)

Value

A sparse matrix

Author(s)

Sacha Epskamp

Examples

```
# Duplication matrix for 10 variables:  
duplicationMatrix(10)  
  
# Elimination matrix for 10 variables:  
eliminationMatrix(10)  
  
# Diagonalization matrix for 10 variables:  
diagonalizationMatrix(10)
```

emergencystart

Reset starting values to simple defaults

Description

This function overwrites the starting values to simple defaults. This can help in cases where optimization fails.

Usage

```
emergencystart(x)
```

Arguments

x A psychometrics model.

Value

A psychometrics model.

Author(s)

Sacha Epskamp

Description

These functions implement Ergodic Subspace Analysis by von Oertzen, Schmiedek and Voelkle (2020). The functions can be used on the output of a [dlvm1](#) model, or manually by supplying a within persons and between persons variance-covariance matrix.

Usage

```
esa(x, cutoff = 0.1,
    between = c("crosssection", "between"))
esa_manual(sigma_wp, sigma_bp, cutoff = 0.1)
## S3 method for class 'esa'
print(x, printref = TRUE, ...)
## S3 method for class 'esa_manual'
print(x, printref = TRUE, ...)
## S3 method for class 'esa'
plot(x, plot = c("observed", "latent"), ...)
## S3 method for class 'esa_manual'
plot(x, ...)
```

Arguments

x	Output of a dlvm1 model
sigma_wp	Manual within-person variance-covariance matrix
sigma_bp	Manual between-person variance-covariance matrix
cutoff	Cutoff used to determine ergodicity
printref	Logical, should the reference be printed?
plot	Should ergodicity of observed or latent variables be plotted?
between	Should the between-persons variance-covariance matrix be based on exected cross-sectional or between-person relations
...	Not used

Value

For each group a `esa_manual` object with the following elements:

ergodicity	Ergodicity values of each component
Q_esa	Component loadings
V_bp	Between persons subspace
V_ergodic	Ergodic subspace
V_wp	Within person subspace
cutoff	Cutoff value used

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

References

von Oertzen, T., Schmiedek, F., and Voelkle, M. C. (2020). Ergodic Subspace Analysis. *Journal of Intelligence*, 8(1), 3.

factorscores	<i>Compute factor scores</i>
--------------	------------------------------

Description

Currently, only the lvm framework with single group and no missing data is supported.

Usage

```
factorscores(data, model, method = c("bartlett", "regression"))
```

Arguments

data	Dataset to compute factor scores for
model	A psychometrics model
method	The method to use: "regression" or "bartlett"

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

find_penalized_lambda	<i>Automated Lambda Selection for Penalized Estimation</i>
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Description

Selects the optimal penalty strength (λ) for penalized maximum likelihood (PML) or penalized full-information maximum likelihood (PFIML) models via EBIC-based grid search. A log-spaced grid of λ values is searched from λ_{\max} (the smallest λ for which all penalized parameters are exactly zero, derived from KKT conditions) down to $\lambda_{\max} * \lambda_{\min_ratio}$. Each λ is optimized using ISTA (Iterative Shrinkage-Thresholding Algorithm) with warm starts from the previous solution, and the model with the best EBIC is selected. After selection, a β_{\min} threshold is applied to set near-zero penalized parameters exactly to zero.

This function is called automatically by `runmodel` when the model contains auto-select penalty parameters (i.e., `penalty_lambda = NA` in the parameter table, which is the default when using `estimator = "PML"` or `estimator = "PFIML"`). It can also be called directly for more control over the search.

Usage

```
find_penalized_lambda(model, criterion = "ebic.5", nLambda = 50,
                      lambda_min_ratio = 1e-4,
                      beta_min = c("numerical", "theoretical"),
                      verbose)
```

Arguments

model	A psychometrics model with estimator = "PML" or estimator = "PFIML". The model should contain parameters with penalty_lambda = NA (auto-select), which is the default when specifying estimator = "PML" or "PFIML" in model constructors such as ggm , lvm , etc.
criterion	Character string specifying the information criterion used to select the best lambda. One of: "bic" BIC (equivalent to EBIC with gamma = 0) "ebic.25" EBIC with gamma = 0.25 "ebic.5" EBIC with gamma = 0.5 (default; recommended for network models) "ebic.75" EBIC with gamma = 0.75 "ebic1" EBIC with gamma = 1.0 Higher gamma values penalize model complexity more strongly and tend to produce sparser solutions. Gamma = 0.5 is widely recommended for network models (Foygel & Drton, 2010).
nLambda	Integer, the number of lambda values in the search grid. Default is 50. The grid is log-spaced between lambda_max and lambda_max * lambda_min_ratio.
lambda_min_ratio	Numeric, the ratio of the smallest to the largest lambda in the grid. Default is 1e-4, so that lambda_min = lambda_max * 1e-4.
beta_min	Threshold for zeroing out small penalized parameter estimates. Parameters with absolute value below beta_min are set to zero, both during EBIC evaluation (for parameter counting) and in the final model. Can be: "numerical" Uses a fixed threshold of 1e-05 (default). "theoretical" Uses $\sqrt{\log(p) / n}$ where p is the number of penalized parameters and n is the total sample size. numeric value Uses the provided value directly as the threshold.
verbose	Logical, should progress messages be printed? Defaults to the model's verbose setting.

Details

The search proceeds as follows:

1. **Compute lambda_max:** The analytical upper bound is derived from KKT (Karush-Kuhn-Tucker) conditions at the null-penalized model (all penalized parameters fixed at zero). Specifically, $\lambda_{\max} = \max(|\text{grad}_j|) / \alpha$ where grad_j are the gradient elements for penalized parameters at the null solution and α is the elastic net mixing parameter.

2. **Build lambda grid:** A log-spaced sequence of nLambda values from lambda_max down to lambda_max * lambda_min_ratio.
3. **Grid search with warm starts:** For each lambda (from largest to smallest), the model is optimized using ISTA. The solution from the previous lambda serves as the starting point (warm start), which substantially reduces computation time. The EBIC (Extended Bayesian Information Criterion) is computed using the unpenalized log-likelihood (Convention A), counting only parameters with |estimate| >= beta_min as non-zero.
4. **Beta-min thresholding:** Penalized parameters with |estimate| < beta_min are set exactly to zero in the final model.

The EBIC is computed as:

$$EBIC = -2 \cdot LL + k \cdot \log(n) + 4 \cdot k \cdot \gamma \cdot \log(p_v)$$

where LL is the unpenalized log-likelihood, k is the effective number of parameters (unpenalized free parameters plus non-zero penalized parameters), n is the sample size, γ is the EBIC tuning parameter, and p_v is the number of observed variables.

The returned model stores metadata about the search in `model@optim$lambda_search`, a list containing: `criterion`, `gamma`, `best_lambda`, `best_criterion`, `beta_min`, `lambda_max`, `nLambda`, and `n_thresholded`.

Value

An object of class `psychometrics` ([psychometrics-class](#)) with:

- The selected lambda assigned to all auto-select penalty parameters
- Parameters optimized at the best lambda
- Near-zero penalized parameters set to exactly zero (via `beta_min`)
- Search metadata stored in `model@optim$lambda_search`

After running `find_penalized_lambda`, use `refit` to obtain valid standard errors and fit indices via post-selection inference.

Author(s)

Sacha Epskamp

References

Foygel, R., & Drton, M. (2010). Extended Bayesian Information Criteria for Gaussian Graphical Models. In *Advances in Neural Information Processing Systems* (Vol. 23).

See Also

`penalize` for manually setting penalty parameters, `runmodel` for running models (calls `find_penalized_lambda` automatically when auto-select penalties are present), `refit` for post-selection inference after penalized estimation.

Examples

```
# Load bfi data from psych package:
library("psychTools")
library("dplyr")
data(bfi)

ConsData <- bfi %>%
  select(A1:A5) %>%
  na.omit

vars <- names(ConsData)

# Automatic lambda selection (called implicitly by runmodel):
mod <- ggm(ConsData, vars = vars, estimator = "PML")
mod <- mod %>% runmodel # automatically searches for best lambda

# Check lambda search results:
mod@optim$lambda_search

# Post-selection refit for standard errors:
mod_refit <- mod %>% refit
mod_refit %>% parameters

# Direct call with custom criterion:
mod2 <- ggm(ConsData, vars = vars, estimator = "PML")
mod2 <- find_penalized_lambda(mod2, criterion = "ebic1",
                             nLambda = 100)
```

fit

Print fit indices

Description

This function will print all fit indices of the model/

Usage

```
fit(x)
```

Arguments

x A psychometrics model.

Value

Invisibly returns a data frame with fit measure estimates.

Author(s)

Sacha Epskamp

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit an empty GGM:
mod0 <- ggm(ConsData, vars = vars, omega = "zero")

# Run model:
mod0 <- mod0 %>% runmodel

# Inspect fit:
mod0 %>% fit # Pretty bad fit...
```

fixpar

Parameters modification

Description

The `fixpar` function can be used to fix a parameter to some value (Typically zero), and the `freepar` function can be used to free a parameter from being fixed to a value.

Usage

```
fixpar(x, matrix, row, col, value = 0, group, verbose,
       log = TRUE, runmodel = FALSE, ...)
```

```
freepar(x, matrix, row, col, start, group, verbose, log =
       TRUE, runmodel = FALSE, startEPC = TRUE, ...)
```

Arguments

x	A psychometrics model.
matrix	String indicating the matrix of the parameter
row	Integer or string indicating the row of the matrix of the parameter
col	Integer or string indicating the column of the matrix of the parameter
value	Used in fixpar to indicate the value to which a parameters is constrained
start	Used in freepar to indicate the starting value of the parameter
group	Integer indicating the group of the parameter to be constrained
verbose	Logical, should messages be printed?
log	Logical, should the log be updated?
runmodel	Logical, should the model be updated?
startEPC	Logical, should the starting value be set at the expected parameter change?
...	Arguments sent to runmodel

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

 fixstart

Attempt to Fix Starting Values

Description

This function attempts to fix starting values by comparing the analytic gradient to a numerically approximated gradient. Parameters with a difference between the analytic and numeric gradient that exceeds 'maxdiff' will be reduced by a factor of 'reduce' in each iteration until the average absolute difference between analytic and numeric gradients is lower than 'tol'. Only off-diagonal elements in omega, sigma, kappa, lowertri or rho matrices or any element in beta matrices are adjusted.

Usage

```
fixstart(x, reduce = 0.5, maxdiff = 0.1, tol = 0.01, maxtry = 25)
```

Arguments

x	A 'psychonetrics' model
reduce	The factor with which problematic parameters are reduced in each iteration.
maxdiff	Maximum difference between analytic and numeric gradient to be considered problematic.
tol	Average absolute difference between analytic and numeric gradient that is considered acceptable.
maxtry	Maximum number of iterations to attempt to fix starting values.

Author(s)

Sacha Epskamp

generate

Generate data from a fitted psychonetrics object

Description

This function will generate new data from the estimated mean and variance-covariance structure of a psychonetrics model.

Usage

```
generate(x, n = 500)
```

Arguments

x	A psychonetrics model.
n	Number of cases to sample per group.

Value

A data frame with simulated data

Author(s)

Sacha Epskamp

getmatrix *Extract an estimated matrix*

Description

This function will extract an estimated matrix, and will either return a single matrix for single group models or a list of such matrices for multiple group models.

Usage

```
getmatrix(x, matrix, group, threshold = FALSE, alpha = 0.01,
          adjust = c("none", "holm", "hochberg", "hommel",
                    "bonferroni", "BH", "BY", "fdr"), mode = c("tested",
                    "all"), diag = TRUE)
```

Arguments

x	A psychometrics model.
matrix	String indicating the matrix to be extracted.
group	Integer indicating the group for the matrix to be extracted.
threshold	Logical. Should the matrix be thresholded (non-significant values set to zero? Can also be a value with an absolute threshold below which parameters are set to zero.)
alpha	Significance level to use.
adjust	p-value adjustment method to use. See p.adjust.
mode	Mode for adjusting for multiple comparisons. Should all parameters be considered as the total number of tests or only the tested parameters (parameters of interest)?
diag	Set to FALSE to set diagonal elements to zero.

Value

A matrix of parameter estimates, of a list of such matrices for multiple group models.

Author(s)

Sacha Epskamp

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")
```

```
# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars, omega = "full")

# Run model:
mod <- mod %>% runmodel

# Obtain network:
mod %>% getmatrix("omega")
```

getVCOV

Obtain the asymptotic covariance matrix

Description

This function can be used to obtain the estimated asymptotic covariance matrix from a psychometrics object.

Usage

```
getVCOV(model, approximate_SEs = FALSE)
```

Arguments

`model` A psychometrics model.

`approximate_SEs` Logical, should standard errors be approximated? If true, an approximate matrix inverse of the Fischer information is used to obtain the standard errors.

Value

This function returns a matrix.

Author(s)

Sacha Epskamp

groupequal	<i>Group equality constrains</i>
------------	----------------------------------

Description

The `groupequal` function constrains parameters equal across groups, and the `groupfree` function frees equality constrains across groups.

Usage

```
groupequal(x, matrix, row, col, verbose, log = TRUE, runmodel =  
          FALSE, identify = TRUE, ...)
```

```
groupfree(x, matrix, row, col, verbose, log = TRUE, runmodel =  
          FALSE, identify = TRUE, ...)
```

Arguments

<code>x</code>	A psychometrics model.
<code>matrix</code>	String indicating the matrix of the parameter
<code>row</code>	Integer or string indicating the row of the matrix of the parameter
<code>col</code>	Integer or string indicating the column of the matrix of the parameter
<code>verbose</code>	Logical, should messages be printed?
<code>log</code>	Logical, should the log be updated?
<code>runmodel</code>	Logical, should the model be updated?
<code>identify</code>	Logical, should the model be identified?
<code>...</code>	Arguments sent to <code>runmodel</code>

Value

An object of the class `psychometrics` ([psychometrics-class](#))

Author(s)

Sacha Epskamp

Ising

*Ising model***Description**

This is the family of Ising models fit to dichotomous datasets. Note that the input matters (see also <https://arxiv.org/abs/1811.02916>) in this model! Models based on a dataset that is encoded with -1 and 1 are not entirely equivalent to models based on datasets encoded with 0 and 1 (non-equivalences occur in multi-group settings with equality constrains).

Usage

```
Ising(data, omega = "full", tau, beta, beta_model =
      c("beta", "log_beta"), vars, groups, covs, means,
      nob, covtype = c("choose", "ML", "UB"), responses,
      missing = "listwise", equal = "none",
      baseline_saturated = TRUE, estimator = "default",
      optimizer, storedata = FALSE, WLS.W, sampleStats,
      identify = TRUE, verbose = FALSE, maxNodes = 20,
      min_sum = -Inf, bootstrap = FALSE, boot_sub,
      boot_resample, penalty_lambda = NA,
      penalty_alpha = 1, penalize_matrices)
```

Arguments

data	A data frame encoding the data used in the analysis. Can be missing if covs and nob are supplied.
omega	The network structure. Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions nNode x nNode with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
tau	Optional vector encoding the threshold/intercept structure. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
beta	Optional scalar encoding the inverse temperature. 1 indicate free beta parameters, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such scalars.
beta_model	How should beta be modeled? Set beta_model = "log_beta" to model the log of beta rather than beta directly.
vars	An optional character vector encoding the variables used in the analysis. Must equal names of the dataset in data.
groups	An optional character vector encoding the variables used in the analysis. Must equal names of the dataset in data.

<code>covs</code>	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. Make sure <code>covtype</code> argument is set correctly to the type of covariances used.
<code>means</code>	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
<code>nobs</code>	The number of observations used in <code>covs</code> and <code>means</code> , or a vector of such numbers of observations for multiple groups.
<code>covtype</code>	If <code>'covs'</code> is used, this is the type of covariance (maximum likelihood or unbiased) the input covariance matrix represents. Set to <code>"ML"</code> for maximum likelihood estimates (denominator <code>n</code>) and <code>"UB"</code> to unbiased estimates (denominator <code>n-1</code>). The default will try to find the type used, by investigating which is most likely to result from integer valued datasets.
<code>responses</code>	A vector of dichotomous responses used (e.g., <code>c(-1, 1)</code> or <code>c(0, 1)</code>). Only needed when <code>'covs'</code> is used.)
<code>missing</code>	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be <code>"listwise"</code> for listwise deletion, or <code>"pairwise"</code> for pairwise deletion. NOT RECOMMENDED TO BE USED YET IN ISING MODEL.
<code>equal</code>	A character vector indicating which matrices should be constrained equal across groups.
<code>baseline_saturated</code>	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
<code>estimator</code>	The estimator to be used. Currently implemented are <code>"ML"</code> for maximum likelihood estimation, <code>"FIML"</code> for full-information maximum likelihood estimation, <code>"ULS"</code> for unweighted least squares estimation, <code>"WLS"</code> for weighted least squares estimation, and <code>"DWLS"</code> for diagonally weighted least squares estimation. Only ML estimation is currently supported for the Ising model.
<code>optimizer</code>	The optimizer to be used. Can be one of <code>"nlminb"</code> (the default R <code>nlminb</code> function), <code>"ucminf"</code> (from the <code>optimr</code> package), <code>"nloptr_TNEWTON"</code> (preconditioned truncated Newton via <code>nloptr</code>), and <code>"LBFGS++"</code> (pure C++ L-BFGS-B). Defaults to <code>"nlminb"</code> .
<code>storedata</code>	Logical, should the raw data be stored? Needed for bootstrapping (see <code>bootstrap</code>).
<code>WLS.W</code>	Optional WLS weights matrix. CURRENTLY NOT USED.
<code>sampleStats</code>	An optional sample statistics object. Mostly used internally.
<code>identify</code>	Logical, should the model be identified?
<code>verbose</code>	Logical, should messages be printed?
<code>maxNodes</code>	The maximum number of nodes allowed in the analysis. This function will stop with an error if more nodes are used (it is not recommended to set this higher).
<code>min_sum</code>	The minimum sum score that is artificially possible in the dataset. Defaults to <code>-Inf</code> . Set this only if you know a lower sum score is not possible in the data, for example due to selection bias.

<code>bootstrap</code>	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using <code>aggregate_bootstraps!</code> Can be TRUE or FALSE. Can also be "nonparametric" (which sets <code>boot_sub = 1</code> and <code>boot_resample = TRUE</code>) or "case" (which sets <code>boot_sub = 0.75</code> and <code>boot_resample = FALSE</code>).
<code>boot_sub</code>	Proportion of cases to be subsampled (<code>round(boot_sub * N)</code>).
<code>boot_resample</code>	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
<code>penalty_lambda</code>	Numeric penalty strength for penalized ML estimation (PML/PFIML). NA (default) triggers automatic selection via EBIC-based grid search when a penalized estimator is used; set to a specific numeric value to use a fixed penalty strength (0 = no penalty). See <code>find_penalized_lambda</code> and <code>penalize</code> .
<code>penalty_alpha</code>	Elastic net mixing parameter: 1 = LASSO (default), 0 = ridge.
<code>penalize_matrices</code>	Character vector of matrix names to penalize. If missing, defaults are selected based on the model type.

Details

The Ising Model takes the following form:

$$\Pr(\mathbf{Y} = \mathbf{y}) = \frac{\exp(-\beta H(\mathbf{y}; \boldsymbol{\tau}, \boldsymbol{\Omega}))}{Z(\boldsymbol{\tau}, \boldsymbol{\Omega})}$$

With Hamiltonian:

$$H(\mathbf{y}; \boldsymbol{\tau}, \boldsymbol{\Omega}) = -\sum_{i=1}^m \tau_i y_i - \sum_{i=2}^m \sum_{j=1}^{i-1} \omega_{ij} y_i y_j.$$

And Z representing the partition function or normalizing constant.

Value

An object of the class `psychometrics`

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

References

Epskamp, S., Maris, G., Waldorp, L. J., & Borsboom, D. (2018). Network Psychometrics. In: Irving, P., Hughes, D., & Booth, T. (Eds.), *The Wiley Handbook of Psychometric Testing, 2 Volume Set: A Multidisciplinary Reference on Survey, Scale and Test Development*. New York: Wiley.

Examples

```
library("dplyr")
data("Jonas")

# Variables to use:
vars <- names(Jonas)[1:10]
```

```

# Arranged groups to put unfamiliar group first (beta constrained to 1):
Jonas <- Jonas[order(Jonas$group),]

# Form saturated model:
model1 <- Ising(Jonas, vars = vars, groups = "group")

# Run model:
model1 <- model1 %>% runmodel
# Note: SEs are approximated automatically because there are zeroes
# in the crosstables of the "Knows Jonas" group, which makes the
# Fisher information matrix singular. A warning is shown when
# this fallback occurs.

# Prune-stepup to find a sparse model:
model1b <- model1 %>% prune(alpha = 0.05) %>% stepup(alpha = 0.05)

# Equal networks:
model2 <- model1 %>% groupequal("omega") %>% runmodel

# Prune-stepup to find a sparse model:
model2b <- model2 %>% prune(alpha = 0.05) %>% stepup(mi = "mi_equal", alpha = 0.05)

# Equal thresholds:
model3 <- model2 %>% groupequal("tau") %>% runmodel

# Prune-stepup to find a sparse model:
model3b <- model3 %>% prune(alpha = 0.05) %>% stepup(mi = "mi_equal", alpha = 0.05)

# Equal beta:
model4 <- model3 %>% groupequal("beta") %>% runmodel

# Prune-stepup to find a sparse model:
model4b <- model4 %>% prune(alpha = 0.05) %>% stepup(mi = "mi_equal", alpha = 0.05)

# Compare all models:
compare(
  `1. all parameters free (dense)` = model1,
  `2. all parameters free (sparse)` = model1b,
  `3. equal networks (dense)` = model2,
  `4. equal networks (sparse)` = model2b,
  `5. equal networks and thresholds (dense)` = model3,
  `6. equal networks and thresholds (sparse)` = model3b,
  `7. all parameters equal (dense)` = model4,
  `8. all parameters equal (sparse)` = model4b
) %>% arrange(BIC)

```

Description

Responses of 10 attitude items towards a researcher named Jonas. Participants were shown three photos of Jonas with the text: "This is Jonas, a researcher from Germany who is now becoming a PhD in Psychology". Subsequently, the participants had to answer 10 yes / no questions starting with "I believe that Jonas...", as well as rate their familiarity with Jonas. The sample consists of people familiar with Jonas and not familiar with Jonas, and allows for testing Attitudinal Entropy Framework <doi:10.1080/1047840X.2018.1537246>.

Usage

```
data("Jonas")
```

Format

A data frame with 215 observations on the following 12 variables.

scientist ... is a good scientist

jeans ... Is a person that wears beautiful jeans

cares ... really cares about people like you

economics ... would solve our economic problems

hardworking ... is hardworking

honest ... is honest

intouch ... is in touch with ordinary people

knowledgeable ... is knowledgeable

makeupmind ... can't make up his mind

getsthingsdone ... gets things done

familiar Answers to the question "How familiar are you with Jonas?" (three responses possible)

group The question 'familiar' categorized in two groups ("Knows Jonas" and "Doesn't Know Jonas")

Examples

```
data(Jonas)
```

latentgrowth

Latnet growth curve model

Description

Wrapper to lvm to specify a latent growth curve model.

Usage

```
latentgrowth(vars, time = seq_len(ncol(vars)) - 1, covariates =
  character(0), covariates_as = c("regression",
  "covariance"), ...)
```

Arguments

vars	Different from in other psychometrics models, this must be a <i>*matrix*</i> with each row indicating a variable and each column indicating a measurement. The matrix must be filled with names of the variables in the dataset corresponding to variable <i>i</i> at wave <i>j</i> . NAs can be used to indicate missing waves. The rownames of this matrix will be used as variable names.
time	A vector with the encoding of each measurement (e.g., 0, 1, 2, 3).
covariates	A vector with strings indicating names of between-person covariate variables in the data
covariates_as	Should covariates be included as regressions or actual covariates?
...	Arguments sent to lvm

Details

See https://github.com/SachaEpskamp/SEM-code-examples/tree/master/Latent_growth_examples/psychometrics for examples

Value

An object of the class psychometrics ([psychometrics-class](#)). See for an example https://github.com/SachaEpskamp/SEM-code-examples/tree/master/Latent_growth_examples/psychometrics.

Author(s)

Sacha Epskamp

Examples

```
library("dplyr")

# Smoke data cov matrix, based on LISS data panel https://www.dataarchive.lisssdata.nl
smoke <- structure(c(47.2361758611759, 43.5366809116809, 41.0057465682466,
                    43.5366809116809, 57.9789886039886, 47.6992521367521,
                    41.0057465682466,
                    47.6992521367521, 53.0669434731935), .Dim = c(3L, 3L),
                  .Dimnames = list(
                    c("smoke2008", "smoke2009", "smoke2010"), c("smoke2008",
                    "smoke2009", "smoke2010")))

# Design matrix:
design <- matrix(rownames(smoke),1,3)

# Form model:
mod <- latentgrowth(vars = design,
                   covs = smoke, nobs = 352
)

## Not run:
# Run model:
```

```

mod <- mod %>% runmodel

# Evaluate fit:
mod %>% fit

# Look at parameters:
mod %>% parameters

## End(Not run)

```

logbook	<i>Retrieve the psychonetrics logbook</i>
---------	---

Description

This function can be used to retrieve the logbook of a 'psychonetrics' object.

Usage

```
logbook(x, log = TRUE)
```

Arguments

x	A 'psychonetrics' object.
log	Logical, should the entry that the logbook is accessed be added?

Author(s)

Sacha Epskamp

loop_psychonetrics	<i>Parallel loop for psychonetrics</i>
--------------------	--

Description

A convenience wrapper for running repeated psychonetrics analyses in parallel. Replaces the typical boilerplate of `makeCluster` / `clusterExport` / `parLapply` / `stopCluster` with a single function call. Particularly useful for bootstrapping, simulation studies, and cross-validation.

Variables referenced in the expression that exist in the calling environment are automatically exported to parallel workers, so explicit `clusterExport` calls are not needed in most cases.

Usage

```

loop_psychonetrics(expr, reps = 1000, nCores = 1,
  export = character(0),
  packages = c("psychonetrics", "dplyr"),
  verbose = TRUE, envir = parent.frame())

```

Arguments

<code>expr</code>	An expression (code block) to evaluate on each iteration. Typically enclosed in curly braces <code>{...}</code> . The expression should return a single object (e.g., a psychometrics model).
<code>reps</code>	Number of replications (default: 1000).
<code>nCores</code>	Number of CPU cores to use. Set to 1 for sequential execution (default), or higher for parallel execution via <code>makePSOCKcluster</code> .
<code>export</code>	Character vector of additional variable names to export to parallel workers. Usually not needed because variables referenced in <code>expr</code> are auto-detected and exported.
<code>packages</code>	Character vector of packages to load on parallel workers (default: <code>c("psychometrics", "dplyr")</code>).
<code>verbose</code>	Logical; if TRUE (default), shows a progress bar via <code>pbapply</code> .
<code>envir</code>	The environment in which to look up variables for auto-export and to evaluate the expression in sequential mode. Defaults to the calling environment.

Details

When `nCores > 1`, a PSOCK cluster is created and automatically cleaned up when the function exits (even on error). The specified packages are loaded on each worker, and variables referenced in `expr` are automatically detected and exported.

For bootstrapping, use this function together with `bootstrap = "nonparametric"` in the model constructor and [aggregate_bootstraps](#) to summarize the results.

Value

A list of length `reps`, where each element is the result of evaluating `expr` once.

See Also

[aggregate_bootstraps](#), [bootstrap](#), [CIplot](#)

Examples

```
library(dplyr)
data(NA2020)

# Fit the original model:
model <- ggm(NA2020, estimator = "FIML") %>% runmodel

# Bootstrap with pruning (parallel, 100 reps):
bootstraps <- loop_psychometrics({
  ggm(NA2020, bootstrap = "nonparametric",
    estimator = "FIML") %>%
    runmodel %>%
    prune(alpha = 0.05)
}, reps = 100, nCores = 2)
```

```

# Aggregate and inspect:
boot_agg <- aggregate_bootstraps(
  sample = model %>% prune(alpha = 0.05),
  bootstraps = bootstraps
)

# View results:
parameters(boot_agg)
CIplot(boot_agg, "omega", split0 = TRUE)

```

Ivm

Continuous latent variable family of psychometrics models

Description

This is the family of models that models the data as a structural equation model (SEM), allowing the latent and residual variance-covariance matrices to be further modeled as networks. The latent and residual arguments can be used to define what latent and residual models are used respectively: "cov" (default) models a variance-covariance matrix directly, "chol" models a Cholesky decomposition, "prec" models a precision matrix, and "ggm" models a Gaussian graphical model (Epskamp, Rhemtulla and Borsboom, 2017). The wrapper `lnm()` sets latent = "ggm" for the latent network model (LNM), the wrapper `rnm()` sets residual = "ggm" for the residual network model (RNM), and the wrapper `lrnm()` combines the LNM and RNM.

Usage

```

lvm(data, lambda, latent = c("cov", "chol", "prec",
  "ggm"), residual = c("cov", "chol", "prec", "ggm"),
  sigma_zeta = "full", kappa_zeta = "full", omega_zeta =
  "full", lowertri_zeta = "full", delta_zeta = "full",
  sigma_epsilon = "diag", kappa_epsilon = "diag",
  omega_epsilon = "zero", lowertri_epsilon = "diag",
  delta_epsilon = "diag", beta = "zero", nu, nu_eta, tau,
  identify = TRUE, identification = c("loadings",
  "variance"), vars, ordered = character(0), latents,
  groups, groupvar, covs, cors, means, nobs,
  missing = "auto", equal = "none",
  baseline_saturated = TRUE, estimator = "default",
  optimizer, storedata = FALSE, WLS.W, covtype =
  c("choose", "ML", "UB"), corinput, standardize = c("none", "z",
  "quantile"), sampleStats, verbose = FALSE,
  simplelambdastart = FALSE, start = "default",
  bootstrap = FALSE,
  boot_sub, boot_resample, penalty_lambda = NA,
  penalty_alpha = 1, penalize_matrices)

```

```
lnm(...)
```

```
rnm(...)
lrnm(...)
```

Arguments

data	A data frame encoding the data used in the analysis. Can be missing if covs and nobS are supplied.
lambda	A model matrix encoding the factor loading structure. Each row indicates an indicator and each column a latent. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
latent	The type of latent model used. See description.
residual	The type of residual model used. See description.
sigma_zeta	Only used when latent = "cov". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_zeta	Only used when latent = "prec". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega_zeta	Only used when latent = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_zeta	Only used when latent = "chol". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_zeta	Only used when latent = "ggm". Either "diag" or "zero", or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
sigma_epsilon	Only used when residual = "cov". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions

	node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_epsilon	Only used when residual = "prec". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega_epsilon	Only used when residual = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_epsilon	Only used when residual = "chol". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_epsilon	Only used when residual = "ggm". Either "diag" or "zero", or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
beta	A model matrix encoding the structural relations between latent variables. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
nu	Optional vector encoding the intercepts of the observed variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
nu_eta	Optional vector encoding the intercepts of the latent variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
tau	Optional matrix encoding the threshold parameters for ordinal variables. Each column corresponds to an observed variable and each row to a threshold. Use 0 for fixed elements, 1 for free elements, and higher integers for equality constrains. Typically set up automatically when ordered is specified.
identify	Logical, should the model be automatically identified?

identification	Type of identification used. "loadings" to fix the first factor loadings to 1, and "variance" to fix the diagonal of the latent variable model matrix (sigma_zeta, lowertri_zeta, delta_zeta or kappa_zeta) to 1.
vars	An optional character vector encoding the variables used in the analysis. Must equal names of the dataset in data.
ordered	Character vector of variable names to treat as ordinal, or TRUE to treat all variables as ordinal. When ordinal variables are specified, polychoric correlations and thresholds are estimated from the data and the model is fit using the Muthen (1984) three-stage estimator. The estimator defaults to "DWLS" for ordinal models; "WLS" and "ULS" are also supported. Identification is automatically set to "variance".
latents	An optional character vector with names of the latent variables.
groups	Deprecated. Use groupvar instead. An optional string indicating the name of the group variable in data.
groupvar	An optional string indicating the name of the group variable in data. Replaces the deprecated groups argument; if both are supplied, groupvar takes precedence with a warning.
covs	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. Make sure covtype argument is set correctly to the type of covariances used.
cors	A sample correlation matrix, or a list/array of such matrices for multiple groups. When supplied, the matrix is treated as a covariance matrix with a warning (appropriate for standardized data). Requires nobs. Note: corinput = TRUE is not supported in lvm; use varcov with type = "cor" for correlation-based models.
means	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
nobs	The number of observations used in covs and means, or a vector of such numbers of observations for multiple groups.
corinput	Logical. Only supported for ordinal data in lvm(); defaults to FALSE for continuous data. Setting corinput = TRUE is not supported for lvm() and will produce an error; use varcov with type = "cor" instead.
missing	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be "auto" (default) for automatic detection, "listwise" for listwise deletion, or "pairwise" for pairwise deletion. When "auto", the function checks for missing data and switches ML to FIML, PML to PFIML, or defaults to listwise for LS estimators.
equal	A character vector indicating which matrices should be constrained equal across groups.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
estimator	The estimator to be used. Currently implemented are "ML" for maximum likelihood estimation, "FIML" for full-information maximum likelihood estimation, "PML" for penalized maximum likelihood estimation, "PFIML" for penalized

full-information maximum likelihood estimation, "ULS" for unweighted least squares estimation, "WLS" for weighted least squares estimation, and "DWLS" for diagonally weighted least squares estimation. "WLSMV" is accepted as a synonym for "DWLS" (both use DWLS estimation with a mean-and-variance adjusted scaled test statistic). When missing = "auto" (default), "ML" is automatically switched to "FIML" and "PML" to "PFIML" if missing data is detected. Defaults to "ML" for continuous data and "DWLS" when ordinal variables are specified via ordered.

optimizer	The optimizer to be used. Can be one of "nlminb" (the default R nlminb function), "ucminf" (from the optimr package), "nloptr_TNEWTON" (preconditioned truncated Newton via nloptr), and "LBFGS++" (pure C++ L-BFGS-B). Defaults to "nlminb".
storedata	Logical, should the raw data be stored? Needed for bootstrapping (see bootstrap).
verbose	Logical, should progress be printed to the console?
WLS.W	The weights matrix used in WLS estimation (experimental)
sampleStats	An optional sample statistics object. Mostly used internally.
covtype	If 'covs' is used, this is the type of covariance (maximum likelihood or unbiased) the input covariance matrix represents. Set to "ML" for maximum likelihood estimates (denominator n) and "UB" to unbiased estimates (denominator n-1). The default will try to find the type used, by investigating which is most likely to result from integer valued datasets.
standardize	Which standardization method should be used? "none" (default) for no standardization, "z" for z-scores, and "quantile" for a non-parametric transformation to the quantiles of the marginal standard normal distribution.
simplelambdastart	Logical, should simple start values be used for lambda? Setting this to TRUE can avoid some estimation problems.
start	Controls the starting values for the beta (structural regression) matrix. Can be "default" (default) for OLS-based starting values derived from the factor-analysis-implied latent covariance matrix, "simple" for the previous behavior (all free elements set to 0.001), or a fitted psychometrics object to extract beta estimates as starting values. The OLS-based starts compute regression coefficients between latent variables using the FA-derived covariance matrix, which typically leads to faster convergence for models with structural paths. Guard rails automatically fall back to simple starts if numerical issues are detected (e.g., near-singular matrices, extreme coefficients).
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets boot_sub = 1 and boot_resample = TRUE) or "case" (which sets boot_sub = 0.75 and boot_resample = FALSE).
boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)

penalty_lambda	Numeric penalty strength for penalized ML estimation (PML/PFIML). NA (default) triggers automatic selection via EBIC-based grid search when a penalized estimator is used; set to a specific numeric value to use a fixed penalty strength (0 = no penalty). See find_penalized_lambda and penalize .
penalty_alpha	Elastic net mixing parameter: 1 = LASSO (default), 0 = ridge.
penalize_matrices	Character vector of matrix names to penalize. If missing, defaults are selected based on the model type.
...	Arguments sent to varcov

Details

The model used in this family is:

$$\text{var}(\mathbf{y}) = \mathbf{\Lambda}(\mathbf{I} - \mathbf{B})^{-1}\mathbf{\Sigma}_{\zeta}(\mathbf{I} - \mathbf{B})^{-1\top}\mathbf{\Lambda}^{\top} + \mathbf{\Sigma}_{\varepsilon}$$

$$\mathcal{E}(\mathbf{y}) = \boldsymbol{\nu} + \mathbf{\Lambda}(\mathbf{I} - \mathbf{B})^{-1}\boldsymbol{\nu}_{eta}$$

in which the latent covariance matrix can further be modeled in three ways. With latent = "chol" as Cholesky decomposition:

$$\mathbf{\Sigma}_{\zeta} = \mathbf{L}_{\zeta}\mathbf{L}_{\zeta}^{\top},$$

with latent = "prec" as Precision matrix:

$$\mathbf{\Sigma}_{\zeta} = \mathbf{K}_{\zeta}^{-1},$$

and finally with latent = "ggm" as Gaussian graphical model:

$$\mathbf{\Sigma}_{\zeta} = \mathbf{\Delta}_{\zeta}(\mathbf{I} - \mathbf{\Omega}_{\zeta})^{-1}\mathbf{\Delta}_{\zeta}.$$

Likewise, the residual covariance matrix can also further be modeled in three ways. With residual = "chol" as Cholesky decomposition:

$$\mathbf{\Sigma}_{\varepsilon} = \mathbf{L}_{\varepsilon}\mathbf{L}_{\varepsilon}^{\top},$$

with latent = "prec" as Precision matrix:

$$\mathbf{\Sigma}_{\varepsilon} = \mathbf{K}_{\varepsilon}^{-1},$$

and finally with latent = "ggm" as Gaussian graphical model:

$$\mathbf{\Sigma}_{\varepsilon} = \mathbf{\Delta}_{\varepsilon}(\mathbf{I} - \mathbf{\Omega}_{\varepsilon})^{-1}\mathbf{\Delta}_{\varepsilon}.$$

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

References

- Epskamp, S., Rhemtulla, M., & Borsboom, D. (2017). Generalized network psychometrics: Combining network and latent variable models. *Psychometrika*, 82(4), 904-927.
- Muthen, B. (1984). A general structural equation model with dichotomous, ordered categorical, and continuous latent variable indicators. *Psychometrika*, 49(1), 115-132.

Examples

```
library("dplyr")

### Confirmatory Factor Analysis ###

# Example also shown in https://youtu.be/Hdu5z-fwuk8

# Load data:
data(StarWars)

# Originals only:
Lambda <- matrix(1,4)

# Model:
mod0 <- lvm(StarWars, lambda = Lambda, vars = c("Q1","Q5","Q6","Q7"),
            identification = "variance", latents = "Originals")

# Run model:
mod0 <- mod0 %>% runmodel

# Evaluate fit:
mod0 %>% fit

# Full analysis
# Factor loadings matrix:
Lambda <- matrix(0, 10, 3)
Lambda[1:4,1] <- 1
Lambda[c(1,5:7),2] <- 1
Lambda[c(1,8:10),3] <- 1

# Observed variables:
obsvars <- paste0("Q",1:10)

# Latents:
latents <- c("Prequels","Original","Sequels")

# Make model:
mod1 <- lvm(StarWars, lambda = Lambda, vars = obsvars,
            identification = "variance", latents = latents)

# Run model:
mod1 <- mod1 %>% runmodel

# Look at fit:
mod1

# Look at parameter estimates:
mod1 %>% parameters

# Look at modification indices:
mod1 %>% MIs
```

```
# Add and refit:
mod2 <- mod1 %>% freepar("sigma_epsilon","Q10","Q4") %>% runmodel

# Compare:
compare(original = mod1, adjusted = mod2)

# Fit measures:
mod2 %>% fit

### Path diagrams ###
# semPlot is not (yet) supported by default, but can be used as follows:
# Load packages:
if (requireNamespace("semPlot", quietly = TRUE)){
  library("semPlot")

# Estimates:
lambdaEst <- getmatrix(mod2, "lambda")
psiEst <- getmatrix(mod2, "sigma_zeta")
thetaEst <- getmatrix(mod2, "sigma_epsilon")

# LISREL Model: LY = Lambda (lambda-y), TE = Theta (theta-epsilon), PS = Psi
mod <- lisrelModel(LY = lambdaEst, PS = psiEst, TE = thetaEst)

# Plot with semPlot:
semPaths(mod, "std", "est", as.expression = "nodes")

# We can make this nicer (set whatLabels = "none" to hide labels):
semPaths(mod,

# this argument controls what the color of edges represent. In this case,
# standardized parameters:
  what = "std",

# This argument controls what the edge labels represent. In this case, parameter
# estimates:
  whatLabels = "est",

# This argument draws the node and edge labels as mathematical expressions:
  as.expression = "nodes",

# This will plot residuals as arrows, closer to what we use in class:
  style = "lisrel",

# This makes the residuals larger:
  residScale = 10,

# qgraph colorblind friendly theme:
  theme = "colorblind",

# tree layout options are "tree", "tree2", and "tree3":
  layout = "tree2",
```

```
# This makes the latent covariances connect at a cardinal center point:
  cardinal = "lat cov",

# Changes curve into rounded straight lines:
  curvePivot = TRUE,

# Size of manifest variables:
  sizeMan = 4,

# Size of latent variables:
  sizeLat = 10,

# Size of edge labels:
  edge.label.cex = 1,

# Sets the margins:
  mar = c(9,1,8,1),

# Prevents re-ordering of observed variables:
  reorder = FALSE,

# Width of the plot:
  width = 8,

# Height of plot:
  height = 5,

# Colors according to latents:
  groups = "latents",

# Pastel colors:
  pastel = TRUE,

# Disable borders:
  borders = FALSE
)

# Use arguments filetype = "pdf" and filename = "semPlotExample1" to store PDF
} # end semPlot block

### Latent Network Modeling ###

# Latent network model:
lnm <- lvm(StarWars, lambda = Lambda, vars = obsvars,
          latents = latents, identification = "variance",
          latent = "ggm")

# Run model:
lnm <- lnm %>% runmodel

# Look at parameters:
lnm %>% parameters
```

```

# Remove non-sig latent edge:
lnm <- lnm %>% prune(alpha = 0.05)

# Compare to the original CFA model:
compare(cfa = mod1, lnm = lnm)

# Plot network:
library("qgraph")
qgraph(lnm@modelmatrices[[1]]$omega_zeta, labels = latents,
       theme = "colorblind", vsize = 10)

# A wrapper for the latent network model is the lnm function:
lnm2 <- lnm(StarWars, lambda = Lambda, vars = obsvars,
           latents = latents, identification = "variance")
lnm2 <- lnm2 %>% runmodel %>% prune(alpha = 0.05)
compare(lnm, lnm2) # Is the same as the model before.

# I could also estimate a "residual network model", which adds partial correlations to
# the residual level:
# This can be done using lvm(..., residual = "ggm") or with rnm(...)
rnm <- rnm(StarWars, lambda = Lambda, vars = obsvars,
          latents = latents, identification = "variance")
# Stepup search:
rnm <- rnm %>% stepup

# It will estimate the same model (with link Q10 - Q4) as above. In the case of only one
# partial correlation, There is no difference between residual covariances (SEM) or
# residual partial correlations (RNM).

# For more information on latent and residual network models, see:
# Epskamp, S., Rhemtulla, M.T., & Borsboom, D. Generalized Network Psychometrics:
# Combining Network and Latent Variable Models
# (2017). Psychometrika. doi:10.1007/s11336-017-9557-x

### Gaussian graphical models ###

# All psychonetrics functions (e.g., lvm, lnm, rnm...) allow input via a covariance
# matrix, with the "covs" and "nobs" arguments.
# The following fits a baseline GGM network with no edges:
S <- (nrow(StarWars) - 1) / (nrow(StarWars)) * cov(StarWars[,1:10])
ggmmod <- ggm(covs = S, nobs = nrow(StarWars))

# Run model with stepup search and pruning:
ggmmod <- ggmmod %>% prune %>% modelsearch

# Fit measures:
ggmmod %>% fit

# Plot network:
nodeNames <- c(
  "I am a huge Star Wars\nfan! (star what?)",

```

```

"I would trust this person\nwith my democracy.",
"I enjoyed the story of\nAnakin's early life.",
"The special effects in\nthis scene are awful (Battle of\nGeonosis).",
"I would trust this person\nwith my life.",
"I found Darth Vader's big\nreveal in 'Empire' one of the greatest
moments in movie history.",
"The special effects in\nthis scene are amazing (Death Star\nExplosion).",
"If possible, I would\ndefinitely buy this\nndroid.",
"The story in the Star\nWars sequels is an improvement to\nthe previous movies.",
"The special effects in\nthis scene are marvellous (Starkiller\nBase Firing)."
)
library("qgraph")
qgraph(as.matrix(ggmmmod@modelmatrices[[1]]$omega), nodeNames = nodeNames,
       legend.cex = 0.25, theme = "colorblind", layout = "spring")

# We can actually compare this model statistically (note they are not nested) to the
# latent variable model:
compare(original_cfa = mod1, adjusted_cfa = mod2, exploratory_ggm = ggmmmod)

### Measrement invariance ###
# Let's say we are interested in seeing if people >= 30 like the original trilogy better
# than people < 30.
# First we can make a grouping variable:
StarWars$agegroup <- ifelse(StarWars$Q12 < 30, "young", "less young")

# Let's look at the distribution:
table(StarWars$agegroup) # Pretty even...

# Observed variables:
obsvars <- paste0("Q",1:10)

# Let's look at the mean scores:
StarWars %>% group_by(agegroup) %>% summarize(across(all_of(obsvars), mean))
# Less young people seem to score higher on prequel questions and lower on other
# questions

# Factor loadings matrix:
Lambda <- matrix(0, 10, 3)
Lambda[1:4,1] <- 1
Lambda[c(1,5:7),2] <- 1
Lambda[c(1,8:10),3] <- 1

# Residual covariances:
Theta <- diag(1, 10)
Theta[4,10] <- Theta[10,4] <- 1

# Latents:
latents <- c("Prequels", "Original", "Sequels")

# Make model:
mod_configural <- lvm(StarWars, lambda = Lambda, vars = obsvars,
                    latents = latents, sigma_epsilon = Theta,

```

```

        identification = "variance",
        groups = "agegroup")

# Run model:
mod_configural <- mod_configural %>% runmodel

# Look at fit:
mod_configural
mod_configural %>% fit

# Looks good, let's try weak invariance:
mod_weak <- mod_configural %>% groupequal("lambda") %>% runmodel

# Compare models:
compare(configural = mod_configural, weak = mod_weak)

# weak invariance can be accepted, let's try strong:
mod_strong <- mod_weak %>% groupequal("nu") %>% runmodel
# Means are automatically identified

# Compare models:
compare(configural = mod_configural, weak = mod_weak, strong = mod_strong)

# Questionable p-value and AIC difference, but ok BIC difference. This is quite good, but
# let's take a look. I have not yet implemented LM tests for equality constrains, but we
# can look at something called "equality-free" MIs:
mod_strong %>% MIs(matrices = "nu", type = "free")

# Indicates that Q10 would improve fit. We can also look at residuals:
residuals(mod_strong)

# Let's try freeing intercept 10:
mod_strong_partial <- mod_strong %>% groupfree("nu",10) %>% runmodel

# Compare all models:
compare(configural = mod_configural,
        weak = mod_weak,
        strong = mod_strong,
        strong_partial = mod_strong_partial)

# This seems worth it and lead to an acceptable model! It seems that older people find
# the latest special effects more marvellous!
mod_strong_partial %>% getmatrix("nu")

# Now let's investigate strict invariance:
mod_strict <- mod_strong_partial %>% groupequal("sigma_epsilon") %>% runmodel

# Compare all models:
compare(configural = mod_configural,
        weak = mod_weak,
        strong_partial = mod_strong_partial,
        strict = mod_strict)
# Strict invariance can be accepted!

```

```

# Now we can test for homogeneity!
# Are the latent variances equal?
mod_eqvar <- mod_strict %>% groupequal("sigma_zeta") %>% runmodel

# Compare:
compare(strict = mod_strict, eqvar = mod_eqvar)

# This is acceptable. What about the means? (alpha = nu_eta)
mod_eqmeans <- mod_eqvar %>% groupequal("nu_eta") %>% runmodel

# Compare:
compare(eqvar = mod_eqvar, eqmeans = mod_eqmeans)

# Rejected! We could look at MIs again:
mod_eqmeans %>% MIs(matrices = "nu_eta", type = "free")

# Indicates the strongest effect for prequels. Let's see what happens:
eqmeans2 <- mod_eqvar %>%
  groupequal("nu_eta", row = c("Original", "Sequels")) %>% runmodel

# Compare:
compare(eqvar = mod_eqvar, eqmeans = eqmeans2)
# Questionable, what about the sequels as well?

eqmeans3 <- mod_eqvar %>% groupequal("nu_eta", row = "Original") %>% runmodel

# Compare:
compare(eqvar = mod_eqvar, eqmeans = eqmeans3)

# Still questionable.. Let's look at the mean differences:
mod_eqvar %>% getmatrix("nu_eta")

# Looks like people over 30 like the prequels better and the other two trilogies less!

```

meta_lvm

Meta-analytic latent variable model

Description

Single-stage meta-analytic latent variable model (LVM) for covariance or correlation matrices. Combines a latent variable model (CFA/SEM) for the mean structure with a random-effects model for between-study heterogeneity. Based on the MAGNA framework (Epskamp et al., 2024).

Usage

```

meta_lvm(covs, nobs, data, cors, studyvar, groups, groupvar,
         corinput, Vmats, Vmethod = c("individual", "pooled",
         "metaSEM_individual", "metaSEM_weighted"), Vestimation

```

```

= c("averaged", "per_study"),
lambda, beta = "zero",
latent = c("cov", "chol", "prec", "ggm"),
sigma_zeta = "full", kappa_zeta = "full",
omega_zeta = "full", lowertri_zeta = "full",
delta_zeta = "full",
residual = c("cov", "chol", "prec", "ggm"),
sigma_epsilon = "diag", kappa_epsilon = "diag",
omega_epsilon = "zero", lowertri_epsilon = "diag",
delta_epsilon = "diag",
identify = TRUE,
identification = c("loadings", "variance"),
randomEffects = c("chol", "cov", "prec", "ggm", "cor"),
sigma_randomEffects = "full",
kappa_randomEffects = "full",
omega_randomEffects = "full",
lowertri_randomEffects = "full",
delta_randomEffects = "full",
rho_randomEffects = "full",
SD_randomEffects = "full",
vars, latents,
baseline_saturated = TRUE, optimizer,
estimator = c("FIML", "ML"),
sampleStats, verbose = FALSE,
bootstrap = FALSE, boot_sub, boot_resample)

```

Arguments

covs	A list of covariance or correlation matrices. Must contain rows and columns with NAs for variables not included in a study.
nobs	A vector with the number of observations per study.
data	Optional data frame. When supplied together with studyvar, covariance matrices and sample sizes are computed internally per study. Cannot be used together with covs, cors, or nobs.
cors	A list of correlation matrices. When supplied, the matrices are treated as covariance matrices with a warning (appropriate for standardized data). Requires nobs.
studyvar	A string indicating the column name in data that identifies the study. Required when data is supplied.
groups	Deprecated. Use groupvar instead. Multi-group support is not yet included for meta-analytic models.
groupvar	Not yet supported for meta-analytic models. Supplying this argument will produce an error.
corinput	Logical. Defaults to FALSE. Controls whether the input is treated as correlation matrices.
Vmats	Optional list with 'V' matrices (sampling error variance approximations).

vmethod	Which method should be used to approximate the sampling error variance?
vestimation	How should the sampling error estimates be evaluated?
lambda	A matrix encoding the factor loading structure. Can be a pattern matrix with TRUE/FALSE or 0/1 entries, or a matrix with starting values and higher integer codes for equality constraints.
beta	Structural (regression) matrix among latent variables. Defaults to "zero".
latent	Parameterization of the latent covariance structure. One of "cov", "chol", "prec", or "ggm".
sigma_zeta	Latent covariance matrix specification (used when latent = "cov").
kappa_zeta	Latent precision matrix specification (used when latent = "prec").
omega_zeta	Latent partial correlation matrix specification (used when latent = "ggm").
lowertri_zeta	Latent Cholesky factor specification (used when latent = "chol").
delta_zeta	Latent scaling matrix specification (used when latent = "ggm").
residual	Parameterization of the residual covariance structure. One of "cov", "chol", "prec", or "ggm".
sigma_epsilon	Residual covariance matrix specification (used when residual = "cov"). Defaults to "diag". Diagonal elements are always fixed (derived from the correlation constraint).
kappa_epsilon	Residual precision matrix specification (used when residual = "prec").
omega_epsilon	Residual partial correlation matrix specification (used when residual = "ggm").
lowertri_epsilon	Residual Cholesky factor specification (used when residual = "chol").
delta_epsilon	Residual scaling matrix specification (used when residual = "ggm").
identify	Logical, should the model be identified? Defaults to TRUE.
identification	How to identify the model. "loadings" or "variance".
randomEffects	Parameterization of the random effects covariance structure.
sigma_randomEffects	Random effects covariance matrix specification (used when randomEffects = "cov").
kappa_randomEffects	Random effects precision matrix specification (used when randomEffects = "prec").
omega_randomEffects	Random effects partial correlation matrix specification (used when randomEffects = "ggm").
lowertri_randomEffects	Random effects Cholesky factor specification (used when randomEffects = "chol").
delta_randomEffects	Random effects scaling matrix specification (used when randomEffects = "ggm").
rho_randomEffects	Random effects correlation matrix specification (used when randomEffects = "cor").

SD_randomEffects	Random effects standard deviation matrix specification (used when randomEffects = "cor").
vars	Character vector of observed variable names. If missing, names are taken from the correlation matrices.
latents	Character vector of latent variable names.
baseline_saturated	Logical indicating if baseline and saturated models should be included.
optimizer	The optimizer to be used. Defaults to "nlminb".
estimator	The estimator to be used. "ML" or "FIML" (default).
sampleStats	Optional sample statistics object.
verbose	Logical, should progress be printed?
bootstrap	Should the data be bootstrapped?
boot_sub	Proportion of cases to subsample for bootstrap.
boot_resample	Logical, should bootstrap be with replacement?

Details

This function implements a single-stage meta-analytic latent variable model. The model specifies that the mean of the vectorized correlation matrices follows an LVM-implied correlation structure:

$$\mu = \text{vechs}(\Lambda(I - B)^{-1}\Sigma_{\zeta}(I - B)^{-\top}\Lambda' + \Sigma_{\varepsilon})$$

where diagonal elements of Σ_{ε} are derived from the constraint that μ represents correlations (diagonal of implied covariance = 1).

Between-study heterogeneity is modeled as:

$$\Sigma = \Sigma^{(\text{ran})} + V$$

where V is the known sampling error covariance and $\Sigma^{(\text{ran})}$ captures true between-study variation.

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

References

- Epskamp, S. et al. (2024). Meta-Analytic Gaussian Network Aggregation. *Psychometrika*.
- Jak, S., and Cheung, M. W. L. (2019). Meta-analytic structural equation modeling with moderating effects on SEM parameters. *Psychological Methods*.

See Also

[meta_varcov](#), [lvm](#)

Examples

```
## Not run:
# Generate simulated data
set.seed(42)
nStudies <- 10
nvar <- 6
nlat <- 2

# True factor loading matrix
lambda_true <- matrix(0, nvar, nlat)
lambda_true[1:3, 1] <- c(0.7, 0.8, 0.6)
lambda_true[4:6, 2] <- c(0.7, 0.8, 0.6)

# Generate correlation matrices per study
cors <- list()
nobs <- sample(100:400, nStudies)
for (i in 1:nStudies) {
  sigma <- lambda_true %*% t(lambda_true) + diag(1 - rowSums(lambda_true^2))
  dat <- MASS::mvrnorm(nobs[i], rep(0, nvar), sigma)
  cors[[i]] <- cor(dat)
  colnames(cors[[i]]) <- rownames(cors[[i]]) <- paste0("V", 1:nvar)
}

# Fit meta-analytic CFA
mod <- meta_lvm(covs = cors, nobs = nobs, lambda = lambda_true != 0)
mod <- mod %>% runmodel

# Inspect results
print(mod)
parameters(mod)
getmatrix(mod, "lambda")

## End(Not run)
```

meta_var1

Meta-analytic VAR(1) model

Description

Single-stage meta-analytic vector autoregressive model (VAR(1)) for time-series data collected across multiple studies. Pools temporal (lag-1) and contemporaneous network structures across studies using a random-effects framework. The `meta_gvar` wrapper sets `contemporaneous = "ggm"` for graphical VAR estimation.

Usage

```
meta_var1(data, covs, nobs,
          vars, studyvar, idvar, dayvar, beepvar,
          contemporaneous = c("cov", "chol", "prec", "ggm"),
          beta = "full",
          omega_zeta = "full", delta_zeta = "full",
          kappa_zeta = "full", sigma_zeta = "full",
          lowertri_zeta = "full",
          randomEffects = c("chol", "cov", "prec", "ggm", "cor"),
          sigma_randomEffects = "full",
          kappa_randomEffects = "full",
          omega_randomEffects = "full",
          lowertri_randomEffects = "full",
          delta_randomEffects = "full",
          rho_randomEffects = "full",
          SD_randomEffects = "full",
          Vmats,
          Vmethod = c("individual", "pooled"),
          Vestimation = c("averaged", "per_study"),
          baseline_saturated = TRUE, optimizer,
          estimator = c("FIML", "ML"),
          sampleStats, verbose = FALSE,
          bootstrap = FALSE, boot_sub, boot_resample)

meta_gvar(...)
```

Arguments

data	A data frame containing time-series data for multiple studies. Use together with studyvar to identify studies.
covs	A list of pre-computed Toeplitz covariance matrices (one per study). Each matrix should have dimension $2p \times 2p$ where p is the number of variables, with the first p rows/columns being lagged variables and the second p being current variables. Alternative to data.
nobs	A vector with the number of observations per study. Required when covs is supplied.
vars	Character vector of observed variable names. If missing, inferred from data.
studyvar	A string indicating the column name in data that identifies the study. Required when data is supplied. If not supplied but idvar is, idvar is used as studyvar with a warning.
idvar	Optional string indicating the subject ID column within each study. When both studyvar and idvar are supplied, data is first split by studyvar, then idvar is used within each study to collate time series (as in var1).
dayvar	Optional string indicating the day variable (passed to tsData per study).
beepvar	Optional string indicating the beep variable within day (passed to tsData per study).

contemporaneous	Parameterization of the contemporaneous (residual innovation) covariance structure. One of "cov", "chol", "prec", or "ggm".
beta	Temporal lag-1 regression matrix specification. Defaults to "full" (all elements free, including diagonal autoregressive effects).
omega_zeta	Contemporaneous partial correlation matrix specification (used when contemporaneous = "ggm").
delta_zeta	Contemporaneous scaling matrix specification (used when contemporaneous = "ggm").
kappa_zeta	Contemporaneous precision matrix specification (used when contemporaneous = "prec").
sigma_zeta	Contemporaneous covariance matrix specification (used when contemporaneous = "cov").
lowertri_zeta	Contemporaneous Cholesky factor specification (used when contemporaneous = "chol").
randomEffects	Parameterization of the random effects covariance structure.
sigma_randomEffects	Random effects covariance matrix specification (used when randomEffects = "cov").
kappa_randomEffects	Random effects precision matrix specification (used when randomEffects = "prec").
omega_randomEffects	Random effects partial correlation matrix specification (used when randomEffects = "ggm").
lowertri_randomEffects	Random effects Cholesky factor specification (used when randomEffects = "chol").
delta_randomEffects	Random effects scaling matrix specification (used when randomEffects = "ggm").
rho_randomEffects	Random effects correlation matrix specification (used when randomEffects = "cor").
SD_randomEffects	Random effects standard deviation matrix specification (used when randomEffects = "cor").
Vmats	Optional list with 'V' matrices (sampling error variance approximations).
Vmethod	Which method should be used to approximate the sampling error variance? "individual" or "pooled".
Vestimation	How should the sampling error estimates be evaluated? "averaged" or "per_study".
baseline_saturated	Logical indicating if baseline and saturated models should be included.
optimizer	The optimizer to be used. Defaults to "nlsminb".
estimator	The estimator to be used. "ML" or "FIML" (default).

sampleStats	Optional sample statistics object.
verbose	Logical, should progress be printed?
bootstrap	Should the data be bootstrapped?
boot_sub	Proportion of cases to subsample for bootstrap.
boot_resample	Logical, should bootstrap be with replacement?
...	Arguments sent to meta_var1.

Details

This function implements a single-stage meta-analytic VAR(1) model. For p observed variables, each study's time-series data is transformed into a Toeplitz covariance structure from which two blocks are extracted:

- **Sigma_0**: the $p \times p$ stationary covariance (symmetric, $p(p+1)/2$ unique elements)
- **Sigma_1**: the $p \times p$ lag-1 cross-covariance (non-symmetric, p^2 elements)

The VAR(1) structural model implies:

$$\text{vec}(\Sigma_0) = (I - \beta \otimes \beta)^{-1} \text{vec}(\Sigma_\zeta)$$

$$\Sigma_1 = \beta \Sigma_0$$

The meta-level mean is $\mu = [\text{vech}(\Sigma_0), \text{vec}(\Sigma_1)]$ and heterogeneity is modeled as $\Sigma = \Sigma^{(\text{ran})} + V$.

Note: the exogenous (lagged) covariance block from the full Toeplitz matrix is not modeled, as it is a nuisance parameter at the meta-analytic level.

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

See Also

[var1](#), [meta_varcov](#), [meta_lvm](#)

Examples

```
## Not run:
library(mlVAR)
set.seed(42)
Model <- mlVARsim(nPerson = 50, nNode = 3, nTime = 50, lag = 1)

# Fit meta-analytic VAR(1):
mod <- meta_var1(Model$Data,
                 vars = Model$varNames,
                 studyvar = Model$idvar,
```

```

        estimator = "ML")
mod <- mod %>% runmodel

# Inspect results:
print(mod)
getmatrix(mod, "beta")

# Fit meta-analytic GVAR (with GGM contemporaneous):
mod_gvar <- meta_gvar(Model$Data,
  vars = Model$varNames,
  studyvar = Model$idvar,
  estimator = "ML")
mod_gvar <- mod_gvar %>% runmodel
getmatrix(mod_gvar, "omega_zeta")

## End(Not run)

```

 meta_varcov

Variance-covariance and GGM meta analysis

Description

Meta analysis of correlation matrices to fit a homogenous correlation matrix or Gaussian graphical model. Based on meta-analytic SEM (Jak and Cheung, 2019).

Usage

```

meta_varcov(cors, nobs, data, covs, studyvar, groups, groupvar,
  corinput, Vmats, Vmethod = c("individual", "pooled",
  "metaSEM_individual", "metaSEM_weighted"), Vestimation
  = c("averaged", "per_study"), type = c("cor", "ggm"),
  sigma_y = "full", kappa_y = "full", omega_y = "full",
  lowertri_y = "full", delta_y = "full", rho_y = "full",
  SD_y = "full", randomEffects = c("chol", "cov",
  "prec", "ggm", "cor"), sigma_randomEffects = "full",
  kappa_randomEffects = "full", omega_randomEffects =
  "full", lowertri_randomEffects = "full",
  delta_randomEffects = "full", rho_randomEffects =
  "full", SD_randomEffects = "full", vars,
  baseline_saturated = TRUE, optimizer, estimator =
  c("FIML", "ML"), sampleStats, verbose = FALSE,
  bootstrap = FALSE, boot_sub, boot_resample)

```

```
meta_ggm(...)
```

Arguments

cors A list of correlation matrices. Must contain rows and columns with NAs for variables not included in a study.

nobs	A vector with the number of observations per study.
data	Optional data frame. When supplied together with <code>studyvar</code> , correlation matrices and sample sizes are computed internally per study. Cannot be used together with <code>cors</code> , <code>covs</code> , or <code>nobs</code> .
covs	A list of covariance matrices. Alternative to <code>cors</code> ; when supplied, <code>corinput</code> defaults to <code>FALSE</code> .
studyvar	A string indicating the column name in <code>data</code> that identifies the study. Required when <code>data</code> is supplied.
groups	Deprecated. Use <code>groupvar</code> instead. Multi-group support is not yet included for meta-analytic models.
groupvar	Not yet supported for meta-analytic models. Supplying this argument will produce an error.
corinput	Logical. Defaults to <code>TRUE</code> when <code>cors</code> is used, <code>FALSE</code> when <code>covs</code> is used. Controls whether the input is treated as correlation matrices.
Vmats	Optional list with 'V' matrices (sampling error variance approximations).
Vmethod	Which method should be used to approximate the sampling error variance?
Vestimation	How should the sampling error estimates be evaluated?
type	What to model? Currently only "cor" and "ggm" are supported.
sigma_y	Only used when <code>type = "cov"</code> . Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_y	Only used when <code>type = "prec"</code> . Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega_y	Only used when <code>type = "ggm"</code> . Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_y	Only used when <code>type = "chol"</code> . Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_y	Only used when <code>type = "ggm"</code> . Either "diag" or "zero" (not recommended), or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality

	constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
rho_y	Only used when type = "cor". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
SD_y	Only used when type = "cor". Either "diag" or "zero", or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
randomEffects	What to model for the random effects?
sigma_randomEffects	Only used when type = "cov". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_randomEffects	Only used when randomEffects = "prec". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega_randomEffects	Only used when randomEffects = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_randomEffects	Only used when randomEffects = "chol". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_randomEffects	Only used when randomEffects = "ggm". Either "diag" or "zero", or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

rho_randomEffects	Only used when randomEffects = "cor". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
SD_randomEffects	Only used when randomEffects = "cor". Either "diag" or "zero", or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
vars	Variables to be included.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
optimizer	The optimizer to be used. Can be one of "nlminb" (the default R nlminb function), "ucminf" (from the optimr package), "nloptr_TNEWTON" (preconditioned truncated Newton via nloptr), and "LBFGS++" (pure C++ L-BFGS-B). Defaults to "nlminb".
estimator	The estimator to be used. Currently implemented are "ML" for maximum likelihood estimation or "FIML" for full-information maximum likelihood estimation.
sampleStats	An optional sample statistics object. Mostly used internally.
verbose	Logical, should progress be printed to the console?
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets boot_sub = 1 and boot_resample = TRUE) or "case" (which sets boot_sub = 0.75 and boot_resample = FALSE).
boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
...	Arguments sent to meta_varcov

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

References

Jak, S., and Cheung, M. W. L. (2019). Meta-analytic structural equation modeling with moderating effects on SEM parameters. Psychological methods.

MIs *Print modification indices*

Description

This function prints a list of modification indices (MIs)

Usage

```
MIs(x, all = FALSE, matrices, type = c("normal", "equal", "free"), top = 10,  
    verbose = TRUE, nonZero = FALSE)
```

Arguments

x	A psychometrics model.
all	Logical, should all MIs be printed or only the highest?
matrices	Optional vector of matrices to include in the output.
type	String indicating which kind of modification index should be printed. ("mi" is the typical MI, "mi_free" is the modification index free from equality constraints across groups, and "mi_equal" is the modification index if the parameter is added constrained equal across all groups).
top	Number of MIs to include in output if all = FALSE
verbose	Logical, should messages be printed?
nonZero	Logical, should only MIs be printed of non-zero parameters? Useful to explore violations of group equality.

Value

Invisibly returns a relevant subset of the data frame containing all information on the parameters, or a list of such data frames if multiple types of MIs are requested.

Author(s)

Sacha Epskamp

Examples

```
# Load bfi data from psych package:  
library("psychTools")  
data(bfi)  
  
# Also load dplyr for the pipe operator:  
library("dplyr")  
  
# Let's take the agreeableness items, and gender:  
ConsData <- bfi %>%
```

```

select(A1:A5, gender) %>%
na.omit # Let's remove missingness (otherwise use Estimator = "FIML)

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars, omega = "zero")

# Run model:
mod <- mod %>% runmodel

# Modification indices:
mod %>% MIs

```

ml_lvm

*Multi-level latent variable model family***Description**

This family is the two-level random intercept variant of the [lvm](#) model family. It is mostly a special case of the [dlvm1](#) family, with the addition of structural effects rather than temporal effects in the beta matrix.

Usage

```

ml_lnm(...)
ml_rnm(...)
ml_lrnm(...)
ml_lvm(data, lambda, clusters, within_latent = c("cov",
"chol", "prec", "ggm"), within_residual = c("cov",
"chol", "prec", "ggm"), between_latent = c("cov",
"chol", "prec", "ggm"), between_residual = c("cov",
"chol", "prec", "ggm"), beta_within = "zero",
beta_between = "zero", omega_zeta_within = "full",
delta_zeta_within = "full", kappa_zeta_within =
"full", sigma_zeta_within = "full",
lowertri_zeta_within = "full", omega_epsilon_within =
"zero", delta_epsilon_within = "diag",
kappa_epsilon_within = "diag", sigma_epsilon_within =
"diag", lowertri_epsilon_within = "diag",
omega_zeta_between = "full", delta_zeta_between =
"full", kappa_zeta_between = "full",
sigma_zeta_between = "full", lowertri_zeta_between =
"full", omega_epsilon_between = "zero",
delta_epsilon_between = "diag", kappa_epsilon_between =
"diag", sigma_epsilon_between = "diag",
lowertri_epsilon_between = "diag", nu, nu_eta,

```

```

identify = TRUE, identification = c("loadings",
"variance"), vars, latents, groups, equal = "none",
baseline_saturated = TRUE, estimator = c("FIML",
"MUML"), optimizer, storedata = FALSE, verbose =
FALSE, standardize = c("none", "z", "quantile"),
sampleStats, bootstrap = FALSE, boot_sub,
boot_resample)

```

Arguments

<code>data</code>	A data frame encoding the data used in the analysis. Must be a raw dataset.
<code>lambda</code>	A model matrix encoding the factor loading structure. Each row indicates an indicator and each column a latent. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix. Could also be the result of simplestructure .
<code>clusters</code>	A string indicating the variable in the dataset that describes group membership.
<code>within_latent</code>	The type of within-person latent contemporaneous model to be used.
<code>within_residual</code>	The type of within-person residual model to be used.
<code>between_latent</code>	The type of between-person latent model to be used.
<code>between_residual</code>	The type of between-person residual model to be used.
<code>beta_within</code>	A model matrix encoding the within-cluster structural. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix. Defaults to "zero".
<code>beta_between</code>	A model matrix encoding the between-cluster structural. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix. Defaults to "zero".
<code>omega_zeta_within</code>	Only used when <code>within_latent = "ggm"</code> . Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>delta_zeta_within</code>	Only used when <code>within_latent = "ggm"</code> . Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>kappa_zeta_within</code>	Only used when <code>within_latent = "prec"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating

free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`sigma_zeta_within`

Only used when `within_latent = "cov"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`lowertri_zeta_within`

Only used when `within_latent = "chol"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`omega_epsilon_within`

Only used when `within_residual = "ggm"`. Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`delta_epsilon_within`

Only used when `within_residual = "ggm"`. Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`kappa_epsilon_within`

Only used when `within_residual = "prec"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`sigma_epsilon_within`

Only used when `within_residual = "cov"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`lowertri_epsilon_within`

Only used when `within_residual = "chol"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`omega_zeta_between`

Only used when `between_latent = "ggm"`. Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating

free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`delta_zeta_between`

Only used when `between_latent = "ggm"`. Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`kappa_zeta_between`

Only used when `between_latent = "prec"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`sigma_zeta_between`

Only used when `between_latent = "cov"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`lowertri_zeta_between`

Only used when `between_latent = "chol"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`omega_epsilon_between`

Only used when `between_residual = "ggm"`. Can be "full", "zero", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`delta_epsilon_between`

Only used when `between_residual = "ggm"`. Can be "diag", "zero" (not recommended), or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`kappa_epsilon_between`

Only used when `between_residual = "prec"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

`sigma_epsilon_between`

Only used when `between_residual = "cov"`. Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For

	multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_epsilon_between	Only used when <code>between_residual = "chol"</code> . Can be "full", "diag", or a typical model matrix with 0s indicating parameters constrained to zero, 1s indicating free parameters, and higher integers indicating equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
nu	Optional vector encoding the intercepts of the observed variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
nu_eta	Optional vector encoding the intercepts of the latent variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
identify	Logical, should the model be automatically identified?
identification	Type of identification used. "loadings" to fix the first factor loadings to 1, and "variance" to fix the diagonal of the latent variable model matrix (<code>sigma_zeta</code> , <code>lowertri_zeta</code> , <code>delta_zeta</code> or <code>kappa_zeta</code>) to 1.
vars	An optional character vector with names of the variables used.
latents	An optional character vector with names of the latent variables.
groups	An optional string indicating the name of the group variable in data.
equal	A character vector indicating which matrices should be constrained equal across groups.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
estimator	Estimator used. Currently only "FIML" is supported.
optimizer	The optimizer to be used. Usually either "nllminb" (with box constrains) or "ucminf" (ignoring box constrains), but any optimizer supported by <code>optimr</code> can be used.
storedata	Logical, should the raw data be stored? Needed for bootstrapping (see bootstrap).
verbose	Logical, should progress be printed to the console?
standardize	Which standardization method should be used? "none" (default) for no standardization, "z" for z-scores, and "quantile" for a non-parametric transformation to the quantiles of the marginal standard normal distribution.
sampleStats	An optional sample statistics object. Mostly used internally.
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets <code>boot_sub = 1</code> and <code>boot_resample = TRUE</code>) or "case" (which sets <code>boot_sub = 0.75</code> and <code>boot_resample = FALSE</code>).

boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
...	Arguments sent to 'ml_lvm'

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

ml_tsdlvm1	<i>Multi-level Lag-1 dynamic latent variable model family of psychometrics models for time-series data</i>
------------	--

Description

Deprecated. Use [dlvm1](#) directly with long-format data instead (set vars to a character vector and provide idvar).

This function is a deprecated wrapper around [dlvm1](#) that allows for specifying the model using a long format data and similar input as the mlVAR package. The ml_ts_lvgvar simply sets within_latent = "ggm" and between_latent = "ggm" by default. The ml_gvar and ml_var are simple wrappers with different named defaults for contemporaneous and between-person effects. All these functions now call dlvm1() directly and emit deprecation warnings.

Usage

```
ml_tsdlvm1(data, beepvar, idvar, vars, groups, estimator = "FIML",
  standardize = c("none", "z", "quantile"), ...)
```

```
ml_ts_lvgvar(...)
```

```
ml_gvar(..., contemporaneous = c("ggm", "cov", "chol", "prec"),
  between = c("ggm", "cov", "chol", "prec"))
```

```
ml_var(..., contemporaneous = c("cov", "chol", "prec", "ggm"),
  between = c("cov", "chol", "prec", "ggm"))
```

Arguments

data	The data to be used. Must be raw data in long format (each row indicates one person at one time point).
beepvar	Optional string indicating assessment beep per day. Adding this argument will cause non-consecutive beeps to be treated as missing!

idvar	String indicating the subject ID
vars	Vectors of variables to include in the analysis
groups	An optional string indicating the name of the group variable in data.
estimator	Estimator to be used. Must be "FIML".
standardize	Which standardization method should be used? "none" (default) for no standardization, "z" for z-scores, and "quantile" for a non-parametric transformation to the quantiles of the marginal standard normal distribution.
contemporaneous	The type of within-person latent contemporaneous model to be used.
between	The type of between-person latent model to be used.
...	Arguments sent to dlvm1

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

 modelsearch

Stepwise model search

Description

This function performs stepwise model search to find an optimal model that (locally) minimizes some criterion (by default, the BIC).

Usage

```
modelsearch(x, criterion = "bic", matrices, prunealpha = 0.01,
            addalpha = 0.01, verbose, ...)
```

Arguments

x	A psychometrics model.
criterion	String indicating the criterion to minimize. Any criterion from fit can be used.
matrices	Vector of strings indicating which matrices should be searched. Will default to network structures and factor loadings.
prunealpha	Minimal alpha used to consider edges to be removed
addalpha	Maximum alpha used to consider edges to be added
verbose	Logical, should messages be printed?
...	Arguments sent to runmodel

Details

The full algorithm is as follows:

1. Evaluate all models in which an edge is removed that has $p > \text{prunealpha}$, or an edge is added that has a modification index with $p < \text{addalpha}$
2. If none of these models improve the criterion, return the previous model and stop the algorithm
3. Update the model to the model that improved the criterion the most
4. Evaluate all other considered models that improved the criterion
5. If none of these models improve the criterion, go to 1, else go to 3

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

See Also

[prune](#), [stepup](#)

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML)

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars)

# Run model:
mod <- mod %>% runmodel

# Model search
mod <- mod %>% prune(alpha= 0.01) %>% modelsearch
```

Description

A subset of self-report data collected in 2020 by Adela-Maria Isvoranu as part of a graduate-level course on network analysis at the University of Amsterdam. Students in the course collected the data from a convenience sample. The full dataset contains responses to 87 items covering topics such as sleep, well-being, and social functioning. This subset contains 8 items used in Chapter 6 of the textbook *Network Psychometrics with R* (Epskamp, Isvoranu, & Haslbeck, 2022) to demonstrate Gaussian Graphical Model estimation with psychometrics. All items are rated on a 1–7 Likert scale. The dataset contains some missing values, making it suitable for demonstrating FIML estimation.

Usage

```
data("NA2020")
```

Format

A data frame with 501 observations on 8 variables.

regular_sleep I try to keep a regular sleep pattern (Q10)

worried_sleep I am worried about my current sleeping behavior (Q13)

sleep_interfere My sleep interferes with my daily functioning (Q14)

happy_health I am happy with my physical health (Q68)

optimistic_future I feel optimistic about the future (Q70)

very_happy I am very happy (Q75)

feel_alone I often feel alone (Q77)

happy_love_life I am happy with my love life (Q80)

Source

The full dataset is available at <https://osf.io/45n6d/>. Data collected by Adela-Maria Isvoranu as part of a graduate course on network analysis at the University of Amsterdam (2020).

References

Epskamp, S., Haslbeck, J. M. B., Isvoranu, A. M., & Van Borkulo, C. D. (2022). Pairwise Markov random fields. In A. M. Isvoranu, S. Epskamp, L. J. Waldorp, & D. Borsboom (Eds.), *Network psychometrics with R: A guide for behavioral and social scientists* (pp. 95–118). Routledge.

Examples

```
data(NA2020)

# Estimate a GGM using FIML (as in Chapter 6):
library(dplyr)
mod <- ggm(NA2020, estimator = "FIML") %>% runmodel

# Prune non-significant edges:
mod_pruned <- mod %>% prune(alpha = 0.05)

# Inspect parameters:
mod_pruned %>% parameters
```

parameters

Print parameter estimates

Description

This function will print a list of parameters of the model

Usage

```
parameters(x)
```

Arguments

x A psychometrics model.

Value

Invisibly returns a data frame containing information on all parameters.

Author(s)

Sacha Epskamp

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
```

```

select(A1:A5, gender) %>%
na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars, omega = "zero")

# Run model:
mod <- mod %>% runmodel

# Parameter estimates:
mod %>% parameters

```

parequal

Set equality constraints across parameters

Description

This function can be used to set parameters equal

Usage

```
parequal(x, ..., inds = integer(0), verbose, log = TRUE,
runmodel = FALSE)
```

Arguments

x	A psychometrics model.
...	Arguments sent to runmodel
inds	Parameter indices of parameters to be constrained equal
verbose	Logical, should messages be printed?
log	Logical, should the log be updated?
runmodel	Logical, should the model be updated?

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

 partialprune

Partial pruning of multi-group models

Description

This function will search for a multi-group model with equality constraints on some but not all parameters. This is called partial pruning (Epskamp, Isvoranu, & Cheung, 2020; Haslbeck, 2020). The algorithm is as follows: 1. remove all parameters not significant at alpha in all groups (without equality constraints), 2. create a union model with all parameters included in any group included in all groups and constrained equal. 3. Stepwise free equality constraints of the parameter that features the largest sum of modification indices until BIC can no longer be improved. 4. Select and return (by default) the best model according to BIC (original model, pruned model, union model and partially pruned model).

Usage

```
partialprune(x, alpha = 0.01, matrices, verbose,
             combinefun = c("unionmodel", "intersectionmodel", "identity"),
             return = c("partialprune", "best", "union_equal", "prune"),
             criterion = "bic", best = c("lowest", "highest"),
             final_prune = c("saturated", "partialprune"), ...)
```

Arguments

x	A psychometrics model.
alpha	Significance level to use.
matrices	Vector of strings indicating which matrices should be pruned. Will default to network structures.
verbose	Logical, should messages be printed?
combinefun	Function used to combine models of different groups.
return	What model to return? "best" for best fitting model (according to BIC), "partialprune" for the partialpruned model, "union_equal" for the union model with equality constraints, and "prune" for the originally pruned model without equality constraints.
best	Should the lowest or the highest index of criterion be used to select the final model?
criterion	What criterion to use for the model selection in the last step? Defaults to "bic" for BIC selection.
final_prune	Should the last prune step be based on removing edges not significant in the last model in the partialprune algorithm or the first model (saturated model) in the algorithm? Defaults to "saturated". Set to "partialprune" to mimic psychometrics < 0.13.1 behavior.
...	Arguments sent to prune .

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

References

Epskamp, S., Isvoranu, A. M., & Cheung, M. (2020). Meta-analytic gaussian network aggregation. PsyArXiv preprint. DOI:10.31234/osf.io/236w8.

Haslbeck, J. (2020). Estimating Group Differences in Network Models using Moderation Analysis. PsyArXiv preprint. DOI:10.31234/osf.io/926pv.

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the extroversion items, and gender:
ExData <- bfi %>%
  select(E1:E5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML)

# Define variables:
vars <- names(ExData)[1:5]

# Saturated estimation:
mod_saturated <- ggm(ExData,
  vars = vars,
  groups = "gender")

# Partial prune model:
mod_partial <- mod_saturated
partialprune

# Obtain the networks:
getmatrix(mod_partial, "omega")

# Differences:
getmatrix(mod_partial, "omega")[[1]] -
  getmatrix(mod_partial, "omega")[[2]]

# Difference detected in edge 4 - 5
```

penalize

*Penalized Maximum Likelihood Functions***Description**

Functions for penalized maximum likelihood (PML) and penalized full-information maximum likelihood (PFIML) estimation. `penalize` adds an L1 (LASSO) or elastic net penalty to specified model parameters. `unpenalize` removes the penalty. `penaltyVector` extracts the per-free-parameter penalty strengths. `refit` performs post-selection inference by fixing penalized zeros and re-running with standard ML (or FIML for PFIML) estimation to obtain standard errors.

Usage

```
penalize(x, matrix, row, col, lambda, group, verbose, log = TRUE)
unpenalize(x, matrix, row, col, group, verbose, log = TRUE)
penaltyVector(x)
refit(x, threshold = 1e-8, verbose, log = TRUE, ...)
```

Arguments

<code>x</code>	A psychometrics model.
<code>matrix</code>	Character vector of matrix name(s) to penalize/unpenalize. If missing, defaults to the matrices returned by <code>defaultPenalizeMatrices(x)</code> for <code>penalize</code> , or all currently penalized matrices for <code>unpenalize</code> .
<code>row</code>	Integer or character indicating specific row(s) of the matrix. If missing, the entire matrix is penalized/unpenalized.
<code>col</code>	Integer or character indicating specific column(s) of the matrix. If missing, the entire matrix is penalized/unpenalized.
<code>lambda</code>	Numeric, the penalty strength per parameter. If missing, uses the global <code>x@penalty\$lambda</code> .
<code>group</code>	Integer indicating specific group(s). If missing, all groups are included.
<code>verbose</code>	Logical, should messages be printed?
<code>log</code>	Logical, should the log be updated?
<code>threshold</code>	For <code>refit</code> : threshold below which penalized parameters are considered zero and fixed.
<code>...</code>	Additional arguments passed to <code>runmodel</code> in <code>refit</code> .

Details

Penalized ML estimation uses the "PML" estimator, which wraps the standard ML fit function and gradient with an additional penalty term. For models with missing data, the "PFIML" estimator wraps the FIML fit function and gradient with the same penalty. The ISTA (Iterative Shrinkage-Thresholding Algorithm) proximal gradient optimizer is used automatically for both PML and PFIML models.

For symmetric matrices (e.g., ω), only off-diagonal elements are penalized by default. For non-symmetric matrices (e.g., β), all elements are penalized.

The `refit` function is used for post-selection inference: after a penalized model is estimated, `refit` fixes parameters that were shrunk to zero and re-estimates the model with standard ML (for PML) or FIML (for PFIML) to obtain valid standard errors and fit indices.

Model constructors (e.g., `ggm`, `lvm`, `var1`) accept `penalty_lambda`, `penalty_alpha`, and `penalize_matrices` arguments for convenient PML/PFIML setup. When `missing = "auto"` (the default), specifying `estimator = "PML"` with missing data will automatically switch to "PFIML".

Value

`penalize`, `unpenalize`, and `refit` return an object of class `psychometrics` ([psychometrics-class](#)). `penaltyVector` returns a numeric vector of penalty strengths per free parameter.

Author(s)

Sacha Epskamp

See Also

[find_penalized_lambda](#) for automatic lambda selection via EBIC grid search, [runmodel](#) for running models (calls `find_penalized_lambda` automatically when `penalty_lambda = NA`), [ggm](#) and other model constructors for the `penalty_lambda`, `penalty_alpha`, and `penalize_matrices` arguments.

Examples

```
# Load bfi data from psych package:
library("psychTools")
library("dplyr")
data(bfi)

ConsData <- bfi %>%
  select(A1:A5) %>%
  na.omit

vars <- names(ConsData)

# Penalized GGM estimation with manual lambda:
mod <- ggm(ConsData, vars = vars, estimator = "PML", penalty_lambda = 0.1)
mod <- mod %>% runmodel

# Post-selection refit for standard errors:
mod_refit <- mod %>% refit
mod_refit %>% parameters

# Manual unpenalize: free a specific edge from the penalty:
mod2 <- ggm(ConsData, vars = vars, estimator = "PML", penalty_lambda = 0.1)
mod2 <- unpenalize(mod2, "omega", row = 1, col = 2)
mod2 <- mod2 %>% runmodel
```

prune

*Stepdown model search by pruning non-significant parameters.***Description**

This function will (recursively) remove parameters that are not significant and refit the model.

Usage

```
prune(x, alpha = 0.01, adjust = c("none", "holm",
  "hochberg", "hommel", "bonferroni", "BH", "BY",
  "fdr"), matrices, runmodel = TRUE, recursive = FALSE,
  verbose, log = TRUE, identify = TRUE, startreduce = 1,
  limit = Inf, mode = c("tested","all"), ...)
```

Arguments

<code>x</code>	A psychometrics model.
<code>alpha</code>	Significance level to use.
<code>adjust</code>	p-value adjustment method to use. See <code>p.adjust</code> .
<code>matrices</code>	Vector of strings indicating which matrices should be pruned. Will default to network structures.
<code>runmodel</code>	Logical, should the model be evaluated after pruning?
<code>recursive</code>	Logical, should the pruning process be repeated?
<code>verbose</code>	Logical, should messages be printed?
<code>log</code>	Logical, should the log be updated?
<code>identify</code>	Logical, should models be identified automatically?
<code>startreduce</code>	A numeric value indicating a factor with which the starting values should be reduced. Can be useful when encountering numeric problems.
<code>limit</code>	The maximum number of parameters to be pruned.
<code>mode</code>	Mode for adjusting for multiple comparisons. Should all parameters be considered as the total number of tests or only the tested parameters (parameters of interest)?
<code>...</code>	Arguments sent to <code>runmodel</code>

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

See Also[stepup](#)**Examples**

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars, omega = "full")

# Run model:
mod <- mod %>% runmodel

# Prune model:
mod <- mod %>% prune(adjust = "fdr", recursive = FALSE)
```

psychometrics-class *Class "psychometrics"*

Description

Main class for psychometrics results.

Objects from the Class

Objects can be created by calls of the form `new("psychometrics", ...)`.

Slots

model: Object of class "character" ~~
submodel: Object of class "character" ~~
parameters: Object of class "data.frame" ~~
matrices: Object of class "data.frame" ~~
meanstructure: Object of class "logical" ~~

```
computed: Object of class "logical" ~~
sample: Object of class "psychonetrics_samplestats" ~~
modelmatrices: Object of class "list" ~~
log: Object of class "psychonetrics_log" ~~
optim: Object of class "list" ~~
fitmeasures: Object of class "list" ~~
baseline_saturated: Object of class "list" ~~
equal: Object of class "character" ~~
objective: Object of class "numeric" ~~
information: Object of class "matrix" ~~
identification: Object of class "character" ~~
optimizer: Object of class "character" ~~
optim.args: Object of class "list" ~~
estimator: Object of class "character" ~~
distribution: Object of class "character" ~~
extramatrices: Object of class "list" ~~
rawts: Object of class "logical" ~~
Drawts: Object of class "list" ~~
types: Object of class "list" ~~
cpp: Object of class "logical" ~~
verbose: Object of class "logical" ~~
```

Methods

```
resid signature(object = "psychonetrics"): ...
residuals signature(object = "psychonetrics"): ...
show signature(object = "psychonetrics"): ...
```

Author(s)

Sacha Epskamp

Examples

```
showClass("psychonetrics")
```

```
psychonetrics_bootstrap-class  
  Class "psychonetrics_bootstrap"
```

Description

Class for aggregated bootstrap results.

Objects from the Class

Objects can be created by calls of the form `new("psychonetrics_bootstrap", ...)`.

Slots

```
model: Object of class "character" ~~  
submodel: Object of class "character" ~~  
parameters: Object of class "data.frame" ~~  
models: Object of class "list" ~~  
matrices: Object of class "data.frame" ~~  
fitmeasures: Object of class "data.frame" ~~  
distribution: Object of class "character" ~~  
verbose: Object of class "logical" ~~  
boot_sub: Object of class "numeric" ~~  
boot_resample: Object of class "logical" ~~  
n_fail: Object of class "numeric" ~~  
n_success: Object of class "numeric" ~~  
types: Object of class "list" ~~
```

Methods

```
show signature(object = "psychonetrics_bootstrap"): ...
```

Author(s)

Sacha Epskamp

Examples

```
showClass("psychonetrics_bootstrap")
```

psychonetrics_log-class
Class "psychonetrics"

Description

A logbook entry in the psychonetrics logbook

Objects from the Class

Objects can be created by calls of the form `new("psychonetrics_log", ...)`.

Slots

event: Object of class "character" ~~

time: Object of class "POSIXct" ~~

sessionInfo: Object of class "sessionInfo" ~~

Methods

`show` signature(object = "psychonetrics_log"): ...

Author(s)

Sacha Epskamp

Examples

```
showClass("psychonetrics_log")
```

psychonetrics_update *Model updating functions*

Description

These functions update a psychonetrics model. Typically they are not required.

Usage

```

addMIs(x, matrices = "all", type = c("normal", "free",
                                     "equal"), verbose, analyticFisher = TRUE)

addSEs(x, verbose, approximate_SEs = FALSE)

addfit(x, verbose)

identify(x)

```

Arguments

x	A psychometrics model.
matrices	Optional vector of matrices to include in MIs.
type	String indicating which modification indices should be updated.
verbose	Logical, should messages be printed?
analyticFisher	Logical indicating if an analytic Fisher information matrix should be used.
approximate_SEs	Logical, should standard errors be approximated? If true, an approximate matrix inverse of the Fischer information is used to obtain the standard errors.

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

ri_clpm	<i>Random intercept cross-lagged panel models</i>
---------	---

Description

Function to run random intercept cross-lagged panel models under the lvm framework.

Usage

```

ri_clpm(data, vars, lambda,
        type = c("cov", "chol", "prec", "ggm"),
        verbose = FALSE, ...)
ri_clpm_stationary(x,
                  stationary = c("intercepts",
                                "contemporaneous",
                                "innovation",
                                "temporal"))

```

Arguments

x	A psychometrics model.
stationary	The part of the model to implement stationarity on.
data	A data frame encoding the data used in the analysis. Can be missing if covs and nobs are supplied.
vars	Required argument. Different from in other psychometrics models, this must be a <i>*matrix*</i> with each row indicating a variable and each column indicating a measurement. The matrix must be filled with names of the variables in the dataset corresponding to variable i at wave j. NAs can be used to indicate missing waves. The rownames of this matrix will be used as variable names.
lambda	A model matrix encoding the factor loading structure. Each row indicates an indicator and each column a latent. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
type	The type of model to model innovation
verbose	Logical, should progress be printed to the console?
...	Arguments sent to lvm

Value

A single psychometrics object

Author(s)

Sacha Epskamp

Examples

```
# to be made
```

```
runmodel
```

```
Run a psychometrics model
```

Description

This is the main function used to run a psychometrics model.

Usage

```
runmodel(x, level = c("gradient", "fitfunction"), addfit =
  TRUE, addMIs = TRUE, addSEs = TRUE, addInformation =
  TRUE, log = TRUE, verbose, optim.control,
  analyticFisher = TRUE, warn_improper = FALSE,
  warn_gradient = TRUE, warn_bounds = TRUE,
  return_improper = TRUE, bounded = TRUE,
  approximate_SEs = FALSE,
  criterion = "ebic.5",
  beta_min = c("numerical", "theoretical"))
```

Arguments

x	A psychometrics model.
level	Level at which the model should be estimated. Defaults to "gradient" to indicate the analytic gradient should be used.
addfit	Logical, should fit measures be added?
addMIs	Logical, should modification indices be added?
addSEs	Logical, should standard errors be added?
addInformation	Logical, should the Fisher information be added?
log	Logical, should the log be updated?
verbose	Logical, should messages be printed?
optim.control	A list with options for optimr
analyticFisher	Logical, should the analytic Fisher information be used? If FALSE, numeric information is used instead.
return_improper	Should a result in which improper computation was used be return? Improper computation can mean that a pseudoinverse of small spectral shift was used in computing the inverse of a matrix.
warn_improper	Logical. Should a warning be given when at some point in the estimation a pseudoinverse was used?
warn_gradient	Logical. Should a warning be given when the average absolute gradient is > 1?
bounded	Logical. Should bounded estimation be used (e.g., variances should be positive)?
approximate_SEs	Logical, should standard errors be approximated? If true, an approximate matrix inverse of the Fischer information is used to obtain the standard errors.
warn_bounds	Should a warning be given when a parameter is estimated near its bounds?
criterion	Character string specifying the information criterion for automatic lambda selection in PML/PFIML models. One of "bic", "ebic.25", "ebic.5" (default), "ebic.75", or "ebic1". Only used when the model contains auto-select penalty parameters (penalty_lambda = NA). See find_penalized_lambda for details.

`beta_min` Threshold for zeroing out small penalized parameters during automatic lambda selection. Can be "numerical" (default; uses $1e-05$), "theoretical" (uses $\sqrt{\log(p)/n}$), or a numeric value. See [find_penalized_lambda](#) for details.

Details

For penalized models (PML or PFIML) with auto-select penalty parameters (`penalty_lambda = NA`, the default), `runmodel` automatically calls [find_penalized_lambda](#) to select the optimal penalty strength via EBIC grid search before fitting. The `criterion` and `beta_min` arguments control this search. After the lambda search, use [refit](#) for post-selection inference with standard errors.

Value

An object of the class `psychometrics` ([psychometrics-class](#))

Author(s)

Sacha Epskamp

See Also

[find_penalized_lambda](#) for manual control over the lambda search, [penalize](#) for setting penalty parameters, [refit](#) for post-selection inference after penalized estimation.

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars, omega = "full")

# Run model:
mod <- mod %>% runmodel
```

setestimator	<i>Convenience functions</i>
--------------	------------------------------

Description

These functions can be used to change some estimator options.

Usage

```
setestimator(x, estimator)
```

```
setoptimizer(x, optimizer = c("default", "nlminb", "ucminf",
                              "nloptr_TNEWTON", "LBFGS++"), optim.args)
```

```
usecpp(x, use = TRUE)
```

Arguments

`x` A psychometrics model.

`estimator` A string indicating the estimator to be used

`optimizer` The optimizer to be used. The following optimizers are available:

R-based optimizers:

"nlminb" Port trust-region Newton-like optimizer from base R. Uses analytic gradients and a trust-region method that adaptively controls step sizes, making it very stable for SEM problems. This is the default and recommended optimizer for most models.

"ucminf" Unconstrained minimization using a quasi-Newton method (via the `optimr` package). Does not support box constraints. Fast for unconstrained problems.

C++ based optimizer:

"LBFGS++" L-BFGS-B optimizer from the `LBFGSpp` library (Yixuan Qiu). This is a pure C++ implementation that computes the objective function and gradient in a single combined call, avoiding redundant computation via internal caching. Supports box constraints. Recommended when speed is important.

NLOpt-based optimizer (via `nloptr`):

"nloptr_TNEWTON" Preconditioned truncated Newton with restarts from the `NLOpt` library (via the `nloptr` package). Uses the `NLOPT_LD_TNEWTON_PRECOND_RESTART` algorithm, which builds a local quadratic model of the objective and solves the Newton system approximately using a preconditioned conjugate-gradient method. Supports box constraints and uses analytic gradients. Can handle large parameter spaces efficiently. See Dembo & Steihaug (1982) for the underlying method.

	Defaults to "nlminb".
use	Logical indicating if C++ should be used (currently only used in FIML)
optim.args	List of arguments to sent to the optimizer.

Details

The default optimizer is nlminb with the following arguments:

- eval.max=20000L
- iter.max=10000L
- trace=0L
- abs.tol=sqrt(.Machine\$double.eps)
- rel.tol=sqrt(.Machine\$double.eps)
- step.min=1.0
- step.max=1.0
- x.tol=1.5e-8
- xf.tol=2.2e-14

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

setverbose	<i>Should messages of computation progress be printed?</i>
------------	--

Description

This function controls if messages should be printed for a psychometrics model.

Usage

```
setverbose(x, verbose = TRUE)
```

Arguments

x	A psychometrics model.
verbose	Logical indicating if verbose should be enabled

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

`simplestructure`*Generate factor loadings matrix with simple structure*

Description

This function generates the input for lambda arguments in latent variable models using a simple structure. The input is a vector with an element for each variable indicating the factor the variable loads on.

Usage`simplestructure(x)`**Arguments**

`x` A vector indicating which factor each indicator loads on.

Author(s)

Sacha Epskamp <mail@sachaepskamp.com>

`StarWars`*Star Wars dataset*

Description

This questionnaire was constructed by Carolin Katzera, Charlotte Tanis, Esther Niehoff, Myrthe Veenman, and Jason Nak as part of an assignment for a course on confirmatory factor analysis (<http://sachaepskamp.com/SEM2018>). They also collected the data among fellow psychology students as well as through social media.

Usage`data("StarWars")`

Format

A data frame with 271 observations on the following 13 variables.

- Q1 I am a huge Star Wars fan! (star what?)
- Q2 I would trust this person with my democracy <picture of Jar Jar Binks>
- Q3 I enjoyed the story of Anakin's early life
- Q4 The special effects in this scene are awful <video of the Battle of Geonosis>
- Q5 I would trust this person with my life <picture of Han Solo>
- Q6 I found Darth Vader's big reveal in "Empire" one of the greatest moments in movie history
- Q7 The special effects in this scene are amazing <video of the Death Star explosion>
- Q8 If possible, I would definitely buy this droid <picture of BB-8>
- Q9 The story in the Star Wars sequels is an improvement to the previous movies
- Q10 The special effects in this scene are marvellous <video of the Starkiller Base firing>
- Q11 What is your gender?
- Q12 How old are you?
- Q13 Have you seen any of the Star Wars movies?

Details

The questionnaire is online at https://github.com/SachaEpskamp/SEM-code-examples/blob/master/CFA_fit_examples/StarWars. The authors of the questionnaire defined a measurement model before collecting data: Q2 - Q4 are expected to load on a "prequel" factor, Q5 - Q7 are expected to load in a "originals" factor, and Q8 - Q10 are expected to load on a "sequal" factor. Finally, Q1 is expected to load on all three.

Source

https://github.com/SachaEpskamp/SEM-code-examples/blob/master/CFA_fit_examples

Examples

```
data(StarWars)
```

stepup

Stepup model search along modification indices

Description

This function automatically performs step-up search by adding the parameter with the largest modification index until some criterion is reached or no modification indices are significant at alpha.

Usage

```
stepup(x, alpha = 0.01, criterion = "bic", matrices, mi =
      c("mi", "mi_free", "mi_equal"), greedyadjust =
      c("bonferroni", "none", "holm", "hochberg", "hommel",
        "fdr", "BH", "BY"), stopif, greedy = FALSE, verbose,
      checkinformation = TRUE, singularinformation =
      c("tryfix", "skip", "continue", "stop"), startEPC =
      TRUE, ...)
```

Arguments

x	A psychometrics model.
alpha	Significance level to use.
criterion	String indicating the criterion to minimize. Any criterion from fit can be used.
matrices	Vector of strings indicating which matrices should be searched. Will default to network structures and factor loadings.
mi	String indicating which kind of modification index should be used ("mi" is the typical MI, "mi_free" is the modification index free from equality constraints across groups, and "mi_equal" is the modification index if the parameter is added constrained equal across all groups).
greedyadjust	String indicating which p-value adjustment should be used in greedy start. Any method from p.adjust can be used.
stopif	An R expression, using objects from fit , which will break stepup search if it evaluates to TRUE. For example, <code>stopif = rmsea < 0.05</code> will lead to search to stop if rmsea is below 0.05.
greedy	Logical, should a greedy start be used? If TRUE, the first step adds any parameter that is significant (after adjustment)
verbose	Logical, should messages be printed?
checkinformation	Logical, should the Fisher information be checked for potentially non-identified models?
singularinformation	String indicating how to proceed if the information matrix is singular. "tryfix" will adjust starting values to try to fix the problem, "skip" will lead to the algorithm to skip the current parameter, "continue" will ignore the situation, and "stop" will break the algorithm and return a list with the last two models.
startEPC	Logical, should the starting value be set at the expected parameter change?
...	Arguments sent to runmodel

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

See Also[prune](#)**Examples**

```

# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML)

# Define variables:
vars <- names(ConsData)[1:5]

# Let's fit a full GGM:
mod <- ggm(ConsData, vars = vars, omega = "full")

# Run model:
mod <- mod %>%runmodel %>%prune(alpha = 0.05)

# Remove an edge (example):
mod <- mod %>%fixpar("omega",1,2) %>%runmodel

# Stepup search
mod <- mod %>%stepup(alpha = 0.05)

```

transmod*Transform between model types*

Description

This function allows to transform a model variance–covariance structure from one type to another. Its main uses are to (1) use a Cholesky decomposition to estimate a saturated covariance matrix or GGM, and (2) to transform between conditional (ggm) and marginal associations (cov).

Usage

```

transmod(x, ..., verbose, keep_computed = FALSE, log = TRUE,
         identify = TRUE)

```

Arguments

x	A psychometrics model
...	Named arguments with the new types to use (e.g., between = "ggm" or y = "cov")
verbose	Logical, should messages be printed?
keep_computed	Logical, should the model be stated to be uncomputed adter the transformation? In general, a model does not need to be re-computed as transformed parameters should be at the maximum likelihood estimate.
log	Logical, should a logbook entry be made?
identify	Logical, should the model be identified after transforming?

Details

Transformations are only possible if the model is diagonal (e.g., no partial correlations) or saturated (e.g., all covariances included).

Author(s)

Sacha Epskamp

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (otherwise use Estimator = "FIML")

# Define variables:
vars <- names(ConsData)[1:5]

# Model with Cholesky decompositon:
mod <- varcov(ConsData, vars = vars, type = "chol")

# Run model:
mod <- mod %>% runmodel

# Transform to GGM:
mod_trans <- transmod(mod, type = "ggm") %>% runmodel
# Note: runmodel often not needed

# Obtain thresholded GGM:
getmatrix(mod_trans, "omega", threshold = TRUE)
```

tsdlvm1	<i>Lag-1 dynamic latent variable model family of psychometrics models for time-series data</i>
---------	--

Description

This is the family of models that models a dynamic factor model on time-series. There are two covariance structures that can be modeled in different ways: contemporaneous for the contemporaneous model and residual for the residual model. These can be set to "cov" for covariances, "prec" for a precision matrix, "ggm" for a Gaussian graphical model and "chol" for a Cholesky decomposition. The `ts_lvlgvar` wrapper function sets contemporaneous = "ggm" for the graphical VAR model.

Usage

```
tsdlvm1(data, lambda, contemporaneous = c("cov", "chol",
    "prec", "ggm"), residual = c("cov", "chol", "prec",
    "ggm"), beta = "full", omega_zeta = "full", delta_zeta
    = "diag", kappa_zeta = "full", sigma_zeta = "full",
    lowertri_zeta = "full", omega_epsilon = "zero",
    delta_epsilon = "diag", kappa_epsilon = "diag",
    sigma_epsilon = "diag", lowertri_epsilon = "diag", nu,
    mu_eta, identify = TRUE, identification =
    c("loadings", "variance"), latents, beepvar, dayvar,
    idvar, vars, groups, groupvar, covs, cors,
    means, nob, corinput, missing =
    "auto", equal = "none", baseline_saturated = TRUE,
    estimator = "ML", optimizer, storedata = FALSE,
    sampleStats, covtype = c("choose", "ML", "UB"),
    centerWithin = FALSE, standardize = c("none", "z",
    "quantile"), verbose = FALSE, bootstrap = FALSE,
    boot_sub, boot_resample, penalty_lambda = NA,
    penalty_alpha = 1, penalize_matrices)
```

```
ts_lvlgvar(...)
```

Arguments

data	A data frame encoding the data used in the analysis. Can be missing if covs and nob are supplied.
lambda	A model matrix encoding the factor loading structure. Each row indicates an indicator and each column a latent. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
contemporaneous	The type of contemporaneous model used. See description.

residual	The type of residual model used. See description.
beta	A model matrix encoding the temporal relationships (transpose of temporal network) between latent variables. A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix. Can also be "full" for a full temporal network or "zero" for an empty temporal network.
omega_zeta	Only used when contemporaneous = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_zeta	Only used when contemporaneous = "ggm". Either "diag" or "zero" (not recommended), or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_zeta	Only used when contemporaneous = "prec". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
sigma_zeta	Only used when contemporaneous = "cov". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_zeta	Only used when contemporaneous = "chol". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega_epsilon	Only used when residual = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_epsilon	Only used when residual = "ggm". Either "diag" or "zero" (not recommended), or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.

<code>kappa_epsilon</code>	Only used when <code>residual = "prec"</code> . Either <code>"full"</code> to estimate every element freely, <code>"diag"</code> to only include diagonal elements, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>sigma_epsilon</code>	Only used when <code>residual = "cov"</code> . Either <code>"full"</code> to estimate every element freely, <code>"diag"</code> to only include diagonal elements, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>lowertri_epsilon</code>	Only used when <code>residual = "chol"</code> . Either <code>"full"</code> to estimate every element freely, <code>"diag"</code> to only include diagonal elements, or a matrix of the dimensions <code>node x node</code> with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
<code>nu</code>	Optional vector encoding the intercepts of the observed variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
<code>mu_eta</code>	Optional vector encoding the means of the latent variables. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free intercepts, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
<code>identify</code>	Logical, should the model be automatically identified?
<code>identification</code>	Type of identification used. <code>"loadings"</code> to fix the first factor loadings to 1, and <code>"variance"</code> to fix the diagonal of the latent variable model matrix (<code>sigma_zeta</code> , <code>lowertri_zeta</code> , <code>delta_zeta</code> or <code>kappa_zeta</code>) to 1.
<code>latents</code>	An optional character vector with names of the latent variables.
<code>beepvar</code>	Optional string indicating assessment beep per day. Adding this argument will cause non-consecutive beeps to be treated as missing!
<code>dayvar</code>	Optional string indicating assessment day. Adding this argument makes sure that the first measurement of a day is not regressed on the last measurement of the previous day. IMPORTANT: only add this if the data has multiple observations per day.
<code>idvar</code>	Optional string indicating the subject ID
<code>vars</code>	An optional character vector encoding the variables used in the analysis. Must equal names of the dataset in data.
<code>groups</code>	Deprecated. Use <code>groupvar</code> instead. An optional string indicating the name of the group variable in data.
<code>groupvar</code>	An optional string indicating the name of the group variable in data. Replaces the deprecated <code>groups</code> argument; if both are supplied, <code>groupvar</code> takes precedence with a warning.

covs	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. Make sure covtype argument is set correctly to the type of covariances used.
cors	A sample correlation matrix, or a list/array of such matrices for multiple groups. When supplied, the matrix is treated as a covariance matrix with a warning (appropriate for standardized data). Requires nobs. Note: corinput = TRUE is not supported in tsdlvm1.
means	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
nobs	The number of observations used in covs and means, or a vector of such numbers of observations for multiple groups.
corinput	Logical. Not supported in tsdlvm1() and will produce an error if set to TRUE.
missing	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be "auto" (default) for automatic detection, "listwise" for listwise deletion, or "pairwise" for pairwise deletion. When "auto", the function checks for missing data and switches ML to FIML, PML to PFIML, or defaults to listwise for LS estimators.
equal	A character vector indicating which matrices should be constrained equal across groups.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
estimator	The estimator to be used. Currently implemented are "ML" for maximum likelihood estimation, "FIML" for full-information maximum likelihood estimation, "PML" for penalized maximum likelihood estimation, "PFIML" for penalized full-information maximum likelihood estimation, "ULS" for unweighted least squares estimation, "WLS" for weighted least squares estimation, and "DWLS" for diagonally weighted least squares estimation. When missing = "auto" (default), "ML" is automatically switched to "FIML" and "PML" to "PFIML" if missing data is detected.
optimizer	The optimizer to be used. Can be one of "nlminb" (the default R nlminb function), "ucminf" (from the optimr package), "nloptr_TNEWTON" (preconditioned truncated Newton via nloptr), and "LBFGS++" (pure C++ L-BFGS-B). Defaults to "nlminb".
storedata	Logical, should the raw data be stored? Needed for bootstrapping (see bootstrap).
standardize	Which standardization method should be used? "none" (default) for no standardization, "z" for z-scores, and "quantile" for a non-parametric transformation to the quantiles of the marginal standard normal distribution.
sampleStats	An optional sample statistics object. Mostly used internally.
centerWithin	Logical, should data be within-person centered?
covtype	If 'covs' is used, this is the type of covariance (maximum likelihood or unbiased) the input covariance matrix represents. Set to "ML" for maximum likelihood estimates (denominator n) and "UB" to unbiased estimates (denominator n-1). The default will try to find the type used, by investigating which is most likely to result from integer valued datasets.

verbose	Logical, should messages be printed?
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets boot_sub = 1 and boot_resample = TRUE) or "case" (which sets boot_sub = 0.75 and boot_resample = FALSE).
boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
penalty_lambda	Numeric penalty strength for penalized ML estimation (PML/PFIML). NA (default) triggers automatic selection via EBIC-based grid search when a penalized estimator is used; set to a specific numeric value to use a fixed penalty strength (0 = no penalty). See find_penalized_lambda and penalize .
penalty_alpha	Elastic net mixing parameter: 1 = LASSO (default), 0 = ridge.
penalize_matrices	Character vector of matrix names to penalize. If missing, defaults are selected based on the model type.
...	Arguments sent to tsdlvm1

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

Examples

```
# Note: this example is wrapped in a dontrun environment because the data is not
# available locally.
## Not run:
# Obtain the data from:
#
# Epskamp, S., van Borkulo, C. D., van der Veen, D. C., Servaas, M. N., Isvoranu, A. M.,
# Riese, H., & Cramer, A. O. (2018). Personalized network modeling in psychopathology:
# The importance of contemporaneous and temporal connections. Clinical Psychological
# Science, 6(3), 416-427.
#
# Available here: https://osf.io/c8wjz/
tsdata <- read.csv("Supplementary2_data.csv")

# Encode time variable in a way R understands:
tsdata$time <- as.POSIXct(tsdata$time, tz = "Europe/Amsterdam")

# Extract days:
tsdata$Day <- as.Date(tsdata$time, tz = "Europe/Amsterdam")

# Variables to use:
```

```

vars <- c("relaxed", "sad", "nervous", "concentration", "tired", "rumination",
         "bodily.discomfort")

# Create lambda matrix (in this case: one factor):
Lambda <- matrix(1,7,1)

# Estimate dynamical factor model:
model <- tsdlvm1(
  tsdata,
  lambda = Lambda,
  vars = vars,
  dayvar = "Day",
  estimator = "FIML"
)

# Run model:
model <- model %>% runmodel

# Look at fit:
model %>% print
model %>% fit # Pretty bad fit

## End(Not run)

```

unionmodel

Unify models across groups

Description

The unionmodel will add all parameters to all groups that are free in at least one group, and the intersectionmodel will constrain all parameters across groups to zero unless they are free to estimate in all groups.

Usage

```
unionmodel(x, runmodel = FALSE, verbose, log = TRUE, identify =
          TRUE, matrices, ...)
```

```
intersectionmodel(x, runmodel = FALSE, verbose, log = TRUE, identify =
                 TRUE, matrices, ...)
```

Arguments

x	A psychometrics model.
runmodel	Logical, should the model be updated?
verbose	Logical, should messages be printed?
log	Logical, should the log be updated?
identify	Logical, should the model be identified?

matrices Which matrices should be used to form the union/intersection model?
 ... Arguments sent to runmodel

Value

An object of the class psychometrics ([psychometrics-class](#))

Author(s)

Sacha Epskamp

var1 *Lag-1 vector autoregression family of psychometrics models*

Description

This is the family of models that models time-series data using a lag-1 vector autoregressive model (VAR; Epskamp, Waldorp, Mottus, Borsboom, 2018). The model is fitted to the Toeplitz matrix, but unlike typical SEM software the block of covariances of the lagged variables is not used in estimating the temporal and contemporaneous relationships (the block is modeled completely separately using a Cholesky decomposition, and does not enter the model otherwise). The contemporaneous argument can be used to define what contemporaneous model is used: contemporaneous = "cov" (default) models a variance-covariance matrix, contemporaneous = "chol" models a Cholesky decomposition, contemporaneous = "prec" models a precision matrix, and contemporaneous = "ggm" (alias: gvar()) models a Gaussian graphical model, also then known as a graphical VAR model.

Usage

```
var1(data, contemporaneous = c("cov", "chol", "prec",
  "ggm"), beta = "full", omega_zeta = "full", delta_zeta
  = "full", kappa_zeta = "full", sigma_zeta = "full",
  lowertri_zeta = "full", mu, beepvar, dayvar, idvar,
  vars, groups, groupvar, covs, cors, means,
  nobs, corinput, missing = "auto",
  equal = "none", baseline_saturated = TRUE, estimator =
  "ML", optimizer, storedata = FALSE, covtype =
  c("choose", "ML", "UB"), standardize = c("none", "z",
  "quantile"), sampleStats, verbose = FALSE, bootstrap =
  FALSE, boot_sub, boot_resample,
  penalty_lambda = NA, penalty_alpha = 1,
  penalize_matrices)

gvar(...)
```

Arguments

data	A data frame encoding the data used in the analysis. Can be missing if covs and nobs are supplied.
contemporaneous	The type of contemporaneous model used. See description.
beta	A model matrix encoding the temporal relationships (transpose of temporal network). A 0 encodes a fixed to zero element, a 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix. Can also be "full" for a full temporal network or "zero" for an empty temporal network.
omega_zeta	Only used when contemporaneous = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta_zeta	Only used when contemporaneous = "ggm". Either "diag" to estimate all scalings or "zero" (not recommended) to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
kappa_zeta	Only used when contemporaneous = "prec". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
sigma_zeta	Only used when contemporaneous = "cov". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri_zeta	Only used when contemporaneous = "chol". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
mu	Optional vector encoding the mean structure. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free means, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.

beepvar	Optional string indicating assessment beep per day. Adding this argument will cause non-consecutive beeps to be treated as missing!
dayvar	Optional string indicating assessment day. Adding this argument makes sure that the first measurement of a day is not regressed on the last measurement of the previous day. IMPORTANT: only add this if the data has multiple observations per day.
idvar	Optional string indicating the subject ID
vars	An optional character vector encoding the variables used in the analysis. Must equal names of the dataset in data.
groups	Deprecated. Use groupvar instead. An optional string indicating the name of the group variable in data.
groupvar	An optional string indicating the name of the group variable in data. Replaces the deprecated groups argument; if both are supplied, groupvar takes precedence with a warning.
covs	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. Make sure covtype argument is set correctly to the type of covariances used.
cors	A sample correlation matrix, or a list/array of such matrices for multiple groups. When supplied, the matrix is treated as a covariance matrix with a warning (appropriate for standardized data). Requires nobobs. Note: corinput = TRUE is not supported in var1.
means	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
nobs	The number of observations used in covs and means, or a vector of such numbers of observations for multiple groups.
corinput	Logical. Not supported in var1() and will produce an error if set to TRUE.
missing	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be "auto" (default) for automatic detection, "listwise" for listwise deletion, or "pairwise" for pairwise deletion. When "auto", the function checks for missing data and switches ML to FIML, PML to PFIML, or defaults to listwise for LS estimators.
equal	A character vector indicating which matrices should be constrained equal across groups.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
estimator	The estimator to be used. Currently implemented are "ML" for maximum likelihood estimation, "FIML" for full-information maximum likelihood estimation, "PML" for penalized maximum likelihood estimation, "PFIML" for penalized full-information maximum likelihood estimation, "ULS" for unweighted least squares estimation, "WLS" for weighted least squares estimation, and "DWLS" for diagonally weighted least squares estimation. When missing = "auto" (default), "ML" is automatically switched to "FIML" and "PML" to "PFIML" if missing data is detected.

optimizer	The optimizer to be used. Can be one of "nlminb" (the default R nlminb function), "ucminf" (from the optimr package), "nloptr_TNEWTON" (preconditioned truncated Newton via nloptr), and "LBFGS++" (pure C++ L-BFGS-B). Defaults to "nlminb".
storedata	Logical, should the raw data be stored? Needed for bootstrapping (see bootstrap).
standardize	Which standardization method should be used? "none" (default) for no standardization, "z" for z-scores, and "quantile" for a non-parametric transformation to the quantiles of the marginal standard normal distribution.
sampleStats	An optional sample statistics object. Mostly used internally.
covtype	If 'covs' is used, this is the type of covariance (maximum likelihood or unbiased) the input covariance matrix represents. Set to "ML" for maximum likelihood estimates (denominator n) and "UB" to unbiased estimates (denominator n-1). The default will try to find the type used, by investigating which is most likely to result from integer valued datasets.
verbose	Logical, should messages be printed?
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets boot_sub = 1 and boot_resample = TRUE) or "case" (which sets boot_sub = 0.75 and boot_resample = FALSE).
boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
penalty_lambda	Numeric penalty strength for penalized ML estimation (PML/PFIML). NA (default) triggers automatic selection via EBIC-based grid search when a penalized estimator is used; set to a specific numeric value to use a fixed penalty strength (0 = no penalty). See find_penalized_lambda and penalize .
penalty_alpha	Elastic net mixing parameter: 1 = LASSO (default), 0 = ridge.
penalize_matrices	Character vector of matrix names to penalize. If missing, defaults are selected based on the model type.
...	Arguments sent to var1

Details

This will be updated in a later version.

Value

An object of the class psychometrics

Author(s)

Sacha Epskamp

References

Epskamp, S., Waldorp, L. J., Mottus, R., & Borsboom, D. (2018). The Gaussian graphical model in cross-sectional and time-series data. *Multivariate Behavioral Research*, 53(4), 453-480.

See Also

[lvm](#), [varcov](#), [dlvm1](#)

Examples

```
library("dplyr")
library("graphicalVAR")

beta <- matrix(c(
  0,0.5,
  0.5,0
),2,2,byrow=TRUE)
kappa <- diag(2)
simData <- graphicalVARsim(50, beta, kappa)

# Form model:
model <- gvar(simData)

# Evaluate model:
model <- model %>% runmodel

# Parameter estimates:
model %>% parameters

# Plot the CIs:
CIplot(model, "beta")

# Note: this example is wrapped in a dontrun environment because the data is not
# available locally.
## Not run:
# Longer example:
#
# Obtain the data from:
#
# Epskamp, S., van Borkulo, C. D., van der Veen, D. C., Servaas, M. N., Isvoranu, A. M.,
# Riese, H., & Cramer, A. O. (2018). Personalized network modeling in psychopathology:
# The importance of contemporaneous and temporal connections. Clinical Psychological
# Science, 6(3), 416-427.
#
# Available here: https://osf.io/c8wjz/

tsdata <- read.csv("Supplementary2_data.csv")

# Encode time variable in a way R understands:
tsdata$time <- as.POSIXct(tsdata$time, tz = "Europe/Amsterdam")

# Extract days:
```

```

tsdata$Day <- as.Date(tsdata$time, tz = "Europe/Amsterdam")

# Variables to use:
vars <- c("relaxed", "sad", "nervous", "concentration", "tired", "rumination",
         "bodily.discomfort")

# Estimate, prune with FDR, and perform stepup search:
model_FDRprune <- gvar(
  tsdata,
  vars = vars,
  dayvar = "Day",
  estimator = "FIML"
) %>%
runmodel %>%
prune(adjust = "fdr", recursive = FALSE) %>%
stepup(criterion = "bic")

# Estimate with greedy stepup search:
model_stepup <- gvar(
  tsdata,
  vars = vars,
  dayvar = "Day",
  estimator = "FIML",
  omega_zeta = "zero",
  beta = "zero"
) %>%
runmodel %>%
stepup(greedy = TRUE, greedyadjust = "bonferroni", criterion = "bic")

# Compare models:
compare(
  FDRprune = model_FDRprune,
  stepup = model_stepup
)
# Very similar but not identical. Stepup is preferred here according to AIC and BIC

# Stepup results:
temporal <- getmatrix(model_stepup, "PDC") # PDC = Partial Directed Correlations
contemporaneous <- getmatrix(model_stepup, "omega_zeta")

# Average layout:
library("qgraph")
L <- averageLayout(temporal, contemporaneous)

# Labels:
labs <- gsub("\\.", "\\n", vars)

# Plot:
layout(t(1:2))
qgraph(temporal, layout = L, theme = "colorblind", directed=TRUE, diag=TRUE,
       title = "Temporal", vsize = 12, mar = rep(6,4), asize = 5,
       labels = labs)
qgraph(contemporaneous, layout = L, theme = "colorblind",

```

```

title = "Contemporaneous", vsize = 12, mar = rep(6,4), asize = 5,
labels = labs)

## End(Not run)

```

varcov

Variance-covariance family of psychometrics models

Description

This is the family of models that models only a variance-covariance matrix with mean structure. The type argument can be used to define what model is used: type = "cov" (default) models a variance-covariance matrix directly, type = "chol" (alias: cholesky()) models a Cholesky decomposition, type = "prec" (alias: precision()) models a precision matrix, type = "ggm" (alias: ggm()) models a Gaussian graphical model (Epskamp, Rhemtulla and Borsboom, 2017), and type = "cor" (alias: corr()) models a correlation matrix.

Usage

```

varcov(data, type = c("cov", "chol", "prec", "ggm", "cor"),
       sigma = "full", kappa = "full", omega = "full",
       lowertri = "full", delta = "diag", rho = "full", SD =
       "full", mu, tau, vars, ordered = character(0), groups,
       groupvar, covs, cors, means, nobs, missing = "auto", equal =
       "none", baseline_saturated = TRUE, estimator =
       "default", optimizer, storedata = FALSE, WLS.W,
       sampleStats, meanstructure, corinput, verbose = FALSE,
       covtype = c("choose", "ML", "UB"), standardize =
       c("none", "z", "quantile"), fullFIML = FALSE,
       bootstrap = FALSE, boot_sub, boot_resample,
       penalty_lambda = NA, penalty_alpha = 1,
       penalize_matrices)

cholesky(...)
precision(...)
prec(...)
ggm(...)
corr(...)

```

Arguments

data	A data frame encoding the data used in the analysis. Can be missing if covs and nobs are supplied.
type	The type of model used. See description.
sigma	Only used when type = "cov". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups,

	this argument can be a list or array with each element/slice encoding such a matrix.
kappa	Only used when type = "prec". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
omega	Only used when type = "ggm". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
lowertri	Only used when type = "chol". Either "full" to estimate every element freely, "diag" to only include diagonal elements, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
delta	Only used when type = "ggm". Either "diag" or "zero" (not recommended), or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
rho	Only used when type = "cor". Either "full" to estimate every element freely, "zero" to set all elements to zero, or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
SD	Only used when type = "cor". Either "diag" or "zero" (not recommended), or a matrix of the dimensions node x node with 0 encoding a fixed to zero element, 1 encoding a free to estimate element, and higher integers encoding equality constrains. For multiple groups, this argument can be a list or array with each element/slice encoding such a matrix.
mu	Optional vector encoding the mean structure. Set elements to 0 to indicate fixed to zero constrains, 1 to indicate free means, and higher integers to indicate equality constrains. For multiple groups, this argument can be a list or array with each element/column encoding such a vector.
tau	Optional list encoding the thresholds per variable.
vars	An optional character vector encoding the variables used in the analysis. Must equal names of the dataset in data.
groups	Deprecated. Use groupvar instead. An optional string indicating the name of the group variable in data.
groupvar	An optional string indicating the name of the group variable in data. Replaces the deprecated groups argument; if both are supplied, groupvar takes precedence with a warning.

covs	A sample variance–covariance matrix, or a list/array of such matrices for multiple groups. Make sure covtype argument is set correctly to the type of covariances used.
cors	A sample correlation matrix, or a list/array of such matrices for multiple groups. When supplied, corinput defaults to TRUE and the matrix is used in place of covs. Requires nobobs.
means	A vector of sample means, or a list/matrix containing such vectors for multiple groups.
nobs	The number of observations used in covs and means, or a vector of such numbers of observations for multiple groups.
covtype	If 'covs' is used, this is the type of covariance (maximum likelihood or unbiased) the input covariance matrix represents. Set to "ML" for maximum likelihood estimates (denominator n) and "UB" to unbiased estimates (denominator n-1). The default will try to find the type used, by investigating which is most likely to result from integer valued datasets.
missing	How should missingness be handled in computing the sample covariances and number of observations when data is used. Can be "auto" (default) for automatic detection, "listwise" for listwise deletion, or "pairwise" for pairwise deletion. When "auto", the function checks for missing data and switches ML to FIML, PML to PFIML, or defaults to listwise for LS estimators.
equal	A character vector indicating which matrices should be constrained equal across groups.
baseline_saturated	A logical indicating if the baseline and saturated model should be included. Mostly used internally and NOT Recommended to be used manually.
estimator	The estimator to be used. Currently implemented are "ML" for maximum likelihood estimation, "FIML" for full-information maximum likelihood estimation, "PML" for penalized maximum likelihood estimation, "PFIML" for penalized full-information maximum likelihood estimation, "ULS" for unweighted least squares estimation, "WLS" for weighted least squares estimation, and "DWLS" for diagonally weighted least squares estimation. "WLSMV" is accepted as a synonym for "DWLS" (both use DWLS estimation with a mean-and-variance adjusted scaled test statistic). When missing = "auto" (default), "ML" is automatically switched to "FIML" and "PML" to "PFIML" if missing data is detected. Defaults to "ML" for continuous data and "DWLS" when ordinal variables are specified via ordered.
optimizer	The optimizer to be used. Can be one of "nlminb" (the default R nlminb function), "ucminf" (from the optimr package), "nloptr_TNEWTON" (preconditioned truncated Newton via nloptr), and "LBFGS++" (pure C++ L-BFGS-B). Defaults to "nlminb".
storedata	Logical, should the raw data be stored? Needed for bootstrapping (see bootstrap).
standardize	Which standardization method should be used? "none" (default) for no standardization, "z" for z-scores, and "quantile" for a non-parametric transformation to the quantiles of the marginal standard normal distribution.
WLS.W	Optional WLS weights matrix.

sampleStats	An optional sample statistics object. Mostly used internally.
verbose	Logical, should progress be printed to the console?
ordered	A vector with strings indicating the variables that are ordered catagorical, or set to TRUE to model all variables as ordered catagorical.
meanstructure	Logical, should the meanstructure be modeled explicitly?
corinput	Logical, is the input a correlation matrix?
fullFIML	Logical, should row-wise FIML be used? Not recommended!
bootstrap	Should the data be bootstrapped? If TRUE the data are resampled and a bootstrap sample is created. These must be aggregated using aggregate_bootstraps! Can be TRUE or FALSE. Can also be "nonparametric" (which sets boot_sub = 1 and boot_resample = TRUE) or "case" (which sets boot_sub = 0.75 and boot_resample = FALSE).
boot_sub	Proportion of cases to be subsampled (round(boot_sub * N)).
boot_resample	Logical, should the bootstrap be with replacement (TRUE) or without replacement (FALSE)
penalty_lambda	Numeric penalty strength for penalized ML estimation (PML/PFIML). Default is NA, which triggers automatic lambda selection via EBIC grid search when estimator = "PML" or "PFIML" (see find_penalized_lambda). Set to a specific numeric value (e.g., 0.1) for manual lambda, or 0 for no penalty.
penalty_alpha	Elastic net mixing parameter: 1 = LASSO (default), 0 = ridge.
penalize_matrices	Character vector of matrix names to penalize. If missing, defaults are selected based on the model type.
...	Arguments sent to varcov

Details

The model used in this family is:

$$\text{var}(\mathbf{y}) = \Sigma$$

$$\mathcal{E}(\mathbf{y}) = \boldsymbol{\mu}$$

in which the covariance matrix can further be modeled in three ways. With type = "chol" as Cholesky decomposition:

$$\Sigma = \mathbf{L}\mathbf{L}^{\top},$$

with type = "prec" as Precision matrix:

$$\Sigma = \mathbf{K}^{-1},$$

and finally with type = "ggm" as Gaussian graphical model:

$$\Sigma = \mathbf{\Delta}(\mathbf{I} - \mathbf{\Omega})^{-1}\mathbf{\Delta}.$$

Value

An object of the class psychonetrics

Author(s)

Sacha Epskamp

References

Epskamp, S., Rhemtulla, M., & Borsboom, D. (2017). Generalized network psychometrics: Combining network and latent variable models. *Psychometrika*, 82(4), 904-927.

See Also

[lvm](#), [var1](#), [dlvm1](#), [penalize](#) for manual penalty control, [find_penalized_lambda](#) for automatic lambda selection, [refit](#) for post-selection inference after penalized estimation.

Examples

```
# Load bfi data from psych package:
library("psychTools")
data(bfi)

# Also load dplyr for the pipe operator:
library("dplyr")

# Let's take the agreeableness items, and gender:
ConsData <- bfi %>%
  select(A1:A5, gender) %>%
  na.omit # Let's remove missingness (or use missing = "auto" default which auto-selects FIML)

# Define variables:
vars <- names(ConsData)[1:5]

# Saturated estimation:
mod_saturated <- ggm(ConsData, vars = vars)

# Run the model:
mod_saturated <- mod_saturated %>% runmodel

# We can look at the parameters:
mod_saturated %>% parameters

# Labels:
labels <- c(
  "indifferent to the feelings of others",
  "inquire about others' well-being",
  "comfort others",
  "love children",
  "make people feel at ease")

# Plot CIs:
CIplot(mod_saturated, "omega", labels = labels, labelstart = 0.2)
```

```

# We can also fit an empty network:
mod0 <- ggm(ConsData, vars = vars, omega = "zero")

# Run the model:
mod0 <- mod0 %>% runmodel

# We can look at the modification indices:
mod0 %>% MIs

# To automatically add along modification indices, we can use stepup:
mod1 <- mod0 %>% stepup

# Let's also prune all non-significant edges to finish:
mod1 <- mod1 %>% prune

# Look at the fit:
mod1 %>% fit

# Compare to original (baseline) model:
compare(baseline = mod0, adjusted = mod1)

# We can also look at the parameters:
mod1 %>% parameters

# Or obtain the network as follows:
getmatrix(mod1, "omega")

# Penalized GGM estimation with automatic lambda selection:
mod_pml <- ggm(ConsData, vars = vars, estimator = "PML")
mod_pml <- mod_pml %>% runmodel

# Check selected lambda:
mod_pml@optim$lambda_search

# Obtain the sparse network:
getmatrix(mod_pml, "omega")

# Refit for standard errors and fit indices:
mod_pml_refit <- mod_pml %>% refit
mod_pml_refit %>% parameters

```

write_psychometrics *Write comprehensive model output to a text file*

Description

Writes a comprehensive plain-text output file containing all model information, similar to Mplus or LISREL output. The file includes model specification, sample information, parameter estimates, fit measures, model matrices, modification indices, and the model logbook. This file can be shared as supplementary material in publications.

Usage

```
write_psychometrics(x, file = "psychometrics_output.txt",  
                   matrices = TRUE, MIs = TRUE, logbook = TRUE)
```

Arguments

x	A psychometrics model.
file	Character string specifying the path to the output file. Defaults to "psychometrics_output.txt".
matrices	Logical. Should the full estimated model matrices be included in the output? Defaults to TRUE. Set to FALSE for more compact output.
MIs	Logical. Should modification indices be included? Only included if modification indices have been computed via addMIs . Defaults to TRUE.
logbook	Logical. Should the model logbook be included? Defaults to TRUE.

Value

Invisibly returns the file path of the written output.

Author(s)

Sacha Epskamp

Examples

```
library("dplyr")  
  
# Load data:  
data(StarWars)  
  
# Simple CFA:  
Lambda <- matrix(1, 4)  
mod <- lvm(StarWars, lambda = Lambda, vars = c("Q1", "Q5", "Q6", "Q7"),  
          identification = "variance", latents = "Originals")  
mod <- mod %>% runmodel %>% addfit  
  
# Write output:  
write_psychometrics(mod, file = tempfile(fileext = ".txt"))
```

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