Package 'simsem'

September 8, 2012

Type Package

Version 0.3-3

Date 2012-09-08

Title SIMulated Structural Equation Modeling.

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Suggests parallel, Amelia, copula, quantreg, splines, foreign, KernSmooth, semTools Description This package can be used to generate data using the structural equation modeling framework. This package is tailored to use those simulated data for various purposes, such as model fit evaluation, power analysis, or missing data handling and planning. License GPL (>= 2) LazyLoad yes URL http://www.simsem.org R topics documented: analyze anova bind bind bind bind clean cleanSimResult continuousPower 10
work. This package is tailored to use those simulated data for various purposes, such as model fit evaluation, power analysis, or missing data handling and planning. License GPL (>= 2) LazyLoad yes URL http://www.simsem.org R topics documented: analyze anova anova bind bindDist clean clean SimResult
LazyLoad yes URL http://www.simsem.org R topics documented: analyze anova bind bindDist clean cleanSimResult
URL http://www.simsem.org R topics documented: analyze 3 anova 4 bind 5 bindDist 6 clean 6 cleanSimResult 9
analyze 3 anova 4 bind 5 bindDist 8 clean 9 cleanSimResult 9
analyze
anova 4 bind 5 bindDist 8 clean 9 cleanSimResult 9
createData 11 draw 13 estmodel 15 extractLavaanFit 17 find2Dhist 18 findFactorIntercept 18 findFactorMean 19

2

findFactorResidualVar	20
indFactorTotalCov	21
indFactorTotalVar	
findIndIntercept	24
indIndMean	25
findIndResidualVar	
indIndTotalVar	
îndphist	
indPossibleFactorCor	
indPower	
indRecursiveSet	
indRowZero	
indTargetPower	
itMeasuresChi	
generate	
getCondQtile	
getCutoff	
getCutoffNested	
getCutoffNonNested	
getExtraOutput	
getKeywords	
getPopulation	
getPower	
getPowerFit	45
getPowerFitNested	47
getPowerFitNonNested	49
mposeMissing	51
nterpolate	
ikRatioFit	
oadingFromAlpha	
miss	
nodel	
nodel.lavaan	
nultipleAllEqual	
overlapHist	63
blot3DQtile	63 64
blotCutoff	. 03
blotCutoffNested	
blotCutoffNonNested	
plotDist	
plotIndividualScatter	
plotLogisticFit	
plotMisfit	
plotOverHist	73
plotPower	74
olotPowerFit	75
blotPowerFitDf	77
blotPowerFitNested	
plotPowerFitNonNested	
plotPowerSig	
blotQtile	
blotScatter	

analyze 3

analy	/ze	TBA																		
Index																				119
	whichMonotonic .		٠.	٠	 •	 	 •	 ٠	•	 •	 •	٠		٠	•	•	•		•	117
	validatePath																			
	validateObject																			
	validateCovariance																			
	twoTailedPValue .																			
	summaryShort																			
	summaryPopulation																			
	summaryParam																			
	summaryMisspec .																			
	summaryFit																			
	summaryConverge																			
	sortList																			
	SimVector-class																			
	SimSem-class																			
	SimResult-class																			
	SimMissing-class																			
	SimDataDist-class . SimMatrix-class																			
	sim																			
	setPopulation																			
	revText																			95
	pValueVariedCutoff																			94
	pValueNonNested .																			92
	pValueNested																			91
	pValueCondCutoff																			90
	pValue					 														88
	printIfNotNull					 														87
	predProb					 														86
	popMisfitMACS																			85
	popDiscrepancy					 														84

Description

TBA

Usage

```
analyze(model, data, package="lavaan", miss=NULL, indLab=NULL, aux=NULL, ...)
```

Arguments

model	TBA
data	TBA
package	TBA
miss	TBA

4 anova

```
indLab TBA aux TBA
```

Value

TBA

Author(s)

Examples

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
LY <- bind(loading, 0.7)

latent.cor <- matrix(NA, 2, 2)
diag(latent.cor) <- 1
RPS <- binds(latent.cor, 0.5)

RTE <- binds(diag(6))

VY <- bind(rep(NA,6),2)

CFA.Model <- model(LY = LY, RPS = RPS, RTE = RTE, modelType = "CFA")
dat <- generate(CFA.Model,200)
out <- analyze(CFA.Model,dat)</pre>
```

anova

Provide a comparison of nested models and nonnested models across replications

Description

This function will provide averages of model fit statistics and indices for nested models. It will also provide average differences of fit indices and power for likelihood ratio tests of nested models.

Arguments

object	SimResult object being described. Currently at least two objects must be included as arguments
• • •	any additional arguments, such as additional objects or for the function with result object

bind 5

Value

A data frame that provides the statistics described above from all parameters. For using with linkS4class{SimResult}, the result is a list with two or three elements:

- summary: Average of fit indices across all replications
- diff: Average of the differences in fit indices across all replications
- varyParam: The statistical power of chi-square difference test given values of varying parameters (such as sample size or percent missing)

Author(s)

Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>), Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

SimResult for the object input

Examples

```
loading1 <- matrix(0, 6, 1)</pre>
loading1[1:6, 1] <- NA
loading2 <- loading1</pre>
loading2[6,1] <- 0
LX1 <- bind(loading1, 0.7)
LX2 <- bind(loading2, 0.7)
RPH <- binds(diag(1))</pre>
RTD <- binds(diag(6))
CFA.Model1 <- model(LY = LX1, RPS = RPH, RTE = RTD, modelType="CFA")
CFA.Model2 <- model(LY = LX2, RPS = RPH, RTE = RTD, modelType="CFA")
# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
# Need to make sure that both simResult calls have the same seed!
Output1 <- sim(5, n=500, model=CFA.Model1, generate=CFA.Model1, seed=123567)
Output2 <- sim(5, n=500, model=CFA.Model2, generate=CFA.Model1, seed=123567)
anova(Output1, Output2)
Output1b <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model1, generate=CFA.Model1, seed=123567)
Output2b <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model2, generate=CFA.Model1, seed=123567)
anova(Output1b, Output2b)
```

bind

Specify matrices for Monte Carlo simulation of structural equation models

Description

Create SimMatrix or SimVector object that specifies

- 1. Pattern of fixed/freed parameters for analysis
- 2. Population parameter values for data generation

6 bind

3. Any model misspecification (true population parameter is different than the one specified) for these parameters.

Each matrix in the Lisrel-style notation is specified in this way (e.g. LY, PS, and TE) and is used to create a model analysis template and a data generation template for simulation through the model function.

Usage

```
bind(free = NULL, popParam = NULL, misspec = NULL, symmetric = FALSE)
binds(free = NULL, popParam = NULL, misspec = NULL, symmetric = TRUE)
```

Arguments

free

Required matrix or vector where each element represents a fixed or freed parameter used for analysis with structural equation models. Parameters can be freed by setting the corresponding element in the matrix to NA, and can be fixed by setting the value of the element to any number (e.g. 0). Parameters can be labeled using any character string. Any labeled parameter is considered to be free, and parameters with identical labels will be constrained to equality for analysis.

popParam

Optional matrix or vector of identical dimension to the free matrix whose elements contain population parameter values for data generation in simulation. For similutation, each free parameter requires a population parameter value, which is a quoted numeric value. Parameters that don't have population values are left as empty strings. Population parameters can also be drawn from a distribution. This is done by wrapping a call to create 1 value from an existing random generation function in quotes: e.g "runif(1,0,1)", "rnorm(1,0,.01)" Every replication in the simulation will draw a parameter value from this distribution. The function checks that what is quoted is valid R.

If a random population parameter is constrained to equality in the free matrix, each drawn population parameter value will be the same. More details on data generation is available in ?generate, ?createData, and ?draw.

To simplify the most common case, popParam can take 1 value or distribution and create a matrix or vector that assigns that population parameter or distribution to all freed parameters. These population values are used as starting values for analysis by default.

misspec

Optional matrix or vector of identical dimension to the free matrix whose elements contain population parameter values for specifying misspecification. Elements of the misspec matrix contain population parameters that are added to parameters that are fixed or have an existing population value. These parameters are also quoted numeric strings, and can optionally be drawn from distributions as described above. To simplify the most common case, misspec can take 1 value or distribution and create a matrix or vector that assigns that value or distribution to all previously specified fixed parameters. Details about misspecification are included the data generation functions.

symmetric

Set as TRUE if the matrix created is symmetric (RPS/PS, RTE/TE). The function binds can also be used, which defaults to symmetric = TRUE

Details

Bind is the first step in the bind -> model -> sim workflow of *simsem*, and this document outlines the user interface or language used to describe these simulations. This interface, while complex,

bind 7

enables a wide array of simulation specifications for structural equation models by building on LISREL-style parameter specifications.

In simulations supported by *simsem*, a given parameter may be either fixed or freed for analysis, but may optionally also have a population value or distribution for data generation, or a value or distribution of misspecification. The purpose of bind is to stack these multiple meanings of a parameter into an object recognized by *simsem*, a SimMatrix. Each matrix in the Lisrel notation (e.g. LY, PS, TE, BE) becomes a SimMatrix, and is passed to the function model, which builds the data generation template and an analysis template (a lavaan parameter table), collectively forming a SimSem object, which can be passed to the function sim for simulation.

Value

SimMatrix or SimVector object that used for model specification for analysis and data generation in simsem.

Author(s)

See Also

- model To combine simMatrix objects into a complete data analysis and data generation template, which is a SimSem object
- generate To generate data using the simsem template.
- analyze To analyze real or generated data using the simsem template.

```
loading <- matrix(0, 6, 2)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loadingValues <- matrix(0, 6, 2)</pre>
loadingValues[1:3, 1] <- 0.7</pre>
loadingValues[4:6, 2] <- 0.7</pre>
LY <- bind(loading, loadingValues)
summary(LY)
# Set both factor correlations to .05
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPS <- binds(latent.cor, 0.5)</pre>
# Misspecify all error covarainces
error.cor <- matrix(0, 6, 6)
diag(error.cor) <- NA</pre>
RTE <- binds(error.cor,1,"runif(1,-.05,.05)")</pre>
```

8 bindDist

bindDist	Create a data distribution object.

Description

Create a data distribution object

Usage

```
bindDist(margins, ..., p = NULL, keepScale = TRUE, reverse = FALSE)
```

Arguments

margins	A character vector specifying all the marginal distributions. See the description of margins attribute of the Mvdc function for further details.
	A list whose each component is a list of named components, giving the parameter values of the marginal distributions. See the description of paramMargins attribute of the Mvdc function for further details.
p	Number of variables. If only one distribution object is listed, the p will make the same distribution objects for all variables.
keepScale	A vector representing whether each variable is transformed its mean and standard deviation or not. If TRUE, transform back to retain the mean and standard deviation of a variable equal to the model implied mean and standard deviation (with sampling error)
reverse	A vector representing whether each variable is mirrored or not. If TRUE, reverse the distribution of a variable (e.g., from positive skewed to negative skewed. If one logical value is specified, it will apply to all variables.

Value

SimDataDist that saves analysis result from simulate data.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimResult for the type of resulting object

```
d1 <- list(df=2)
d2 <- list(df=3)
d3 <- list(df=4)
d4 <- list(df=5)
d5 <- list(df=3)
d6 <- list(df=4)
d7 <- list(df=5)
d8 <- list(df=6)</pre>
dist <- bindDist(c(rep("t", 4), rep("chisq", 8)), d1, d2, d3, d4, d5, d6, d7, d8, d5, d6, d7, d8)
```

clean 9

clean

Extract only converged replications in the result objects

Description

Extract only the replications that are convergent in all supplied result objects (SimResult)

Usage

```
clean(...)
```

Arguments

... The target result objects (SimResult)

Value

The cleaned result objects

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

No example

cleanSimResult

Extract only converged replications in the result object

Description

Extract only the replications that are convergeent in a result object (SimResult)

Usage

```
cleanSimResult(object, converged=NULL)
```

Arguments

object The target result object (SimResult)

converged The replications to be extracted. If NULL, the converged slot in the result object

will be used

Value

The cleaned result object

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

10 continuousPower

Examples

No example

continuousPower Find power of model parameters when simulations have randomly varying parameters

Description

A function to find the power of parameters in a model when one or more of the simulations parameters vary randomly across replications.

Usage

```
continuousPower(simResult, contN = TRUE, contMCAR = FALSE, contMAR = FALSE,
contParam = NULL, alpha = .05, powerParam = NULL, pred = NULL)
```

Arguments

simResult	SimResult that includes at least one randomly varying parameter (e.g. sample size, percent missing, model parameters)
contN	Logical indicating if N varies over replications.
contMCAR	Logical indicating if the percentage of missing data that is MCAR varies over replications.
contMAR	Logical indicating if the percentage of missing data that is MAR varies over replications.
contParam	Vector of parameters names that vary over replications.
alpha	Alpha level to use for power analysis.
powerParam	Vector of parameters names that the user wishes to find power for. This can be a vector of names (e.g., "LY1_1", "LY2_2"), or the name of a matrix (e.g. "PS"), if the name of a matrix is used power for all parameters in that matrix will be returned. If parameters are not specified, power for all parameters in the model will be returned.

A list of varying parameter values that users wish to find statistical power from.

Details

pred

A common use of simulations is to conduct power analyses, especially when using SEM (Muthen & Muthen, 2002). Here, researchers select values for each parameter and a sample size and run a simulation to determine power in those conditions (the proportion of generated datasets in which a particular parameter of interest is significantly different from zero). To evaluate power at multiple sample sizes, one simulation for each sample size must be run. By continuously varying sample size across replications, only a single simulation is needed. In this simulation, the sample size for each replication varies randomly across plausible sample sizes (e.g., sample sizes between 200 and 500). For each replication, the sample size and significance of each parameter (0 = not significant, 1 = significant) are recorded. When the simulation is complete, parameter significance is regressed on sample size using logistic regression. For a given sample size, the predicted probability from the logistic regression equation is the power to detect an effect at that sample size. This approach can be extended to other randomly varying simulation parameters such as the percentage of missing data, and model parameters.

createData 11

Value

Data frame containing columns representing values of the randomly varying simulation parameters, and power for model parameters of interest.

Author(s)

Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>), Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

References

Muthen, L. K., & Muthen, B. O. (2002). How to use a Monte Carlo study to decide on sample size and determine power. *Structural Equation Modeling*, 4, 599-620.

See Also

• SimResult to see how to create a simResult object with randomly varying parameters.

Examples

```
## Not run:
# Specify Sample Size by n
loading <- matrix(0, 6, 1)</pre>
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
dat <- generate(CFA.Model, 50)</pre>
out <- analyze(CFA.Model, dat)</pre>
# We will use only 5 replications to save time.
# In reality, more replications are needed.
# Specify both sample size and percent missing completely at random
Output <- sim(NULL, CFA.Model, n=seq(100, 200, 20), pmMCAR=c(0, 0.1, 0.2))
summary(Output)
Cpow <- continuousPower(Output, contN = TRUE, contMCAR = TRUE)</pre>
Cpow
Cpow2 <- continuousPower(Output, contN = TRUE, contMCAR = TRUE, pred=list(N = 200, pmMCAR = 0.3))</pre>
Cpow2
## End(Not run)
```

createData

Create data from a set of drawn parameters.

12 createData

Description

This function can be used to create data from a set of parameters created from draw, called a codeparamSet. This function is used internally to create data, and is available publicly for accessibility and debugging.

Usage

createData(paramSet, n, indDist=NULL, sequential=FALSE, facDist=NULL,
errorDist=NULL, indLab=NULL, modelBoot=FALSE, realData=NULL)

Arguments

paramSet Set of drawn parameters from draw. Integer of desired sample size. indDist A SimDataDist object or list of objects for a distribution of indicators. If one object is passed, each indicator will have the same distribution. Use when sequential is FALSE. sequential If TRUE, use a sequential method to create data such that the data from factor are generated first and apply to a set of equations to obtain the data of indicators. If FALSE, create data directly from model-implied mean and covariance of indicators. facDist A SimDataDist object or list of objects for the distribution of factors. If one object is passed, all factors will have the same distribution. Use when sequential errorDist An object or list of objects of type SimDataDist indicating the distribution of errors. If a single SimDataDist is specified, each error will be genrated with that distribution. indLab A vector of indicator labels. When not specified, the variable names are x1, x2, ... xN. modelBoot When specified, a model-based bootstrap is used for data generation. See details for further information. This argument requires real data to be passed to realData A data frame containing real data. The data generated will follow the distribu-

Details

This function will use mvrnorm function in MASS package to create data from model implied covariance matrix if the data distribution object (SimDataDist) is not specified. It the data distribution object is specified, the Gaussian copula model is used. See SimDataDist for further details. For the model-based bootstrap, the transformation proposed by Yung & Bentler (1996) is used. This procedure is the expansion from the Bollen and Stine (1992) bootstrap including a mean structure. The model-implied mean vector and covariance matrix with trivial misspecification will be used in the model-based bootstrap if misspec is specified. See page 133 of Bollen and Stine (1992) for a reference.

tion of this data set.

Internally, parameters are first drawn, and data is then created from these parameters. Both of these steps are available via the draw and createData functions respectively.

Value

A data frame containing simulated data from the data generation template. A variable "group" is appended indicating group membership.

draw 13

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>), Patrick Miller (University of Notre Dame; <pmille13@nd.edu>)

References

- Bollen, K. A., & Stine, R. A. (1992). Bootstrapping goodness-of-fit measures in structural equation models. *Sociological Methods and Research*, 21, 205-229.
- Yung, Y.-F., & Bentler, P. M. (1996). Bootstrapping techniques in analysis of mean and covariance structures. In G. A. Marcoulides & R. E. Schumacker (Eds.), *Advanced structural equation modeling: Issues and techniques* (pp. 195-226). Mahwah, NJ: Erlbaum.

Examples

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
LY <- bind(loading, 0.7)

latent.cor <- matrix(NA, 2, 2)
diag(latent.cor) <- 1
RPS <- binds(latent.cor, 0.5)

RTE <- binds(diag(6))

VY <- bind(rep(NA,6),2)

CFA.Model <- model(LY = LY, RPS = RPS, RTE = RTE, modelType = "CFA")
param <- draw(CFA.Model)

# Generate data from the first group in the paramList.
dat <- createData(param[[1]], n = 200)</pre>
```

draw

Draw parameters from a SimSem object.

Description

This function draws parameters from a SimSem template, for debugging or other use. Used internally to create data. Data can be created in one step from a SimSem object using generate.

Usage

```
draw(model, maxDraw=50, misfitBounds=NULL, averageNumMisspec=FALSE,
optMisfit = NULL, optDraws=50, misfitType="f0")
```

Arguments

model A SimSem object.

maxDraw Integer specifying the maximum number of attempts to draw a valid set of parameters (no negative error variance, standardized coefficients over 1).

14 draw

misfitBounds Vector that contains upper and lower bounds of the misfit measure. Sets of parameters drawn that are not within these bounds are rejected.

averageNumMisspec

TRUE or FALSE. ??

optMisfit Character vector of either "min" or "max" indicating either maximum or mini-

mum optimized misfit. If not null, the set of parameters out of the number of draws in "optDraws" that has either the maximum or minimum misfit of the

given misfit type will be returned.

optDraws Number of parameter sets to draw if optMisfit is not null. The set of parameters

with the maximum or minimum misfit will be returned.

misfitType Character vector indicating the fit measure used to assess the misfit of a set of

parameters. Can be "f0", "rmsea", "srmr", or "all".

Value

Nested list of drawn parameters in the form [[Group]][[param,misspec,misOnly]][[SimMatrix]]. E.g. The LY parameter matrix of the first group would be indexed as obj[[1]]\$param\$LY. The values in \$param are the raw parameter values with no misspecification. The values in \$misspec are raw parameter values + misspecification. The values in \$misOnly are only the misspecification values.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>), Patrick Miller (University of Notre Dame; <pmille13@nd.edu>)

See Also

createData To generate random data using a set of parameters from draw

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
LY <- bind(loading, 0.7)

latent.cor <- matrix(NA, 2, 2)
diag(latent.cor) <- 1
RPS <- binds(latent.cor, 0.5)

RTE <- binds(diag(6))

VY <- bind(rep(NA,6),2)

CFA.Model <- model(LY = LY, RPS = RPS, RTE = RTE, modelType = "CFA")
param <- draw(CFA.Model)</pre>
```

estmodel 15

estmodel	Shortcut for data analysis template for simulation.	

Description

Creates a data analysis template (lavaan parameter table) for simulations with structural equation models based on Y-side LISREL design matrices. Each corresponds to a LISREL matrix, but must be a matrix or a vector. In addition to the usual Y-side matrices in LISREL, both PS and TE can be specified using correlations (RPS, RTE) and scaled by a vector of residual variances (VTE, VPS) or total variances (VY, VE). Multiple groups are supported by passing lists of matrices or vectors to arguments, or by specifying the number of groups.

Usage

```
estmodel(LY = NULL, PS = NULL, RPS = NULL, TE = NULL, RTE = NULL, BE = NULL,
VTE = NULL, VY = NULL, VPS = NULL, VE=NULL, TY = NULL, AL = NULL,
MY = NULL, ME = NULL, modelType, indLab=NULL, facLab=NULL, groupLab="group",
ngroups=1, smartStart=TRUE)
estmodel.cfa(LY = NULL,PS = NULL,RPS = NULL, TE = NULL,RTE = NULL, VTE = NULL,
VY = NULL, VPS = NULL, VE=NULL, TY = NULL, AL = NULL, MY = NULL, ME = NULL,
indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
estmodel.path(PS = NULL, RPS = NULL, BE = NULL, VPS = NULL, VE=NULL, AL = NULL,
ME = NULL, indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
estmodel.sem(LY = NULL,PS = NULL,RPS = NULL, TE = NULL,RTE = NULL, BE = NULL,
VTE = NULL, indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
```

Arguments

LY	Factor loading matrix from endogenous factors to Y indicators (need to be a matrix or a list of matrices).
PS	Residual covariance matrix among endogenous factors (need to be a symmetric matrix or a list of symmetric matrices).
RPS	Residual correlation matrix among endogenous factors (need to be a symmetric matrix or a list of symmetric matrices).
TE	Measurement error covariance matrix among Y indicators (need to be a symmetric matrix or a list of symmetric matrices).
RTE	Measurement error correlation matrix among Y indicators (need to be a symmetric matrix or a list of symmetric matrices).
BE	Regression coefficient matrix among endogenous factors (need to be a matrix or a list of matrices).
VTE	Measurement error variance of indicators (need to be a vector or a list of vectors).
VY	Total variance of indicators (need to be a vector or a list of vectors). NOTE: Either measurement error variance or indicator variance is specified. Both cannot be simultaneously specified.
VPS	Residual variance of factors (need to be a vector or a list of vectors).

16 estmodel

VE	Total variance of of factors (need to be a vector or a list of vectors). NOTE: Either residual variance of factors or total variance of factors is specified. Both cannot be simulatneously specified.
TY	Measurement intercepts of Y indicators. (need to be a vector or a list of vectors).
AL	Endogenous factor intercept (need to be a vector or a list of vectors).
MY	Overall Y indicator means. (need to be a vector or a list of vectors). NOTE: Either measurement intercept of indicator mean can be specified. Both cannot be specified simultaneously.
ME	Total mean of endogenous factors (need to be a vector or a list of vectors). NOTE: Either endogenous factor intercept or total mean of endogenous factor is specified. Both cannot be simultaneously specified.
modelType	"CFA", "Sem", or "Path". This is specified to ensure that the analysis and data generation template created based on specified matrices in model correspond to what the user intends.
indLab	Character vector of indicator labels. If left blank, automatic labels will be generated as $y1, y2, \dots yy$.
facLab	Character vector of factor labels. If left blank, automatic labels will be generated as $f1, f2, \dots ff$
groupLab	Character of group-variable label (not the names of each group). If left blank, automatic labels will be generated as group
ngroups	Integer. Number of groups for data generation, defaults to 1. If larger than one, all specified matrices will be repeated for each additional group. If any matrix argument is a list, the length of this list will be the number of groups and ngroups is ignored.
smartStart	Defaults to FALSE. If TRUE, population parameter values that are real numbers will be used as starting values.

Details

This function contains default settings:

For modelType="CFA", LY is required. As the default, the on-diagonal elements of PS are fixed as 1 and the off-diagonal elements of PS are freely estimated. The off-diagonal elements of TE are freely estimated and the off-diagonal elements of TE are fixed to 0. The AL elements are fixed to 0. The TY elements are freely estimated.

For modelType="Path", BE is required. As the default, the on-diagonal elements of PS are freely estimated, the off-diagonal elements between exogenous variables (covariance between exogenous variables) are freely estimated, and the other off-diagonal elements are fixed to 0. The AL elements are freely estimated.

For modelType="SEM", LY and BE are required. As the default, the on-diagonal elements of PS are fixed to 1, the off-diagonal elements between exogenous factors (covariance between exogenous factors) are freely estimated, and the other off-diagonal elements are fixed to 0. The off-diagonal elements of TE are freely estimated and the off-diagonal elements of TE are fixed to 0. The AL elements are fixed to 0. The TY elements are freely estimated.

The estmodel.cfa, estmodel.path, and estmodel.sem are the shortcuts for the estmodel function when modelType are "CFA", "Path", and "SEM", respectively.

Value

SimSem object that contains the data generation template (@dgen) and analysis template (@pt).

extractLavaanFit 17

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- model To build data generation and data analysis template for simulation.
- sim For simulations using the SimSem template.
- generate To generate data using the SimSem template.
- analyze To analyze real or generated data using the SimSem template.
- draw To draw parameters using the SimSem template.

Examples

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA

CFA.Model <- estmodel(LY = loading, modelType = "CFA")</pre>
```

extractLavaanFit

Extract fit indices from the lavaan object

Description

Extract fit indices from the lavaan object

Usage

```
extractLavaanFit(Output)
```

Arguments

Output

The lavaan object

Value

The renamed vector of fit measures

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

```
# No example
```

18 findFactorIntercept

find2Dhist

Fit the 2D Kernel Density Estimate

Description

Fit the 2D Kernel Density Estimate to a pair of variables

Usage

```
find2Dhist(vec1, vec2)
```

Arguments

vec1 Variable 1 vec2 Variable 2

Value

The 2D Kernel Density Estimate based on each pair of values in vec1 and vec2

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

No example

findFactorIntercept

Find factor intercept from regression coefficient matrix and factor total means

Description

Find factor intercept from regression coefficient matrix and factor total means for latent variable models. In the path analysis model, this function will find indicator intercept from regression coefficient and indicator total means.

Usage

```
findFactorIntercept(beta, factorMean = NULL)
```

Arguments

beta Regression coefficient matrix

factorMean Total (model-implied) factor (indicator) means. The default is that all total factor

means are 0.

findFactorMean 19

Value

A vector of factor (indicator) intercepts

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

Examples

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- 0.6
path[5, 2] <- path[8, 5] <- 0.6
path[6, 3] <- path[9, 6] <- 0.6
path[5, 1] <- path[8, 4] <- 0.4
path[6, 2] <- path[9, 5] <- 0.4
factorMean <- c(5, 2, 3, 0, 0, 0, 0, 0, 0)
findFactorIntercept(path, factorMean)</pre>
```

findFactorMean

Find factor total means from regression coefficient matrix and factor intercept

Description

Find factor total means from regression coefficient matrix and factor intercepts for latent variable models. In the path analysis model, this function will find indicator total means from regression coefficient and indicator intercept.

Usage

```
findFactorMean(beta, alpha = NULL)
```

Arguments

beta Regression coefficient matrix

alpha Factor (indicator) intercept. The default is that all factor intercepts are 0.

20 findFactorResidualVar

Value

A vector of factor (indicator) total means

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

Examples

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- 0.6
path[5, 2] <- path[8, 5] <- 0.6
path[6, 3] <- path[9, 6] <- 0.6
path[5, 1] <- path[8, 4] <- 0.4
path[6, 2] <- path[9, 5] <- 0.4
intcept <- c(5, 2, 3, 0, 0, 0, 0, 0, 0)
findFactorMean(path, intcept)</pre>
```

findFactorResidualVar Find factor residual variances from regression coefficient matrix, factor (residual) correlations, and total factor variances

Description

Find factor residual variances from regression coefficient matrix, factor (residual) correlation matrix, and total factor variances for latent variable models. In the path analysis model, this function will find indicator residual variances from regression coefficient, indicator (residual) correlation matrix, and total indicator variances.

Usage

```
findFactorResidualVar(beta, corPsi, totalVarPsi = NULL)
```

Arguments

beta Regression coefficient matrix

corPsi Factor or indicator residual correlations.

totalVarPsi Factor or indicator total variances. The default is that all factor or indicator total

variances are 1.

findFactorTotalCov 21

Value

A vector of factor (indicator) residual variances

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

Examples

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- 0.6
path[5, 2] <- path[8, 5] <- 0.6
path[6, 3] <- path[9, 6] <- 0.6
path[5, 1] <- path[8, 4] <- 0.4
path[6, 2] <- path[9, 5] <- 0.4
facCor <- diag(9)
facCor[1, 2] <- facCor[2, 1] <- 0.4
facCor[1, 3] <- facCor[3, 1] <- 0.4
facCor[2, 3] <- facCor[3, 2] <- 0.4
totalVar <- rep(1, 9)
findFactorResidualVar(path, facCor, totalVar)</pre>
```

find Factor Total Cov

Find factor total covariance from regression coefficient matrix, factor residual covariance

Description

Find factor total covariances from regression coefficient matrix, factor residual covariance matrix. The residual covariance matrix might be derived from factor residual correlation, total variance, and error variance. This function can be applied for path analysis model as well.

Usage

```
findFactorTotalCov(beta, psi=NULL, corPsi=NULL, totalVarPsi = NULL, errorVarPsi=NULL)
```

22 findFactorTotalCov

Arguments

beta	Regression coefficient matrix
psi	Factor or indicator residual covariances. This argument can be skipped if factor residual correlation and either total variances or error variances are specified.
corPsi	Factor or indicator residual correlation. This argument must be specified with total variances or error variances.
totalVarPsi	Factor or indicator total variances.
errorVarPsi	Factor or indicator residual variances.

Value

A matrix of factor (model-implied) total covariance

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- 0.6
path[5, 2] <- path[8, 5] <- 0.6
path[6, 3] <- path[9, 6] <- 0.6
path[5, 1] <- path[8, 4] <- 0.4
path[6, 2] <- path[9, 5] <- 0.4
facCor <- diag(9)
facCor[1, 2] <- facCor[2, 1] <- 0.4
facCor[1, 3] <- facCor[3, 1] <- 0.4
facCor[2, 3] <- facCor[3, 2] <- 0.4
residualVar <- c(1, 1, 1, 0.64, 0.288, 0.288, 0.64, 0.29568, 0.21888)
findFactorTotalCov(path, corPsi=facCor, errorVarPsi=residualVar)</pre>
```

findFactorTotalVar 23

findFactorTotalVar	Find factor total variances from regression coefficient matrix, factor
	(residual) correlations, and factor residual variances

Description

Find factor total variances from regression coefficient matrix, factor (residual) correlation matrix, and factor residual variances for latent variable models. In the path analysis model, this function will find indicator total variances from regression coefficient, indicator (residual) correlation matrix, and indicator residual variances.

Usage

```
findFactorTotalVar(beta, corPsi, residualVarPsi)
```

Arguments

```
beta Regression coefficient matrix

corPsi Factor or indicator residual correlations.

residualVarPsi Factor or indicator residual variances.
```

Value

A vector of factor (indicator) total variances

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalCov to find factor covariances

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- 0.6
path[5, 2] <- path[8, 5] <- 0.6
path[6, 3] <- path[9, 6] <- 0.6
path[5, 1] <- path[8, 4] <- 0.4
path[6, 2] <- path[9, 5] <- 0.4
facCor <- diag(9)
facCor[1, 2] <- facCor[2, 1] <- 0.4</pre>
```

24 findIndIntercept

```
facCor[1, 3] <- facCor[3, 1] <- 0.4
facCor[2, 3] <- facCor[3, 2] <- 0.4
residualVar <- c(1, 1, 1, 0.64, 0.288, 0.288, 0.64, 0.29568, 0.21888)
findFactorTotalVar(path, facCor, residualVar)</pre>
```

findIndIntercept

Find indicator intercepts from factor loading matrix, total factor mean, and indicator mean.

Description

Find indicator (measurement) intercepts from a factor loading matrix, total factor mean, and indicator mean.

Usage

```
findIndIntercept(lambda, factorMean = NULL, indicatorMean = NULL)
```

Arguments

lambda Factor loading matrix

factorMean Total (model-implied) mean of factors. As a default, all total factor means are 0.

indicatorMean Total indicator means. As a default, all total indicator means are 0.

Value

A vector of indicator (measurement) intercepts.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- c(0.6, 0.7, 0.8)
loading[4:6, 2] <- c(0.6, 0.7, 0.8)
facMean <- c(0.5, 0.2)
indMean <- rep(1, 6)
findIndIntercept(loading, facMean, indMean)</pre>
```

findIndMean 25

findIndMean	Find indicator total means from factor loading matrix, total factor mean, and indicator intercept.
	•

Description

Find indicator total means from a factor loading matrix, total factor means, and indicator (measurement) intercepts.

Usage

```
findIndMean(lambda, factorMean = NULL, tau = NULL)
```

Arguments

lambda Factor loading matrix

factorMean Total (model-implied) mean of factors. As a default, all total factor means are 0.

tau Indicator (measurement) intercepts. As a default, all intercepts are 0.

Value

A vector of indicator total means.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndResidualVar to find indicator (measurement) residual variances
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- c(0.6, 0.7, 0.8)
loading[4:6, 2] <- c(0.6, 0.7, 0.8)
facMean <- c(0.5, 0.2)
intcept <- rep(0, 6)
findIndMean(loading, facMean, intcept)</pre>
```

26 findIndResidualVar

findIndResidualVar	Find indicator residual variances from factor loading matrix, total fac-
	tor covariance, and total indicator variances.

Description

Find indicator (measurement) residual variances from a factor loading matrix, total factor covariance matrix, and total indicator variances.

Usage

```
findIndResidualVar(lambda, totalFactorCov, totalVarTheta = NULL)
```

Arguments

```
lambda Factor loading matrix
totalFactorCov Total (model-implied) covariance matrix among factors.
totalVarTheta Indicator total variances. As a default, all total variances are 1.
```

Value

A vector of indicator residual variances.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndTotalVar to find indicator (measurement) total variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- c(0.6, 0.7, 0.8)
loading[4:6, 2] <- c(0.6, 0.7, 0.8)
facCov <- matrix(c(1, 0.5, 0.5, 1), 2, 2)
totalVar <- rep(1, 6)
findIndResidualVar(loading, facCov, totalVar)</pre>
```

findIndTotalVar 27

findIndTotalVar Find indicator total variances from factor loading matrix, total covariance, and indicator residual variances.	factor
--	--------

Description

Find indicator total variances from a factor loading matrix, total factor covariance matrix, and indicator (measurement) residual variances.

Usage

```
findIndTotalVar(lambda, totalFactorCov, residualVarTheta)
```

Arguments

```
lambda Factor loading matrix
totalFactorCov Total (model-implied) covariance matrix among factors.
residualVarTheta
Indicator (measurement) residual variances.
```

Value

A vector of indicator total variances.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- findIndIntercept to find indicator (measurement) intercepts
- findIndMean to find indicator (measurement) total means
- findIndResidualVar to find indicator (measurement) residual variances
- findFactorIntercept to find factor intercepts
- findFactorMean to find factor means
- findFactorResidualVar to find factor residual variances
- findFactorTotalVar to find factor total variances
- findFactorTotalCov to find factor covariances

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- c(0.6, 0.7, 0.8)
loading[4:6, 2] <- c(0.6, 0.7, 0.8)
facCov <- matrix(c(1, 0.5, 0.5, 1), 2, 2)
resVar <- c(0.64, 0.51, 0.36, 0.64, 0.51, 0.36)
findIndTotalVar(loading, facCov, resVar)</pre>
```

28 findPossibleFactorCor

findphist	Find the density (likelihood) of a pair value in 2D Kernel Density Es-
	timate

Description

Find the density (likelihood) of a pair value in 2D Kernel Density Estimate

Usage

```
findphist(value, hist)
```

Arguments

value A target pair of values

hist A 2D Binned Kernel Density Estimate

Value

The probability (density) of the target pair of value

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

No example

findPossibleFactorCor Find the appropriate position for freely estimated correlation (or covariance) given a regression coefficient matrix

Description

Find the appropriate position for freely estimated correlation (or covariance) given a regression coefficient matrix. The appropriate position is the pair of variables that are not causally related.

Usage

findPossibleFactorCor(beta)

Arguments

beta The regression coefficient in path analysis.

Value

The symmetric matrix containing the appropriate position for freely estimated correlation.

findPower 29

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• findRecursiveSet to group variables regarding the position in mediation chain.

Examples

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- NA
path[5, 2] <- path[8, 5] <- NA
path[6, 3] <- path[9, 6] <- NA
path[5, 1] <- path[8, 4] <- NA
path[6, 2] <- path[9, 5] <- NA
findPossibleFactorCor(path)</pre>
```

findPower

Find a value of independent variables that provides a given value of power.

Description

Find a value of independent variable that provides a given value of power. If there are more than one varying parameters, this function will find the value of the target varying parameters given the values of the other varying parameters.

Usage

```
findPower(powerTable, iv, power)
```

Arguments

powerTable A data.frame providing varying parameters and powers of each parameter.

This table is obtained by getPower or continuousPower function.

iv The target varying parameter that users would like to find the value providing

a given power from. This argument can be specified as the index of the target

column or the name of target column (i.e., "iv.N" or "N")

power A desired power.

Value

There are five possible types of values provided:

- *Value* The varying parameter value that provides the power just over the specified power value (the adjacent value of varying parameter provides lower power than the specified power value).
- *Minimum value* The minimum value has already provided enough power (way over the specified power value). The value of varying parameters that provides exact desired power may be lower than the minimum value. The example of varying parmaeter that can provides the minimum value is sample size.

30 findRecursiveSet

• *Maximum value* The maximum value has already provided enough power (way over the specified power value). The value of varying parameters that provides exact desired power may be higher than the maximum value. The example of varying parmaeter that can provides the maximum value is percent missing.

- NA There is no value in the domain of varying parameters that provides the power greater than the desired power.
- NaN The power of all values in the varying parameters is 1 (specifically more than 0.9999) and any values of the varying parameters can be picked and still provide enough power.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- getPower to find the power of parameter estimates
- continuousPower to find the power of parameter estimates for the result object (linkS4class{SimResult}) with varying parameters.

Examples

```
## Not run:
# Specify Sample Size by n
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, 0.4)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")

# Specify both sample size and percent missing completely at random
Output <- sim(NULL, model=CFA.Model, n=seq(100, 200, 20), pmMCAR=c(0, 0.1, 0.2))
pow <- getPower(Output)
findPower(pow, "N", 0.80)

## End(Not run)</pre>
```

findRecursiveSet

Group variables regarding the position in mediation chain

Description

In mediation analysis, variables affects other variables as a chain. This function will group variables regarding the chain of mediation analysis.

Usage

```
findRecursiveSet(beta)
```

Arguments

beta

The regression coefficient in path analysis.

findRowZero 31

Value

The list of set of variables in the mediation chain. The variables in position 1 will be the independent variables. The variables in the last variables will be the end of the chain.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

 findPossibleFactorCor to find the possible position for latent correlation given a regression coefficient matrix

Examples

```
path <- matrix(0, 9, 9)
path[4, 1] <- path[7, 4] <- NA
path[5, 2] <- path[8, 5] <- NA
path[6, 3] <- path[9, 6] <- NA
path[5, 1] <- path[8, 4] <- NA
path[6, 2] <- path[9, 5] <- NA
findRecursiveSet(path)</pre>
```

findRowZero

Find rows in a matrix that all elements are zero in non-fixed subset rows and columns.

Description

Find rows in a matrix that all elements are zero in non-fixed subset rows and columns. This function will be used in the findRecursiveSet function

Usage

```
findRowZero(square.matrix, is.row.fixed = FALSE)
```

Arguments

```
square.matrix Any square matrix
is.row.fixed A logical vector with the length equal to the dimension of the square.matrix.
If TRUE, the function will skip examining this row.
```

Value

A vector of positions that contain rows of all zeros

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

```
# No example
```

32 findTargetPower

findTargetPower	Find a value of varying parameters that provides a given value of power.

Description

Find a value of varying parameters that provides a given value of power. This function can deal with only one varying parameter only (findPower can deal with more than one varying parameter).

Usage

```
findTargetPower(iv, dv, power)
```

Arguments

iv A vector of the target varying parameter

dv A data. frame of the power table of target parameters

power A desired power.

Value

The value of the target varying parameter providing the desired power. If the value is NA, there is no value in the domain of varying parameters that provide the target power. If the value is the minimum value of the varying parameters, it means that the minimum value has already provided enough power. The value of varying parameters that provides exact desired power may be lower than the minimum value.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- getPower to find the power of parameter estimates
- continuousPower to find the power of parameter estimates for the result object (linkS4class{SimResult}) with varying parameters.
- findPower to find a value of varying parameters that provides a given value of power, which can deal with more than one varying parameter

Examples

No example

fitMeasuresChi 33

fitMeasuresChi Find fit indices from the discrepancy values of the target model and null models.	fit	:MeasuresChi		
--	-----	--------------	--	--

Description

Find fit indices from the discrepancy values of the target model and null models. This function is modified from the fitMeasures function in lavaan package

Usage

```
fitMeasuresChi(X2, df, p, X2.null, df.null, p.null, N, fit.measures="all")
```

Arguments

X2	The chi-square value of the target model
df	The degree of freedom of the target model
p	The p vlaue of the target model
X2.null	The chi-square value of the null model
df.null	The degree of freedom of the null model
p.null	The p value of the null model
N	Sample size
fit.measures	The list of selected fit measures

Value

A vector of fit measures

Author(s)

Yves Rosseel in the lavaan package Modified by Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

No example

Description

This function can be used to generate random data based on the SimSem template. Some notable features include fine control of misspecification and misspecification optimization, as well as the ability to generate non-normal data. When using *simsem* for simulations, this function is used internally to generate data in the function sim, and can be helpful for debugging, or in creating data for use with other analysis programs.

34 generate

Usage

```
generate(model, n, maxDraw=50, misfitBounds=NULL, misfitType="f0",
averageNumMisspec=FALSE, optMisfit=NULL, optDraws=50,
indDist=NULL, sequential=FALSE, facDist=NULL, errorDist=NULL,
indLab=NULL, modelBoot=FALSE, realData=NULL, params=FALSE)
```

Arguments

n

model A SimSem object. Integer of sample size.

maxDraw Integer specifying the maximum number of attempts to draw a valid set of pa-

rameters (no negative error variance, standardized coefficients over 1).

Vector that contains upper and lower bounds of the misfit measure. Sets of misfitBounds

parameters drawn that are not within these bounds are rejected.

misfitType Character vector indicating the fit measure used to assess the misfit of a set of

parameters. Can be "f0", "rmsea", "srmr", or "all".

averageNumMisspec

TRUE or FALSE. ??

optMisfit Character vector of either "min" or "max" indicating either maximum or mini-

> mum optimized misfit. If not null, the set of parameters out of the number of draws in "optDraws" that has either the maximum or minimum misfit of the

given misfit type will be returned.

Number of parameter sets to draw if optMisfit is not null. The set of parameters optDraws

with the maximum or minimum misfit will be returned.

indDist A SimDataDist object or list of objects for a distribution of indicators. If

one object is passed, each indicator will have the same distribution. Use when

sequential is FALSE.

sequential If TRUE, use a sequential method to create data such that the data from factor

> are generated first and apply to a set of equations to obtain the data of indicators. If FALSE, create data directly from model-implied mean and covariance of

indicators.

facDist A SimDataDist object or list of objects for the distribution of factors. If one ob-

ject is passed, all factors will have the same distribution. Use when sequential

is TRUE.

errorDist An object or list of objects of type SimDataDist indicating the distribution of

errors. If a single SimDataDist is specified, each error will be genrated with

that distribution.

indLab A vector of indicator labels. When not specified, the variable names are x1, x2, ... xN.

modelBoot When specified, a model-based bootstrap is used for data generation. See de-

tails for further information. This argument requires real data to be passed to

readData.

realData A data frame containing real data. The data generated will follow the distribu-

tion of this data set.

params If TRUE, return the parameters drawn along with the generated data set. Default

is FALSE.

generate 35

Details

This function will use mvrnorm function in MASS package to create data from model implied covariance matrix if the data distribution object (SimDataDist) is not specified. It the data distribution object is specified, the Gaussian copula model is used. See SimDataDist for further details. For the model-based bootstrap, the transformation proposed by Yung & Bentler (1996) is used. This procedure is the expansion from the Bollen and Stine (1992) bootstrap including a mean structure. The model-implied mean vector and covariance matrix with trivial misspecification will be used in the model-based bootstrap if misspec is specified. See page 133 of Bollen and Stine (1992) for a reference.

Internally, parameters are first drawn, and data is then created from these parameters. Both of these steps are available via the draw and createData functions respectively.

Value

A data frame containing simulated data from the data generation template. A variable "group" is appended indicating group membership.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>), Patrick Miller (University of Notre Dame; <pmille13@nd.edu>)

References

- Bollen, K. A., & Stine, R. A. (1992). Bootstrapping goodness-of-fit measures in structural equation models. *Sociological Methods and Research*, *21*, 205-229.
- Yung, Y.-F., & Bentler, P. M. (1996). Bootstrapping techniques in analysis of mean and covariance structures. In G. A. Marcoulides & R. E. Schumacker (Eds.), *Advanced structural equation modeling: Issues and techniques* (pp. 195-226). Mahwah, NJ: Erlbaum.

See Also

- draw To draw parameters using the SimSem template.
- createData To generate random data using a set of parameters from draw

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
LY <- bind(loading, 0.7)

latent.cor <- matrix(NA, 2, 2)
diag(latent.cor) <- 1
RPS <- binds(latent.cor, 0.5)

RTE <- binds(diag(6))

VY <- bind(rep(NA,6),2)

CFA.Model <- model(LY = LY, RPS = RPS, RTE = RTE, modelType = "CFA")
dat <- generate(CFA.Model,200)</pre>
```

36 getCondQtile

getCondQtile Get a quantile of a variable given values of predictors	
--	--

Description

Find a quantile of a variable. If the predictors are specified, the result will provide the conditional quantile given specified value of predictors. The quantreg package is used to find conditional quantile.

Usage

```
getCondQtile(y, x=NULL, xval=NULL, df = 0, qtile = 0.5)
```

Arguments

У	The variable that users wish to find a quantile from
Х	The predictors variables. If NULL, the unconditional quantile of the y is provided.
xval	The vector of predictors' values that users wish to find the conditional quantile from. If "all" is specified, the function will provide the conditional quantile of every value in x.
df	The degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.
qtile	The quantile rank.

Value

A (conditional) quantile value

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• getCutoff for finding fit indices cutoffs using conditional quantiles

```
# No example
```

getCutoff 37

getCutoff	Find fit indices cutoff given a priori alpha level

Description

Extract fit indices information from the SimResult and getCutoff of fit indices given a priori alpha level

Usage

```
getCutoff(object, alpha, revDirec = FALSE, usedFit = NULL, ...)
```

Arguments

object
SimResult that saves the analysis results from multiple replications
A priori alpha level

The default is to find criticl point on the side that indicates worse fit (the right side of RMSEA or the left side of CFI). If specifying as TRUE, the directions are reversed.

UsedFit
Vector of names of fit indices that researchers wish to getCutoffs from. The default is to getCutoffs of all fit indices.

.. Additional arguments.

Value

One-tailed cutoffs of several fit indices with a priori alpha level

Methods

signature(object=''data.frame'') This method will find the fit indices cutoff given a specified alpha level. The additional arguments are predictor, predictorVal, and df, which allows the fit indices predicted by any arbitrary independent variables (such as sample size or percent MCAR). The predictor is the data.frame of the predictor values. The number of rows of the predictor argument should be equal to the number of rows in the object. The predictorVal is the values of predictor that researchers would like to find the fit indices cutoffs from. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

signature(object="matrix") The details are similar to the method for data. frame

signature(object="SimResult") This method will find the fit indices cutoff given a specified alpha level. The additional arguments are nVal, pmMCARval, pmMARval, and df, which are needed when using varying sample sizes or percent missing across replications in SimResult. The nVal is the sample size value that researchers wish to find the fit indices cutoffs from. The pmMCARval is the percent missing completely at random value that researchers wish to find the fit indices cutoffs from. The pmMARval is the percent missing at random value that researchers wish to find the fit indices cutoffs from. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

38 getCutoffNested

See Also

SimResult for a detail of simResult

Examples

```
## Not run:
loading <- matrix(0, 6, 2)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loadingValues <- matrix(0, 6, 2)
loadingValues[1:3, 1] <- 0.7</pre>
loadingValues[4:6, 2] <- 0.7</pre>
LX <- bind(loading, loadingValues)
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, 0.5)
error.cor <- matrix(0, 6, 6)</pre>
diag(error.cor) <- 1</pre>
RTD <- binds(error.cor)</pre>
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output < sim(5, n = 200, model=CFA.Model)
getCutoff(Output, 0.05)
# Finding the cutoff when the sample size is varied.
Output2 <- sim(NULL, model=CFA.Model, n=seq(50, 100, 10))
getCutoff(Output2, 0.05, nVal = 75)
## End(Not run)
```

getCutoffNested

Find fit indices cutoff for nested model comparison given a priori alpha level

Description

Extract fit indices information from the simulation of parent and nested models and getCutoff of fit indices given a priori alpha level

Usage

```
getCutoffNested(nested, parent, alpha = 0.05, usedFit = NULL, nVal = NULL, pmMCARval = NULL, pmM
```

Arguments

nested	SimResult that saves the analysis results of nested model from multiple replications
parent	SimResult that saves the analysis results of parent model from multiple replications
alpha	A priori alpha level

getCutoffNested 39

usedFit	Vector of names of fit indices that researchers wish to getCutoffs from. The default is to getCutoffs of all fit indices.
nVal	The sample size value that researchers wish to find the fit indices cutoffs from.
pmMCARval	The percent missing completely at random value that researchers wish to find the fit indices cutoffs from.
pmMARval	The percent missing at random value that researchers wish to find the fit indices cutoffs from.
df	The degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

Value

One-tailed cutoffs of several fit indices with a priori alpha level

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

SimResult for a detail of simResult getCutoff for a detail of finding cutoffs for absolute fit

```
## Not run:
loading.null <- matrix(0, 6, 1)</pre>
loading.null[1:6, 1] <- NA</pre>
LX.NULL <- bind(loading.null, 0.7)
RPH.NULL <- binds(diag(1))</pre>
error.cor.mis <- matrix("rnorm(1, 0, 0.1)", 6, 6)
diag(error.cor.mis) <- 1</pre>
RTD <- binds(diag(6), misspec=error.cor.mis)</pre>
CFA.Model.NULL <- model(LY = LX.NULL, RPS = RPH.NULL, RTE = RTD, modelType="CFA")
loading.alt <- matrix(0, 6, 2)</pre>
loading.alt[1:3, 1] <- NA</pre>
loading.alt[4:6, 2] <- NA</pre>
LX.ALT <- bind(loading.alt, 0.7)
latent.cor.alt <- matrix(NA, 2, 2)</pre>
diag(latent.cor.alt) <- 1</pre>
\label{eq:RPH.ALT} \textit{RPH.ALT} <- binds(latent.cor.alt, "runif(1, 0.7, 0.9)")
CFA.Model.ALT <- model(LY = LX.ALT, RPS = RPH.ALT, RTE = RTD, modelType="CFA")
# The actual number of replications should be greater than 10.
Output.NULL.NULL <- sim(10, n=500, model=CFA.Model.NULL, generate=CFA.Model.NULL)
Output.NULL.ALT <- sim(10, n=500, model=CFA.Model.ALT, generate=CFA.Model.NULL)
getCutoffNested(Output.NULL.NULL, Output.NULL.ALT)
## End(Not run)
```

40 getCutoffNonNested

getCutoffNonNested	Find fit indices cutoff for non-nested model comparison given a priori alpha level
--------------------	--

Description

Extract fit indices information from the simulation of two models fitting on the datasets created from both models and getCutoff of fit indices given a priori alpha level

Usage

```
\label{eq:cutoffNonNested} $$ getCutoffNonNested(dat1Mod1, dat1Mod2, dat2Mod1=NULL, dat2Mod2=NULL, alpha=.05, usedFit=NULL, onetailed=FALSE, nVal = NULL, pmMCARval = NULL, pmMCARval = NULL, df = 0)
```

Arguments

dat1Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 1
dat1Mod2	SimResult that saves the simulation of analyzing Model 2 by datasets created from Model 1
dat2Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 2
dat2Mod2	${\tt SimResult}$ that saves the simulation of analyzing Model 2 by datasets created from Model 2
alpha	A priori alpha level
usedFit	Vector of names of fit indices that researchers wish to get cutoffs from. The default is to get cutoffs of all fit indices.
onetailed	If TRUE, the function will find the cutoff from one-tail test. If FALSE, the function will find the cutoff from two-tailed test.
nVal	The sample size value that researchers wish to find the fit indices cutoffs from.
pmMCARval	The percent missing completely at random value that researchers wish to find the fit indices cutoffs from.
pmMARval	The percent missing at random value that researchers wish to find the fit indices cutoffs from.
df	The degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

Value

One- or two-tailed cutoffs of several fit indices with a priori alpha level. The cutoff is based on the fit indices from Model 1 subtracted by the fit indices from Model 2.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

getExtraOutput 41

See Also

SimResult for a detail of simResult getCutoff for a detail of finding cutoffs for absolute fit getCutoffNested for a detail of finding cutoffs for nested model comparison plotCutoffNonNested Plot cutoffs for non-nested model comparison

Examples

```
## Not run:
loading.A <- matrix(0, 8, 2)</pre>
loading.A[1:3, 1] <- NA
loading.A[4:8, 2] <- NA
LX.A <- bind(loading.A, 0.7)
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, "runif(1, 0.7, 0.9)")
RTD <- binds(diag(8))
CFA.Model.A <- model(LY = LX.A, RPS = RPH, RTE = RTD, modelType="CFA")
loading.B <- matrix(0, 8, 2)</pre>
loading.B[1:4, 1] <- NA
loading.B[5:8, 2] <- NA
LX.B <- bind(loading.B, 0.7)
CFA.Model.B <- model(LY = LX.B, RPS = RPH, RTE = RTD, modelType="CFA")</pre>
# The actual number of replications should be greater than 10.
Output.A.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.A)
Output.A.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.A)
Output.B.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.B)</pre>
Output.B.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.B)
getCutoffNonNested(Output.A.A, Output.A.B, Output.B.A, Output.B.B)
getCutoffNonNested(Output.A.A, Output.A.B)
getCutoffNonNested(Output.B.B, Output.B.A)
## End(Not run)
```

getExtraOutput

Get extra outputs from the result of simulation

Description

Get extra outputs from a simulation result object (SimResult). Users can ask this package to extra output from the lavaan object in each iteration by setting the outfun argument (in the sim function). See the example below.

Usage

```
getExtraOutput(object)
```

Arguments

object

SimResult that have the extra output extracted by the function defined in the outfun argument (in the sim function)

42 getKeywords

Value

A list of extra outputs

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• sim A function to run a Monte Carlo simulation

Examples

```
## Not run:
# Specify Sample Size by n
loading <- matrix(0, 6, 1)</pre>
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
# We will use only 5 replications to save time.
# In reality, more replications are needed.
outfun <- function(out) {</pre>
result <- inspect(out, "mi")</pre>
# Specify both sample size and percent missing completely at random
Output <- sim(5, n=200, model=CFA.Model, outfun=outfun)
getExtraOutput(Output)
## End(Not run)
```

getKeywords

List of all keywords used in the simsem package

Description

List of all keywords used in the simsem package

Usage

```
getKeywords()
```

Value

A list of all keywords used in this package

- usedFit Fit indices used as the default for providing output
- usedFitPop Population fit indices used as the default for providing input
- optMin The method picking the minimum value of misfit across misspecification sets

getPopulation 43

- optMax The method picking the maximum value of misfit across misspecification sets
- optNone Not using the optimization method

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

```
# This function is not a public function.
# getKeywords()
```

getPopulation

Extract the data generation population model underlying a result object

Description

 $This function will extract the data generation population model underlying a result object (linkS4class\{SimResult\}).$

Usage

```
getPopulation(object)
```

Arguments

object

The result object that you wish to extract the data generation population model from (linkS4class{SimResult}).

Value

A data frame contained the population of each replication

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimResult for result object

```
## Not run:
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, "runif(1, 0.4, 0.9)")
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
# We will use only 10 replications to save time.</pre>
```

44 getPower

```
# In reality, more replications are needed.
Output <- sim(10, n=200, model=CFA.Model)
getPopulation(Output)
## End(Not run)</pre>
```

getPower

Find power of model parameters

Description

A function to find the power of parameters in a model when none, one, or more of the simulations parameters vary randomly across replications.

Usage

```
getPower(simResult, alpha = 0.05, contParam = NULL, powerParam = NULL,
nVal = NULL, pmMCARval = NULL, pmMARval = NULL, paramVal = NULL)
```

Arguments

simResult SimResult that may include at least one randomly varying parameter (e.g. sam-

ple size, percent missing, model parameters)

alpha Alpha level to use for power analysis.

contParam Vector of parameters names that vary over replications.

powerParam Vector of parameters names that the user wishes to find power for. This can be a

vector of names (e.g., "LY1_1", "LY2_2"), or the name of a matrix (e.g. "PS"), if the name of a matrix is used power for all parameters in that matrix will be returned. If parameters are not specified, power for all parameters in the model

will be returned.

nVal The sample size values that users wish to find power from.

pmMCARval The percent completely missing at random values that users wish to find power

from.

pmMARval The percent missing at random values that users wish to find power from.

A list of varying parameter values that users wish to find power from.

Details

A common use of simulations is to conduct power analyses, especially when using SEM (Muthen & Muthen, 2002). Here, researchers could select values for each parameter and a sample size and run a simulation to determine power in those conditions (the proportion of generated datasets in which a particular parameter of interest is significantly different from zero). To evaluate power at multiple sample sizes, one simulation for each sample size must be run. This function not only calculate power for each sample size but also calculate power for multiple sample sizes varying continuously. By continuously varying sample size across replications, only a single simulation is needed. In this simulation, the sample size for each replication varies randomly across plausible sample sizes (e.g., sample sizes between 200 and 500). For each replication, the sample size and significance of each parameter (0 = not significant, 1 = significant) are recorded. When the simulation is complete, parameter significance is regressed on sample size using logistic regression. For a given sample size, the predicted probability from the logistic regression equation is the power to detect an effect at that sample size. This approach can be extended to other randomly varying simulation parameters such as the percentage of missing data, and model parameters.

getPowerFit 45

Value

Data frame containing columns representing values of the randomly varying simulation parameters, and power for model parameters of interest.

Author(s)

Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>), Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

References

Muthen, L. K., & Muthen, B. O. (2002). How to use a Monte Carlo study to decide on sample size and determine power. *Structural Equation Modeling*, 4, 599-620.

See Also

• SimResult to see how to create a simResult object with randomly varying parameters.

Examples

```
## Not run:
# Specify Sample Size by n
loading <- matrix(0, 6, 1)</pre>
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")</pre>
# We will use only 5 replications to save time.
# In reality, more replications are needed.
# Specify both sample size and percent missing completely at random
Output <- sim(NULL, model=CFA.Model, n=seq(100, 200, 20), pmMCAR=c(0, 0.1, 0.2))
summary(Output)
getPower(Output)
getPower(Output, nVal=c(100, 200), pmMCARval=c(0, 0.1, 0.2))
## End(Not run)
```

getPowerFit

Find power in rejecting alternative models based on fit indices criteria

Description

Find the proportion of fit indices that indicate worse fit than a specified cutoffs. The cutoffs may be calculated from getCutoff of the null model.

Usage

```
getPowerFit(altObject, cutoff, revDirec = FALSE, usedFit=NULL, ...)
```

46 getPowerFit

Arguments

altObject SimResult that indicates alternative model that users wish to reject

cutoff Fit indices cutoffs from null model or users. This should be a vector with a

specified fit indices names as the name of vector elements. This argument can be missing if the SimResult is specified in the altObject and the SimResult

of the null model is specified.

revDirec The default is to count the proportion of fit indices that indicates lower fit to the

model, such as how many RMSEA in the alternative model that is worse than

cutoffs. The direction can be reversed by setting as TRUE.

usedFit Vector of names of fit indices that researchers wish to getCutoffs from. The

default is to getCutoffs of all fit indices.

... Additional arguments

Value

List of power given different fit indices.

Methods

signature(altObject="data.frame", cutoff="vector") This method will find the fit indices indicated in the altObject that provides worse fit than the cutoff. The additional arguments are predictor, predictorVal, condCutoff, and df, which allows the fit indices predicted by any arbitrary independent variables (such as sample size or percent MCAR). The predictor is the data. frame of the predictor values. The number of rows of the predictor argument should be equal to the number of rows in the object. The predictorVal is the values of predictor that researchers would like to find the power from. The condCutoff is a logical. If TRUE, the cutoff is applicable only a given value of predictorVal. If FALSE, the cutoff is applicable in any values of predictor. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

signature(altObject="matrix", cutoff="vector") The details are similar to the method for altObject="data.frame"
and cutoff="vector".

signature(altObject="SimResult", cutoff="vector") This method will find the fit indices indicated in the altObject that provides worse fit than the cutoff. The additional arguments are nVal, pmMCARval, pmMARval, condCutoff, and df, which are needed when using varying sample sizes or percent missing across replications in SimResult. The nVal is the sample size value that researchers wish to find the fit indices cutoffs from. The pmMCARval is the percent missing completely at random value that researchers wish to find the fit indices cutoffs from. The pmMARval is the percent missing at random value that researchers wish to find the fit indices cutoffs from. The condCutoff is a logical. If TRUE, the cutoff is applicable only a given set of nVal, pmMCARval, and pmMARval. If FALSE, the cutoff is applicable in any values of sample size and percent missing. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

signature(altObject="SimResult", cutoff="missing") The details are similar to the method for altObject="SimResult" and cutoff="vector". The cutoff argument must not be specified. Rather, the nullObject, which is an additional argument of this method, is required. The nullObject is the SimResult that contains the simulation result from fitting the null model by the data from the null model.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

getPowerFitNested 47

See Also

- getCutoff to find the cutoffs from null model.
- SimResult to see how to create simResult

Examples

```
## Not run:
loading.null <- matrix(0, 6, 1)</pre>
loading.null[1:6, 1] <- NA</pre>
LX.NULL <- bind(loading.null, 0.7)
RPH.NULL <- binds(diag(1))</pre>
RTD <- binds(diag(6))
CFA.Model.NULL <- model(LY = LX.NULL, RPS = RPH.NULL, RTE = RTD, modelType="CFA")
# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output.NULL <- sim(5, n=500, model=CFA.Model.NULL)
Cut.NULL <- getCutoff(Output.NULL, 0.95)</pre>
loading.alt <- matrix(0, 6, 2)</pre>
loading.alt[1:3, 1] <- NA</pre>
loading.alt[4:6, 2] <- NA
LX.ALT <- bind(loading.alt, 0.7)
latent.cor.alt <- matrix(NA, 2, 2)</pre>
diag(latent.cor.alt) <- 1</pre>
RPH.ALT <- binds(latent.cor.alt, "runif(1, 0.7, 0.9)")
CFA.Model.ALT <- model(LY = LX.ALT, RPS = RPH.ALT, RTE = RTD, modelType="CFA")
Output.ALT <- sim(5, n=500, model=CFA.Model.NULL, generate=CFA.Model.ALT)
getPowerFit(Output.ALT, cutoff=Cut.NULL)
Rule.of.thumb <- c(RMSEA=0.05, CFI=0.95, TLI=0.95, SRMR=0.06)
getPowerFit(Output.ALT, cutoff=Rule.of.thumb, usedFit=c("RMSEA", "CFI", "TLI", "SRMR"))
Output.NULL2 <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model.NULL, generate=CFA.Model.NULL)
Output.ALT2 <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model.NULL, generate=CFA.Model.ALT)
getPowerFit(Output.ALT2, nullObject=Output.NULL2, nVal=250)
## End(Not run)
```

 ${\tt getPowerFitNested}$

Find power in rejecting nested models based on the differences in fit indices

Description

Find the proportion of the difference in fit indices that indicate worse fit than a specified (or internally derived) cutoffs.

Usage

```
getPowerFitNested(altNested, altParent, cutoff, ...)
```

48 getPowerFitNested

Arguments

altNested SimResult that saves the simulation result of the nested model when the nested

model is FALSE.

altParent SimResult that saves the simulation result of the parent model when the nested

model is FALSE.

cutoff A vector of priori cutoffs for fit indices.

... Additional arguments

Value

List of power given different fit indices.

Methods

signature(altNested="SimResult", altParent="SimResult", cutoff="vector") This method will find the the differences in fit indices from altNested and altParent that provides worse fit than the cutoff. The additional arguments are revDirec, usedFit, nVal, pmMCARval, pmMARval, condCutoff, and df, which are needed when using varying sample sizes or percent missing across replications in SimResult. The revDirec is whether to reverse a direction. The default is to count the proportion of fit indices that indicates lower fit to the model, such as how many RMSEA in the alternative model that is worse than cutoffs. The direction can be reversed by setting as TRUE. The usedFit is the vector of names of fit indices that researchers wish to get power from. The default is to get the powers of all fit indices. The nVal is the sample size value that researchers wish to find the fit indices cutoffs from. The pmMCARval is the percent missing completely at random value that researchers wish to find the fit indices cutoffs from. The pmMARval is the percent missing at random value that researchers wish to find the fit indices cutoffs from. The condCutoff is a logical. If TRUE, the cutoff is applicable only a given set of nVal, pmMCARval, and pmMARval. If FALSE, the cutoff is applicable in any values of sample size and percent missing. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

signature(altNested="SimResult", altParent="SimResult", cutoff="missing") The details are
similar to the method for altNested="SimResult", altParent="SimResult", and cutoff="vector".
The cutoff argument must not be specified. Rather, the nullNested and nullParent, which
are additional arguments of this method, are required. The nullNested is the SimResult
that saves the simulation result of the nested model when the nested model is TRUE. The
nullParent is the SimResult that saves the simulation result of the parent model when the
nested model is TRUE.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- getCutoff to find the cutoffs from null model.
- SimResult to see how to create simResult

getPowerFitNonNested 49

Examples

```
## Not run:
loading.null <- matrix(0, 6, 1)</pre>
loading.null[1:6, 1] <- NA</pre>
LX.NULL <- bind(loading.null, 0.7)
RPH.NULL <- binds(diag(1))</pre>
RTD <- binds(diag(6))
CFA.Model.NULL <- model(LY = LX.NULL, RPS = RPH.NULL, RTE = RTD, modelType="CFA")
loading.alt <- matrix(0, 6, 2)</pre>
loading.alt[1:3, 1] \leftarrow NA
loading.alt[4:6, 2] <- NA
LX.ALT <- bind(loading.alt, 0.7)
latent.cor.alt <- matrix(NA, 2, 2)</pre>
diag(latent.cor.alt) <- 1</pre>
RPH.ALT <- binds(latent.cor.alt, 0.7)
CFA.Model.ALT <- model(LY = LX.ALT, RPS = RPH.ALT, RTE = RTD, modelType="CFA")</pre>
Output.NULL.NULL <- sim(10, n=500, model=CFA.Model.NULL, generate=CFA.Model.NULL)
\label{eq:output.ALT.NULL} $$\operatorname{sim}(10, \ n=500, \ model=CFA.Model.NULL, \ generate=CFA.Model.ALT)$$
Output.NULL.ALT <- sim(10, n=500, model=CFA.Model.ALT, generate=CFA.Model.NULL)
Output.ALT.ALT <- sim(10, n=500, model=CFA.Model.ALT, generate=CFA.Model.ALT)\\
\tt getPowerFitNested(Output.ALT.NULL,\ Output.ALT.ALT,\ nullNested=Output.NULL.\ NULL,\ nullParent=Output.\ NULL.\ ALT,\ nullNested=Output.\ NULL.\ ALT,\ nullParent=Output.\ NULL.\ ALT,\ nullNested=Output.\ NULL.\ ALT,\ nullParent=Output.\ NULL.\ ALT,\ NULL.\ ALT,
getPowerFitNested(Output.ALT.NULL, Output.ALT.ALT, cutoff=c(Chi=3.84, CFI=-0.10))
Output.NULL.NULL2 <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model.NULL, generate=CFA.Model.NULL)
Output.ALT.NULL2 <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model.NULL, generate=CFA.Model.ALT)
\label{eq:output.NULL.ALT2} Output. \texttt{NULL}. \texttt{ALT2} <- \texttt{sim}(\texttt{NULL}, \ \texttt{n=seq}(50, \ 500, \ 50), \ \texttt{model=CFA.Model.ALT}, \ \texttt{generate=CFA.Model.NULL})
Output.ALT.ALT2 <- sim(NULL, n=seq(50, 500, 50), model=CFA.Model.ALT, generate=CFA.Model.ALT)
getPowerFitNested(Output.ALT.NULL2, Output.ALT.ALT2, nullNested=Output.NULL.NULL2, nullParent=Output.NULL.
getPowerFitNested(Output.ALT.NULL2, Output.ALT.ALT2, cutoff=c(Chi=3.84, CFI=-0.10), nVal = 250)
## End(Not run)
```

getPowerFitNonNested Find power in rejecting non-nested models based on the differences in fit indices

Description

Find the proportion of the difference in fit indices from one model that does not in the range of sampling distribution from another model (reject that the dataset comes from the second model) or indicates worse fit than a specified cutoff.

Usage

```
getPowerFitNonNested(dat2Mod1, dat2Mod2, cutoff, ...)
```

Arguments

dat2Mod1 SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 2

dat2Mod2 SimResult that saves the simulation of analyzing Model 2 by datasets created

from Model 2

cutoff A vector of priori cutoffs for fit indices.

... Additional arguments

Value

List of power given different fit indices.

Methods

signature(dat2Mod1="SimResult", dat2Mod2="SimResult", cutoff="vector") This method will find the the differences in fit indices from dat2Mod1 and dat2Mod2 that provides worse fit than the cutoff. The additional arguments are revDirec, usedFit, nVal, pmMCARval, pmMARval, condCutoff, and df, which are needed when using varying sample sizes or percent missing across replications in SimResult. The revDirec is whether to reverse a direction. The default is to count the proportion of the difference of fit indices that lower than the specified cutoffs, such as how many the difference in RMSEA in the alternative model that is lower than cutoffs. The direction can be reversed by setting as TRUE. The usedFit is the vector of names of fit indices that researchers wish to get power from. The default is to get the powers of all fit indices. The nVal is the sample size value that researchers wish to find the fit indices cutoffs from. The pmMCARval is the percent missing completely at random value that researchers wish to find the fit indices cutoffs from. The pmMARval is the percent missing at random value that researchers wish to find the fit indices cutoffs from. The condCutoff is a logical. If TRUE, the cutoff is applicable only a given set of nVal, pmMCARval, and pmMARval. If FALSE, the cutoff is applicable in any values of sample size and percent missing. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

signature(dat2Mod1="SimResult", dat2Mod2="SimResult", cutoff="missing") The details are similar to the method for dat2Mod1="SimResult", dat2Mod2="SimResult", and cutoff="vector". The cutoff argument must not be specified. Rather, the dat1Mod1 and dat1Mod2, which are additional arguments of this method, are required. The dat1Mod1 is the SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 1. The dat1Mod2 is the SimResult that saves the simulation of analyzing Model 2 by datasets created from Model 1. The another additional argument is onetailed that is to derive the cutoff by using one-tailed test if specified as TRUE.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- getCutoffNonNested to find the cutoffs for non-nested model comparison
- SimResult to see how to create simResult

```
## Not run:
loading.A <- matrix(0, 8, 2)
loading.A[1:3, 1] <- NA
loading.A[4:8, 2] <- NA
LX.A <- bind(loading.A, 0.7)</pre>
```

imposeMissing 51

```
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, "runif(1, 0.7, 0.9)")
RTD <- binds(diag(8))
CFA.Model.A <- model(LY = LX.A, RPS = RPH, RTE = RTD, modelType="CFA")</pre>
loading.B <- matrix(0, 8, 2)</pre>
loading.B[1:4, 1] <- NA
loading.B[5:8, 2] <- NA
LX.B <- bind(loading.B, 0.7)
CFA.Model.B <- model(LY = LX.B, RPS = RPH, RTE = RTD, modelType="CFA")
# The actual number of replications should be greater than 10.
Output.A.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.A)
Output.A.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.A)
Output.B.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.B)
Output.B.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.B)
getPowerFitNonNested(Output.B.A, Output.B.B, dat1Mod1=Output.A.A, dat1Mod2=Output.A.B)
getPowerFitNonNested(Output.B.A, Output.B.B, cutoff=c(AIC=0, BIC=0))
## End(Not run)
```

imposeMissing

Impose MAR, MCAR, planned missingness, or attrition on a data set

Description

Function imposes missing values on a data based on the known missing data types, including MCAR, MAR, planned, and attrition.

Usage

```
impose(miss, data.mat, pmMCAR = NULL, pmMAR = NULL)
imposeMissing(data.mat, cov = 0, pmMCAR = 0, pmMAR = 0, nforms = 0,
itemGroups = list(), twoMethod = 0, prAttr = 0, timePoints = 1,
ignoreCols = 0, threshold = 0, logical = NULL)
```

Arguments

miss	Missing data object (SimMissing) used as the template for impose missing values
data.mat	Data to impose missing upon. Can be either a matrix or a data frame.
cov	Column indices of a covariate to be used to impose MAR missing, or MAR attrition. Will not be included in any removal procedure. See details.
pmMCAR	Decimal percent of missingness to introduce completely at random on all variables.
pmMAR	Decimal percent of missingness to introduce using the listed covariate as predictor. See details.
nforms	The number of forms for planned missing data designs, not including the shared form.

52 imposeMissing

itemGroups List of lists of item groupings for planned missing data forms. Unless specified, items will be divided into groups sequentially (e.g. 1-3,4-6,7-9,10-12) twoMethod With missing on one variable: vector of (column index, percent missing). Will put a given percent missing on that column in the matrix to simulate a two method planned missing data research design. With missing on two or more variables: list of (column indices, percent missing). Probability (or vector of probabilities) of an entire case being removed due to prAttr attrition at a given time point. When a covariate is specified along with this argument, attrition will be predicted by the covariate (MAR attrition). See details. timePoints Number of timepoints items were measured over. For longitudinal data, planned missing designs will be implemented within each timepoint. All methods to impose missing values over time assume an equal number of variables at each time point. The columns not imposed any missing values for any missing data patterns. ignoreCols threshold The threshold of the covariate used to impose missing values. Values on the covariate above this threshold are eligible to be deleted. The default threshold is the mean of the variable. logical A matrix of logical values (TRUE/FALSE). If a value in the dataset is corresponding to the TRUE in the logical matrix, the value will be missing.

Details

Without specifying any arguments, no missing values will be introduced.

A single covariate is required to specify MAR missing - this covariate can be distributed in any way. This covariate can be either continuous or categorical, as long as it is numerical. If the covariate is categorical, the threshold should be specified to one of the levels.

MAR missingness is specified using the threshold method - any value on the covariate that is above the specified threshold indicates a row eligible for deletion. If the specified total amount of MAR missingness is not possible given the total rows eligible based on the threshold, the function iteratively lowers the threshold until the total percent missing is possible.

Planned missingness is parameterized by the number of forms (n). This is used to divide the cases into n groups. If the column groupings are not specified, a naive method will be used that divides the columns into n+1 equal forms sequentially (1-4,5-9,10-13..), where the first group is the shared form. The first list of column indices given will be used as the shared group. If this is not desired, this list can be left empty.

For attrition, the probability can be specified as a single value or as a vector. For a single value, the probability of attrition will be the same across time points, and affects only cases not previously lost due to attrition. If this argument is a vector, this specifies different probabilities of attrition for each time point. Values will be recycled if this vector is smaller than the specified number of time points.

An MNAR processes can be generated by specifying MAR missingness and then dropping the covariate from the subsequent analysis.

Currently, if MAR missing is imposed along with attrition, both processes will use the same covariate and threshold.

Currently, all types of missingness (MCAR, MAR, planned, and attrition) are imposed independently. This means that specified global values of percent missing will not be additive (10 percent MCAR + 10 percent MAR does not equal 20 percent total missing).

interpolate 53

Value

A data matrix with NAs introduced in the way specified by the arguments.

Author(s)

Patrick Miller(University of Kansas; <patr1ckm@ku.edu>) Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>)

See Also

SimMissing for the alternative way to save missing data feature for using in the sim function.

Examples

```
data <- matrix(rep(rnorm(10,1,1),19),ncol=19)
datac <- cbind(data,rnorm(10,0,1),rnorm(10,5,5))

# Imposing Missing with the following arguments produces no missing values
imposeMissing(data)
imposeMissing(data,cov=c(1,2))
imposeMissing(data,pmMCAR=0)
imposeMissing(data,nforms=0)

#Some more usage examples
imposeMissing(data,cov=c(1,2),pmMCAR=.1)

imposeMissing(data,nforms=3)
imposeMissing(data,nforms=3,itemGroups=list(c(1,2,3,4,5),c(6,7,8,9,10),c(11,12,13,14,15),c(16,17,18,19)))
imposeMissing(data,twoMethod=c(19,.8))
imposeMissing(data,cov=21,prAttr=.1,timePoints=5)</pre>
```

interpolate

Find the value of one vector relative to a value of another vector by interpolation

Description

Find the value of the resulting vector that have the position similar to the value of the baseline vector. If the starting value in the baseline vector is in between two elements, the resulting value will be predicted by linear interpolation.

Usage

```
interpolate(baselineVec, val, resultVec=NULL)
```

54 likRatioFit

Arguments

baselineVec The target vector to be used as a baseline. The resulting vector can be attached

as the element names of this vector.

val The value relative to the baseline vector to be used for projecting the resulting

value

resultVec The vector that the resulting value will be used to base their result form

Value

The interpolated value from the resulting vector relative to the value in the baseline vector

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

No Example

likRatioFit Find the likelihood ratio (or Bayes factor) based on the bivariate distribution of fit indices

Description

Find the log-likelihood of the observed fit indices on Model 1 and 2 from the real data on the bivariate sampling distribution of fit indices fitting Model 1 and Model 2 by the datasets from the Model 1 and Model 2. Then, the likelihood ratio is computed (which may be interpreted as posterior odd). If the prior odd is 1 (by default), the likelihood ratio is equivalent to Bayes Factor.

Usage

```
likRatioFit(outMod1, outMod2, dat1Mod1, dat1Mod2, dat2Mod1, dat2Mod2,
usedFit=NULL, prior=1)
```

Arguments

outMod1 outMod2	lavaan that saves the analysis result of the first model from the target dataset lavaan that saves the analysis result of the second model from the target dataset
dat1Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 1
dat1Mod2	SimResult that saves the simulation of analyzing Model 2 by datasets created from Model 1
dat2Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 2
dat2Mod2	SimResult that saves the simulation of analyzing Model 2 by datasets created from Model 2
usedFit	Vector of names of fit indices that researchers wish to getCutoffs from. The default is to getCutoffs of all fit indices.
prior	The prior odds. The prior probability that Model 1 is correct over the prior probability that Model 2 is correct.

likRatioFit 55

Value

The likelihood ratio (Bayes Factor) in preference of Model 1 to Model 2. If the value is greater than 1, Model 1 is preferred. If the value is less than 1, Model 2 is preferred.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

SimResult for a detail of simResult pValueNested for a nested model comparison by the difference in fit indices pValueNonNested for a nonnested model comparison by the difference in fit indices

```
## Not run:
library(lavaan)
loading <- matrix(0, 11, 3)</pre>
loading[1:3, 1] <- NA</pre>
loading[4:7, 2] \leftarrow NA
loading[8:11, 3] <- NA
path.A <- matrix(0, 3, 3)</pre>
path.A[2:3, 1] <- NA
path.A[3, 2] <- NA
model.A <- estmodel(LY=loading, BE=path.A, modelType="SEM", indLab=c(paste("x", 1:3, sep=""), paste("y", 1</pre>
out.A <- analyze(model.A, PoliticalDemocracy)</pre>
path.B <- matrix(0, 3, 3)
path.B[1:2, 3] <- NA
path.B[1, 2] <- NA
model.B <- estmodel(LY=loading, BE=path.B, modelType="SEM", indLab=c(paste("x", 1:3, sep=""), paste("y", 1</pre>
out.B <- analyze(model.B, PoliticalDemocracy)</pre>
loading.mis <- matrix("runif(1, -0.2, 0.2)", 11, 3)</pre>
loading.mis[is.na(loading)] <- 0</pre>
datamodel.A <- model.lavaan(out.A, std=TRUE, LY=loading.mis)</pre>
datamodel.B <- model.lavaan(out.B, std=TRUE, LY=loading.mis)</pre>
n <- nrow(PoliticalDemocracy)</pre>
output.A.A <- sim(20, n=n, model.A, generate=datamodel.A)</pre>
output.A.B <- sim(20, n=n, model.B, generate=datamodel.A)
output.B.A <- sim(20, n=n, model.A, generate=datamodel.B)
output.B.B <- sim(20, n=n, model.B, generate=datamodel.B)</pre>
# The output may contain some warnings here. When the number of replications increases (e.g., 1000), the w
likRatioFit(out.A, out.B, output.A.A, output.A.B, output.B.A, output.B.B)
## End(Not run)
```

56 miss

loadingFromAlpha

Find standardized factor loading from coefficient alpha

Description

Find standardized factor loading from coefficient alpha assuming that all items have equal loadings.

Usage

```
loadingFromAlpha(alpha, ni)
```

Arguments

alpha A desired coefficient alpha value.

ni A desired number of items.

Value

result The standardized factor loadings that make desired coefficient alpha with speci-

fied number of items.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
loadingFromAlpha(0.8, 4)
```

miss

TBA

Description

TBA

Usage

```
miss(cov = 0, pmMCAR = 0, pmMAR = 0, nforms = 0, itemGroups = list(),
timePoints = 1, twoMethod = 0, prAttr = 0, package="default", ignoreCols = 0,
threshold = 0, covAsAux = TRUE, logical = NULL, ...)
```

57 miss

Arguments

cov	Column indices of any normally distributed covariates used in the data set.
pmMCAR	Decimal percent of missingness to introduce completely at random on all variables.
pmMAR	Decimal percent of missingness to introduce using the listed covariates as predictors.
nforms	The number of forms for planned missing data designs, not including the shared form.
itemGroups	List of lists of item groupings for planned missing data forms. Without this, items will be divided into groups sequentially (e.g. 1-3,4-6,7-9,10-12)
timePoints	Number of timepoints items were measured over. For longitudinal data, planned missing designs will be implemented within each timepoint.
twoMethod	With missing on one variable: vector of (column index, percent missing). Will put a given percent missing on that column in the matrix to simulate a two method planned missing data research design. With missing on two or more variables: list of (column indices, percent missing).
prAttr	Probability (or vector of probabilities) of an entire case being removed due to attrition at a given time point. See <pre>imposeMissing</pre> for further details.
package	TBA
ignoreCols	The columns not imposed any missing values for any missing data patterns
threshold	The threshold of covariates that divide between the area to impose missing and the area not to impose missing. The default threshold is the mean of the covariate.
covAsAux	If TRUE, the covariate listed in the object will be used as auxiliary variables when putting in the model object. If FALSE, the covariate will be included in the analysis.
logical	A matrix of logical values (TRUE/FALSE). If a value in the dataset is corresponding to the TRUE in the logical matrix, the value will be missing.
	TBA

Details

TBA

Value

TBA

Author(s)

Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>), Patrick Miller (Univeristy of Notre Dame; <pmille13@nd.edu>), Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimMissing for the alternative way to save missing data feature for using in the sim function.

58 model

Examples

```
#Example of imposing 10% MCAR missing in all variables with no imputations (FIML method)
Missing <- miss(pmMCAR=0.1)</pre>
summary(Missing)
loading <- matrix(0, 6, 1)</pre>
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))</pre>
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
#Create data
dat <- generate(CFA.Model, n = 20)</pre>
#Impose missing
#dat <- run(Missing, dat)</pre>
#Analyze data
#out <- run(SimModel, dat)</pre>
#summary(out)
#Example to create simMissing object for 3 forms design at 3 timepoints with 10 imputations
Missing <- miss(nforms=3, timePoints=3, numImps=10)</pre>
```

model

Data generation template and analysis template for simulation.

Description

Creates a data generation and analysis template (lavaan parameter table) for simulations with structural equation models based on Y-side LISREL design matrices. Each corresponds to a LISREL matrix, but must be a SimMatrix or SimVector built using bind. In addition to the usual Y-side matrices in LISREL, both PS and TE can be specified using correlations (RPS, RTE) and scaled by a vector of residual variances (VTE, VPS) or total variances (VY, VE). Multiple groups are supported by passing lists of SimMatrix or SimVector to arguments, or by specifying the number of groups.

Usage

```
model(LY = NULL, PS = NULL, RPS = NULL, TE = NULL, RTE = NULL, BE = NULL,
VTE = NULL, VY = NULL, VPS = NULL, VE = NULL, TY = NULL, AL = NULL, MY = NULL,
ME = NULL, modelType, indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
model.cfa(LY = NULL, PS = NULL, RPS = NULL, TE = NULL, RTE = NULL, VTE = NULL,
VY = NULL, VPS = NULL, VE=NULL, TY = NULL, AL = NULL, MY = NULL, ME = NULL,
indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
model.path(PS = NULL, RPS = NULL, BE = NULL, VPS = NULL, VE=NULL, AL = NULL,
ME = NULL, indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
model.sem(LY = NULL, PS = NULL, RPS = NULL, TE = NULL, RTE = NULL, BE = NULL,
VTE = NULL, VY = NULL, VPS = NULL, VE=NULL, TY = NULL, AL = NULL,
ME = NULL, indLab=NULL, facLab=NULL, groupLab="group", ngroups=1, smartStart=TRUE)
```

model 59

Arguments

LY Factor loading matrix from endogenous factors to Y indicators (need to be SimMatrix object). Residual covariance matrix among endogenous factors (need to be SimMatrix PS object). **RPS** Residual correlation matrix among endogenous factors (need to be SimMatrix object). ΤE Measurement error covariance matrix among Y indicators (need to be SimMatrix object). Measurement error correlation matrix among Y indicators (need to be SimMatrix **RTE** object). BF Regression coefficient matrix among endogenous factors (need to be SimMatrix object). VTE Measurement error variance of indicators (need to be SimVector object). Total variance of indicators (need to be SimVector object). NOTE: Either mea-VY surement error variance or indicator variance is specified. Both cannot be simultaneously specified. **VPS** Residual variance of factors (need to be SimVector object). ٧E Total variance of of factors (need to be SimVector object). NOTE: Either residual variance of factors or total variance of factors is specified. Both cannot be simulatneously specified. TY Measurement intercepts of Y indicators. (need to be SimVector object). Endogenous factor intercept (need to be SimVector object). ΑL MY Overall Y indicator means. (need to be SimVector object). NOTE: Either measurement intercept of indicator mean can be specified. Both cannot be specified simultaneously. ME Total mean of endogenous factors (need to be SimVector object). NOTE: Either endogenous factor intercept or total mean of endogenous factor is specified. Both cannot be simultaneously specified. "CFA", "Sem", or "Path". This is specified to ensure that the analysis and data modelType generation template created based on specified matrices in model correspond to what the user intends. indLab Character vector of indicator labels. If left blank, automatic labels will be generated as y1, y2, ... yy. facLab Character vector of factor labels. If left blank, automatic labels will be generated as f1, f2, ... ff Character of group-variable label (not the names of each group). If left blank, groupLab automatic labels will be generated as group Integer. Number of groups for data generation, defaults to 1. If larger than one, ngroups all specified matrices will be repeated for each additional group. If any matrix argument is a list, the length of this list will be the number of groups and ngroups is ignored. Defaults to FALSE. If TRUE, population parameter values that are real numbers smartStart

will be used as starting values.

60 model

Details

The *simsem* package is intricately tied to the *lavaan* package for analysis of structural equation models. The analysis template that is generated by model is a lavaan parameter table, a low-level access point to lavaan that allows repeated analyses to happen more rapidly. If desired, the parameter table generated can be used directly with lavaan for many analyses.

The data generation template is simply a list of SimMatrix or SimVector. The SimSem object can be passed to the function generate to generate data.

If multiple group data is desired, the user can optionally either specify the number of groups argument, or pass a list of SimMatrix or SimVector to any of the matrix arguments. The length of this list will be the number of groups. If only one argument is a list, all other arguments will be automatically replicated to that length, parameters will be identified in the same way, have the same population parameter value/distribution, and have the same misspecification. If only ngroups is specified, all arguments will be replicated in this fashion. If equality constraints are present during the automatic replication, these parameters will be constrained to be equal across groups.

The model.cfa, model.path, and model.sem are the shortcuts for the model function when modelType are "CFA", "Path", and "SEM", respectively.

Value

SimSem object that contains the data generation template (@dgen) and analysis template (@pt).

Author(s)

See Also

- sim for simulations using the SimSem template.
- generate To generate data using the SimSem template.
- analyze To analyze real or generated data using the SimSem template.
- draw To draw parameters using the SimSem template.

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
LY <- bind(loading, 0.7)

latent.cor <- matrix(NA, 2, 2)
diag(latent.cor) <- 1
RPS <- binds(latent.cor, 0.5)

RTE <- binds(diag(6))

VY <- bind(rep(NA,6),2)

CFA.Model <- model(LY = LY, RPS = RPS, RTE = RTE, modelType = "CFA")</pre>
```

model.lavaan 61

model.lavaan	Build the data generation template and analysis template from the lavaan result

Description

Creates a data generation and analysis template (lavaan parameter table) for simulations with the lavaan result. Model misspecification may be added into the template by a vector, a matrix, or a list of vectors or matrices (for multiple groups).

Usage

```
model.lavaan(object, std = FALSE, LY = NULL, PS = NULL, RPS = NULL,
TE = NULL, RTE = NULL, BE = NULL, VTE = NULL, VY = NULL, VPS = NULL,
VE=NULL, TY = NULL, AL = NULL, MY = NULL, ME = NULL, smartStart=TRUE)
```

Arguments

object	A lavaan object to be used to build the data generation and analysis template.
std	If TRUE, use the resulting standardized parameters for data generation. If FALSE, use the unstandardized parameters for data generation.
LY	Model misspecification in factor loading matrix from endogenous factors to Y indicators (need to be a matrix or a list of matrices).
PS	Model misspecification in residual covariance matrix among endogenous factors (need to be a symmetric matrix or a list of symmetric matrices).
RPS	Model misspecification in residual correlation matrix among endogenous factors (need to be a symmetric matrix or a list of symmetric matrices).
TE	Model misspecification in measurement error covariance matrix among Y indicators (need to be a symmetric matrix or a list of symmetric matrices).
RTE	Model misspecification in measurement error correlation matrix among Y indicators (need to be a symmetric matrix or a list of symmetric matrices).
BE	Model misspecification in regression coefficient matrix among endogenous factors (need to be a symmetric matrix or a list of symmetric matrices).
VTE	Model misspecification in measurement error variance of indicators (need to be a vector or a list of vectors).
VY	Model misspecification in total variance of indicators (need to be a vector or a list of vectors). NOTE: Either measurement error variance or indicator variance is specified. Both cannot be simultaneously specified.
VPS	Model misspecification in residual variance of factors (need to be a vector or a list of vectors).
VE	Model misspecification in total variance of of factors (need to be a vector or a list of vectors). NOTE: Either residual variance of factors or total variance of factors is specified. Both cannot be simulatneously specified.
TY	Model misspecification in measurement intercepts of Y indicators. (need to be a vector or a list of vectors).
AL	Model misspecification in endogenous factor intercept (need to be a vector or a list of vectors).

62 multipleAllEqual

MY	Model misspecification in overall Y indicator means. (need to be a vector or a
	list of vectors). NOTE: Either measurement intercept of indicator mean can be
	specified. Both cannot be specified simultaneously.

Model misspecification in total mean of endogenous factors (need to be a vector or a list of vectors). NOTE: Either endogenous factor intercept or total mean of

endogenous factor is specified. Both cannot be simultaneously specified.

smartStart Defaults to FALSE. If TRUE, population parameter values that are real numbers

will be used as starting values.

Value

ME

SimSem object that contains the data generation template (@dgen) and analysis template (@pt).

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- model To build data generation and data analysis template for simulation.
- sim for simulations using the SimSem template.
- generate To generate data using the SimSem template.
- analyze To analyze real or generated data using the SimSem template.
- draw To draw parameters using the SimSem template.

Examples

multipleAllEqual

Test whether all objects are equal

Description

Test whether all objects are equal. The test is based on the all.equal function.

Usage

```
multipleAllEqual(...)
```

Arguments

... The target objects

overlapHist 63

Value

TRUE if all objects are equal.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
multipleAllEqual(1:5, 1:5, seq(2, 10, 2)/2)
multipleAllEqual(1:5, 1:6, seq(2, 10, 2)/2)
```

overlapHist

Plot overlapping histograms

Description

Plot overlapping histograms

Usage

```
overlapHist(a, b, colors=c("red","blue","purple"), breaks=NULL, xlim=NULL,
ylim=NULL, main=NULL, xlab=NULL, swap=FALSE)
```

Arguments

a	Data for the first histogram
b	Data for the second histogram
colors	Colors for the first histogram, the second histogram, and the overlappling areas.
breaks	How many breaks users used in each histogram (should not be used)
xlim	The range of x-axis
ylim	The range of y-axis
main	The title of the figure
xlab	The labels of x-axis
swap	Specify TRUE to plot b first and then a. The default is FALSE to plot a first and then b.

Value

None. This function will plot only.

Author(s)

Chris Miller provided this code on http://chrisamiller.com/science/2010/07/20/transparent-overlapping-h The code is modified by Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

64 plot3DQtile

Examples

```
# This function is not a public function.
# a <- rnorm(10000, 0, 1)
# b <- rnorm(10000, 1, 1.5)
# overlapHist(a, b, main="Example")</pre>
```

plot3DQtile

Build a persepctive plot or contour plot of a quantile of predicted val-

ues

Description

Build a persepctive plot or contour plot of a quantile of predicted values

Usage

```
plot3DQtile(x, y, z, df=0, qtile=0.5, useContour=TRUE, xlab=NULL,
ylab=NULL, zlab=NULL, main=NULL)
```

Arguments

x	The values of the first variable (e.g., a vector of sample size)
у	The values of the second variable (e.g., a vector of percent missing)
z	The values of the dependent variable
df	The degree of freedom in spline method
qtile	The quantile values used to plot a graph
useContour	If TRUE, use contour plot. If FALSE, use perspective plot.
xlab	The labels of x-axis
ylab	The labels of y-axis
zlab	The labels of z-axis
main	The title of the graph

Value

None. This function will plot only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

```
# No example
```

plotCutoff 65

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Plot sampling distributions of fit indices with fit indices cutoffs

Description

This function will plot sampling distributions of null hypothesis fit indices. The users may add cutoffs by specifying the alpha level.

Usage

```
plotCutoff(object, ...)
```

Arguments

object

The object (SimResult or data.frame) that contains values of fit indices in

each distribution.

... Other arguments specific to different types of object you pass in the function.

Value

NONE. Only plot the fit indices distributions.

Details in ...

- cutoff: A priori cutoffs for fit indices, saved in a vector
- cutoff2: Another set of priori cutoffs for fit indices, saved in a vector
- alpha: A priori alpha level to getCutoffs of fit indices (do not specify when you have cutoff)
- revDirec: The default is to find critical point on the side that indicates worse fit (the right side of RMSEA or the left side of CFI). If specifying as TRUE, the directions are reversed.
- usedFit: The name of fit indices that researchers wish to plot
- useContour: If there are two of sample size, percent completely at random, and percent missing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective plot is used.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- SimResult for simResult that used in this function.
- getCutoff to find values of cutoffs based on null hypothesis sampling distributions only

66 plotCutoffNested

Examples

```
## Not run:
loading <- matrix(0, 6, 2)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loadingValues <- matrix(0, 6, 2)</pre>
loadingValues[1:3, 1] <- 0.7</pre>
loadingValues[4:6, 2] <- 0.7</pre>
LX <- bind(loading, loadingValues)
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, 0.5)
error.cor <- matrix(0, 6, 6)
diag(error.cor) <- 1</pre>
RTD <- binds(error.cor)</pre>
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output <- sim(5, n=200, model=CFA.Model)
plotCutoff(Output,\ 0.05,\ usedFit=c("RMSEA",\ "SRMR",\ "CFI",\ "TLI"))
# Varying N
Output2 <- sim(NULL, n=seq(450, 500, 10), model=CFA.Model)
plotCutoff(Output2, 0.05)
# Varying N and pmMCAR
Output3 <- sim(NULL, n=seq(450, 500, 10), pmMCAR=c(0, 0.05, 0.1, 0.15), model=CFA.Model)
plotCutoff(Output3, 0.05)
## End(Not run)
```

plotCutoffNested

Plot sampling distributions of the differences in fit indices between nested models with fit indices cutoffs

Description

This function will plot sampling distributions of the differences in fit indices between nested models if the nested model is true. The users may add cutoffs by specifying the alpha level.

Usage

```
plotCutoffNested(nested, parent, alpha = 0.05, cutoff = NULL,
usedFit = NULL, useContour = T)
```

Arguments

nested	SimResult that saves the analysis results of nested model from multiple replications
parent	SimResult that saves the analysis results of parent model from multiple replications

alpha A priori alpha level

plotCutoffNested 67

cutoff A priori cutoffs for fit indices, saved in a vector

usedFit Vector of names of fit indices that researchers wish to plot the sampling distri-

bution.

useContour If there are two of sample size, percent completely at random, and percent miss-

ing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective

plot is used.

Value

NONE. Only plot the fit indices distributions.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- SimResult for simResult that used in this function.
- getCutoffNested to find the difference in fit indices cutoffs

```
## Not run:
loading.null <- matrix(0, 6, 1)</pre>
loading.null[1:6, 1] <- NA</pre>
LX.NULL <- bind(loading.null, 0.7)
RPH.NULL <- binds(diag(1))</pre>
RTD <- binds(diag(6))
CFA.Model.NULL <- model(LY = LX.NULL, RPS = RPH.NULL, RTE = RTD, modelType="CFA")
loading.alt <- matrix(0, 6, 2)</pre>
loading.alt[1:3, 1] <- NA</pre>
loading.alt[4:6, 2] <- NA</pre>
LX.ALT <- bind(loading.alt, 0.7)
latent.cor.alt <- matrix(NA, 2, 2)</pre>
diag(latent.cor.alt) <- 1</pre>
RPH.ALT <- binds(latent.cor.alt, "runif(1, 0.7, 0.9)")
CFA.Model.ALT <- model(LY = LX.ALT, RPS = RPH.ALT, RTE = RTD, modelType="CFA")
# The actual number of replications should be greater than 10.
Output.NULL.NULL <- sim(10, n=500, model=CFA.Model.NULL)</pre>
Output.NULL.ALT <- sim(10, n=500, model=CFA.Model.ALT, generate=CFA.Model.NULL)
plotCutoffNested(Output.NULL.NULL, Output.NULL.ALT, alpha=0.05)
## End(Not run)
```

68 plotCutoffNonNested

plotCutoffNonNested	Plot sampling distributions of the differences in fit indices between non-nested models with fit indices cutoffs

Description

This function will plot sampling distributions of the differences in fit indices between non-nested models. The users may add cutoffs by specifying the alpha level.

Usage

```
plotCutoffNonNested(dat1Mod1, dat1Mod2, dat2Mod1=NULL, dat2Mod2=NULL,
alpha=0.05, cutoff = NULL, usedFit = NULL, useContour = T, onetailed=FALSE)
```

Arguments

dat1Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 1
dat1Mod2	SimResult that saves the simulation of analyzing Model 2 by datasets created from Model 1
dat2Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 2
dat2Mod2	SimResult that saves the simulation of analyzing Model 2 by datasets created from Model 2
alpha	A priori alpha level
cutoff	A priori cutoffs for fit indices, saved in a vector
usedFit	Vector of names of fit indices that researchers wish to plot the sampling distribution.
useContour	If there are two of sample size, percent completely at random, and percent missing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective plot is used.
onetailed	If TRUE, the function will find the cutoff from one-tail test. If FALSE, the function will find the cutoff from two-tailed test.

Value

NONE. Only plot the fit indices distributions.

Author(s)

 $Sunthud\ Pornprasert manit\ (University\ of\ Kansas; < psunthud@ku.edu>)$

See Also

- SimResult for simResult that used in this function.
- getCutoffNonNested to find the difference in fit indices cutoffs for non-nested model comparison

plotDist 69

Examples

```
## Not run:
loading.A <- matrix(0, 8, 2)</pre>
loading.A[1:3, 1] <- NA
loading.A[4:8, 2] <- NA
LX.A <- bind(loading.A, 0.7)
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, "runif(1, 0.7, 0.9)")
RTD <- binds(diag(8))
CFA.Model.A <- model(LY = LX.A, RPS = RPH, RTE = RTD, modelType="CFA")
loading.B <- matrix(0, 8, 2)</pre>
loading.B[1:4, 1] <- NA
loading.B[5:8, 2] <- NA
LX.B <- bind(loading.B, 0.7)
CFA.Model.B <- model(LY = LX.B, RPS = RPH, RTE = RTD, modelType="CFA")
# The actual number of replications should be greater than 10.
Output.A.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.A) \\
Output.B.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.B)\\
Output.B.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.B)\\
plotCutoffNonNested(Output.A.A, Output.A.B, Output.B.A, Output.B.B)
plotCutoffNonNested(Output.A.A, Output.A.B)
plotCutoffNonNested(Output.A.A, Output.A.B, onetailed=TRUE)
## End(Not run)
```

plotDist

Plot a distribution of a data distribution object

Description

Plot a distribution of a data distribution object

Usage

```
plotDist(object, xlim = NULL, ylim = NULL, r = 0, var = NULL, contour = TRUE)
```

Arguments

object	The data distribution object (SimDataDist) to plot a distribution
xlim	A numeric vector with two elements specifying the lower and upper limit of the x-axis to be plotted.
ylim	A numeric vector with two elements specifying the lower and upper limit of the y-axis to be plotted. This argument is applicable for the joint distribution of two dimensions only
r	The correlation of two dimensions in the joint distribution
var	A vector of the index of variables to be plotted. The length of vector cannot be greater than 2.

70 plotIndividualScatter

contour

Applicable if two variables are used only. If TRUE, the contour plot is provided. If FALSE, the perspective plot is provided.

Value

No return value. This function will plot a graph only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimDataDist for plotting a data distribution object

Examples

plotIndividualScatter Plot an overlaying scatter plot visualizing the power of rejecting misspecified models

Description

Plot the fit indices value against the value of predictors. The plot will include the fit indices value of the alternative models, the fit indices value of the null model (if specified), and the fit indices cutoffs (if specified).

Usage

```
plotIndividualScatter(altVec, nullVec=NULL, cutoff=NULL, x, main = NULL)
```

Arguments

altVec	The vector saving the fit index distribution when the hypothesized model is FALSE.
nullVec	The vector saving the fit index distribution when the hypothesized model is TRUE.
cutoff	A priori cutoff
X	The data.frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the object.
main	The title of the graph

Value

NONE. Only plot the fit indices distributions.

plotLogisticFit 71

Author(s)

 $Sunthud\ Pornprasert manit\ (University\ of\ Kansas; < psunthud@ku.edu>)$

See Also

• plotPowerFit

Examples

No example

specified model	plotLogisticFit	Plot multiple logistic curves for predicting whether rejecting a misspecified model
-----------------	-----------------	---

Description

This function will find the fit indices cutoff values if not specified, then check whether the hypothesized model is rejected in each dataset, and plot the logistic curve given the value of predictors.

Usage

```
plotLogisticFit(altObject, nullObject=NULL, cutoff=NULL,
  usedFit=NULL, x, xval, alpha=0.05, useContour=TRUE, df=0)
```

Arguments

altObject	The result object (data.frame) saves the simulation result of fitting the hypothesized model when the hypothesized model is FALSE.
nullObject	The result object (data.frame) saves the simulation result of fitting the hypothesized model when the hypothesized model is TRUE. This argument may be not specified if the cutoff is specified.
cutoff	A vector of priori cutoffs for fit indices.
usedFit	Vector of names of fit indices that researchers wish to plot.
alpha	A priori alpha level
X	The data. frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the object.
xval	The values of predictor that researchers would like to find the fit indices cutoffs from.
useContour	If there are two of sample size, percent completely at random, and percent missing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective plot is used.
df	The degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

Value

NONE. Only plot the fit indices distributions.

72 plotMisfit

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

```
• plotPowerFit
```

Examples

```
# No example
```

plotMisfit

Plot the population misfit in the result object

Description

Plot a histogram of the amount of population misfit in parameter result object or the scatter plot of the relationship between misspecified parameter and the population misfit or the fit indices

Usage

```
plotMisfit(object, usedFit="default", misParam=NULL)
```

Arguments

object The result object, SimResult

usedFit The sample fit indices or population misfit used to plot. All sample fit indices

are available. The available population misfit are "pop.fo", "pop.rmsea", and "pop.srmr". If the misParam is not specified, all population misfit are used. If

the misParam is specified, the "pop.rmsea" is used in the plot.

misParam The index or the name of misspecified parameters used to plot.

Value

None. This function will plot only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

```
path.BE <- matrix(0, 4, 4)
path.BE[3, 1:2] <- NA
path.BE[4, 3] <- NA
starting.BE <- matrix("", 4, 4)
starting.BE[3, 1:2] <- "runif(1, 0.3, 0.5)"
starting.BE[4, 3] <- "runif(1, 0.5, 0.7)"
mis.path.BE <- matrix(0, 4, 4)
mis.path.BE[4, 1:2] <- "runif(1, -0.1, 0.1)"
BE <- bind(path.BE, starting.BE, misspec=mis.path.BE)</pre>
```

plotOverHist 73

```
residual.error <- diag(4)
residual.error[1,2] <- residual.error[2,1] <- NA
RPS <- binds(residual.error, "rnorm(1, 0.3, 0.1)")

ME <- bind(rep(NA, 4), 0)

Path.Model <- model(RPS = RPS, BE = BE, ME = ME, modelType="Path")

# The number of replications in actual analysis should be much more than 5
ParamObject <- sim(20, n=500, Path.Model)
plotMisfit(ParamObject)

plotMisfit(ParamObject, misParam=1:2)</pre>
```

plotOverHist

Plot multiple overlapping histograms

Description

Plot multiple overlapping histograms and find the cutoff values if not specified

Usage

```
\verb|plotOverHist(altObject, nullObject, cutoff=NULL, usedFit=NULL, alpha=0.05, cutoff2=NULL, cutoff3=NULL, cutoff4=NULL)|
```

Arguments

altObject	The result object (data.frame) saves the simulation result of fitting the hypothesized model when the hypothesized model is FALSE.
nullObject	The result object (data.frame) saves the simulation result of fitting the hypothesized model when the hypothesized model is TRUE.
cutoff	A vector of priori cutoffs for fit indices.
usedFit	Vector of names of fit indices that researchers wish to plot.
alpha	A priori alpha level
cutoff2	Another vector of priori cutoffs for fit indices.
cutoff3	A vector of priori cutoffs for fit indices for the altObject.
cutoff4	Another vector of priori cutoffs for fit indices for the altObject.

Value

NONE. Only plot the fit indices distributions.

Author(s)

 $Sunthud\ Pornprasert manit\ (University\ of\ Kansas; < psunthud@ku.edu>)$

See Also

• plotPowerFit

74 plotPower

Examples

No example

plotPower	Make a power plot of a parameter given varying parameters

Description

Make a power plot of a parameter given varying parameters (e.g., sample size, percent missing completely at random, or random parameters in the model)

Usage

```
plotPower(object, powerParam, alpha = 0.05, contParam = NULL, contN = TRUE,
contMCAR = TRUE, contMAR = TRUE, useContour=TRUE)
```

Arguments

object	SimResult that includes at least one randomly varying parameter (e.g. sample size, percent missing, model parameters)
powerParam	Vector of parameters names that the user wishes to find power for. This can be a vector of names (e.g., "LY1_1", "LY2_2").
alpha	Alpha level to use for power analysis.
contParam	Vector of parameters names that vary over replications that users wish to use in the plot.
contN	Include the varying sample size in the power plot if available
contMCAR	Include the varying MCAR (missing completely at random percentage) in the power plot if available
contMAR	Include the varying MAR (missing at random percentage) in the power plot if available
useContour	This argument is used when users specify to plot two varying parameters. If TRUE, the contour plot is used. If FALSE, perspective plot is used.

Details

Predicting whether each replication is significant or not by varying parameters using logistic regression (without interaction). Then, plot the logistic curves predicting the probability of significance against the target varying parameters.

Value

Not return any value. This function will plot a graph only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>), Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>)

plotPowerFit 75

See Also

- SimResult to see how to create a simResult object with randomly varying parameters.
- getPower to obtain a statistical power given varying parameters values.

Examples

```
## Not run:
# Specify Sample Size by n
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, 0.4)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")

# Specify both sample size and percent missing completely at random
Output <- sim(NULL, n=seq(100, 200, 20), pmMCAR=c(0, 0.1, 0.2), model=CFA.Model)
plotPower(Output, "1.LY1_1", contMCAR=FALSE)

## End(Not run)</pre>
```

Description

plotPowerFit

This function will plot sampling distributions of fit indices that visualize power in rejecting the misspecified models

ing datasets underlying misspecified models

Plot sampling distributions of fit indices that visualize power of reject-

Usage

```
plotPowerFit(altObject, nullObject = NULL, cutoff = NULL, usedFit = NULL,
alpha = 0.05, contN = TRUE, contMCAR = TRUE, contMAR = TRUE,
useContour = TRUE, logistic = TRUE)
```

Arguments

altObject	The result object (SimResult) saves the simulation result of fitting the hypothesized model when the hypothesized model is FALSE.
nullObject	The result object (SimResult) saves the simulation result of fitting the hypothesized model when the hypothesized model is TRUE. This argument may be not specified if the cutoff is specified.
cutoff	A vector of priori cutoffs for fit indices.
usedFit	Vector of names of fit indices that researchers wish to plot.
alpha	A priori alpha level
contN	Include the varying sample size in the power plot if available
contMCAR	Include the varying MCAR (missing completely at random percentage) in the power plot if available

76 plotPowerFit

contMAR Include the varying MAR (missing at random percentage) in the power plot if

available

useContour If there are two of sample size, percent completely at random, and percent miss-

ing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective

plot is used.

logistic If logistic is TRUE and the varying parameter exists (e.g., sample size or per-

cent missing), the plot based on logistic regression predicting the significance by the varying parameters is preferred. If FALSE, the overlaying scatterplot with

a line of cutoff is plotted.

Value

NONE. Only plot the fit indices distributions.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- SimResult for simResult that used in this function.
- getCutoff to find values of cutoffs based on null hypothesis sampling distributions only
- getPowerFit to find power of rejecting the hypothesized model when the hypothesized model is FALSE.

Examples

```
## Not run:
loading.null <- matrix(0, 6, 1)</pre>
loading.null[1:6, 1] <- NA</pre>
LX.NULL <- bind(loading.null, 0.7)
RPH.NULL <- binds(diag(1))</pre>
RTD <- binds(diag(6))
CFA.Model.NULL <- model(LY = LX.NULL, RPS = RPH.NULL, RTE = RTD, modelType="CFA")
# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output.NULL <- sim(50, n=50, model=CFA.Model.NULL, generate=CFA.Model.NULL)
loading.alt <- matrix(0, 6, 2)</pre>
loading.alt[1:3, 1] \leftarrow NA
loading.alt[4:6, 2] <- NA</pre>
LX.ALT <- bind(loading.alt, 0.7)
latent.cor.alt <- matrix(NA, 2, 2)</pre>
diag(latent.cor.alt) <- 1</pre>
RPH.ALT <- binds(latent.cor.alt, 0.5)</pre>
CFA.Model.ALT <- model(LY = LX.ALT, RPS = RPH.ALT, RTE = RTD, modelType="CFA")</pre>
Output.ALT <- sim(50, n=50, model=CFA.Model.NULL, generate=CFA.Model.ALT)
datNull <- generate(CFA.Model.NULL, n=50, params=TRUE)</pre>
datAlt <- generate(CFA.Model.ALT, n=50, params=TRUE)</pre>
outNull <- analyze(CFA.Model.NULL, datNull)</pre>
outAlt <- analyze(CFA.Model.NULL, datAlt)</pre>
summaryFit(Output.NULL)
```

plotPowerFitDf 77

```
plotPowerFit(Output.ALT, nullObject=Output.NULL, alpha=0.05, usedFit=c("RMSEA", "CFI", "TLI", "SRMR"))
Rule.of.thumb <- c(RMSEA=0.05, CFI=0.95, TLI=0.95, SRMR=0.06)
plotPowerFit(Output.ALT, cutoff=Rule.of.thumb, alpha=0.05, usedFit=c("RMSEA", "CFI", "TLI", "SRMR"))

Output.NULL2 <- sim(NULL, n=seq(50, 250, 25), model=CFA.Model.NULL, generate=CFA.Model.NULL)
Output.ALT2 <- sim(NULL, n=seq(50, 250, 25), model=CFA.Model.NULL, generate=CFA.Model.ALT)

plotPowerFit(Output.ALT2, nullObject=Output.NULL2, alpha=0.05, usedFit=c("RMSEA", "CFI", "TLI", "SRMR"))
plotPowerFit(Output.ALT2, cutoff=Rule.of.thumb, alpha=0.05, usedFit=c("RMSEA", "CFI", "TLI", "SRMR"))

## End(Not run)

Plot sampling distributions of fit indices that visualize power of rejecting datasets underlying misspecified models
```

Description

This function will plot sampling distributions of fit indices that visualize power in rejecting the misspecified models. This function is similar to the plotPowerFit function but the input distributions are data.frame.

Usage

plotPowerFitDf(altObject, nullObject = NULL, cutoff = NULL, usedFit = NULL, alpha = 0.05, x = NULL,

Arguments

altObject	The result object (data.frame) saves the simulation result of fitting the hypothesized model when the hypothesized model is FALSE.
nullObject	The result object (data.frame) saves the simulation result of fitting the hypothesized model when the hypothesized model is TRUE. This argument may be not specified if the cutoff is specified.
cutoff	A vector of priori cutoffs for fit indices.
usedFit	Vector of names of fit indices that researchers wish to plot.
alpha	A priori alpha level
X	The data.frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the object.
xval	The values of predictor that researchers would like to find the fit indices cutoffs from.
useContour	If there are two of sample size, percent completely at random, and percent missing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective plot is used.
logistic	If logistic is TRUE and the varying parameter exists (e.g., sample size or percent missing), the plot based on logistic regression predicting the significance by the varying parameters is preferred. If FALSE, the overlaying scatterplot with a line of cutoff is plotted.

78 plotPowerFitNested

Value

NONE. Only plot the fit indices distributions.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• plotPowerFit

Examples

No example

plotPowerFitNested

Plot power of rejecting a nested model in a nested model comparison by each fit index

Description

This function will plot sampling distributions of the differences in fit indices between parent and nested models. Two sampling distributions will be compared: nested model is FALSE (alternative model) and nested model is TRUE (null model).

Usage

```
plotPowerFitNested(altNested, altParent, nullNested = NULL,
nullParent = NULL, cutoff = NULL, usedFit = NULL, alpha = 0.05,
contN = TRUE, contMCAR = TRUE, contMAR = TRUE, useContour = TRUE,
logistic = TRUE)
```

Arguments

altNested	SimResult that saves the simulation result of the nested model when the nested model is FALSE.
altParent	SimResult that saves the simulation result of the parent model when the nested model is FALSE.
nullNested	SimResult that saves the simulation result of the nested model when the nested model is TRUE. This argument may not be specified if the cutoff is specified.
nullParent	SimResult that saves the simulation result of the parent model when the nested model is TRUE. This argument may not be specified if the cutoff is specified.
cutoff	A vector of priori cutoffs for the differences in fit indices.
usedFit	Vector of names of fit indices that researchers wish to plot.
alpha	A priori alpha level
contN	Include the varying sample size in the power plot if available
contMCAR	Include the varying MCAR (missing completely at random percentage) in the power plot if available

plotPowerFitNested 79

contMAR Include the varying MAR (missing at random percentage) in the power plot if

available

useContour If there are two of sample size, percent completely at random, and percent miss-

ing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective

plot is used.

logistic If logistic is TRUE and the varying parameter exists (e.g., sample size or per-

cent missing), the plot based on logistic regression predicting the significance by the varying parameters is preferred. If FALSE, the overlaying scatterplot with

a line of cutoff is plotted.

Value

NONE. Only plot the fit indices distributions.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- SimResult for simResult that used in this function.
- getCutoffNested to find the cutoffs of the differences in fit indices
- plotCutoffNested to visualize the cutoffs of the differences in fit indices
- getPowerFitNested to find the power in rejecting the nested model by the difference in fit indices cutoffs

Examples

```
## Not run:
loading.null <- matrix(0, 6, 1)</pre>
loading.null[1:6, 1] <- NA</pre>
LX.NULL <- bind(loading.null, 0.7)
RPH.NULL <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model.NULL <- model(LY = LX.NULL, RPS = RPH.NULL, RTE = RTD, modelType="CFA")
loading.alt <- matrix(0, 6, 2)</pre>
loading.alt[1:3, 1] \leftarrow NA
loading.alt[4:6, 2] \leftarrow NA
LX.ALT <- bind(loading.alt, 0.7)
latent.cor.alt <- matrix(NA, 2, 2)</pre>
diag(latent.cor.alt) <- 1</pre>
RPH.ALT <- binds(latent.cor.alt, 0.7)</pre>
CFA.Model.ALT <- model(LY = LX.ALT, RPS = RPH.ALT, RTE = RTD, modelType="CFA")
Output.NULL.NULL <- sim(10, n=500, model=CFA.Model.NULL, generate=CFA.Model.NULL)
Output.ALT.NULL <- sim(10, n=500, model=CFA.Model.NULL, generate=CFA.Model.ALT)
Output.NULL.ALT <- sim(10, n=500, model=CFA.Model.ALT, generate=CFA.Model.NULL)
Output.ALT.ALT <- sim(10, n=500, model=CFA.Model.ALT, generate=CFA.Model.ALT)
plotPowerFitNested(Output.ALT.NULL, Output.ALT.ALT, nullNested=Output.NULL.NULL, nullParent=Output.NULL.AL
```

plotPowerFitNested(Output.ALT.NULL, Output.ALT.ALT, nullNested=Output.NULL.NULL, nullParent=Output.NULL.AL

```
Output.NULL.NULL2 <- sim(NULL, n=seq(50, 500, 25), model=CFA.Model.NULL, generate=CFA.Model.NULL)
Output.ALT.NULL2 <- sim(NULL, n=seq(50, 500, 25), model=CFA.Model.NULL, generate=CFA.Model.ALT)
Output.NULL.ALT2 <- sim(NULL, n=seq(50, 500, 25), model=CFA.Model.ALT, generate=CFA.Model.NULL)
Output.ALT.ALT2 <- sim(NULL, n=seq(50, 500, 25), model=CFA.Model.ALT, generate=CFA.Model.ALT)

plotPowerFitNested(Output.ALT.NULL2, Output.ALT.ALT2, nullNested=Output.NULL.NULL2, nullParent=Output.NULL
plotPowerFitNested(Output.ALT.NULL2, Output.ALT.ALT2, nullNested=Output.NULL.NULL2, nullParent=Output.NULL
plotPowerFitNested(Output.ALT.NULL2, Output.ALT.ALT2, cutoff=c(CFI=-0.1), logistic=FALSE)

## End(Not run)
```

 ${\it plotPowerFitNonNested} \quad {\it Plot power of rejecting \ a \ non-nested \ model \ based \ on \ a \ difference \ in \ fit \ index}$

Description

Plot the proportion of the difference in fit indices from one model that does not in the range of sampling distribution from another model (reject that the dataset comes from the second model) or indicates worse fit than a specified cutoff. This plot can show the proportion in the second model that does not in the range of sampling distribution from the first model too.

Usage

```
plotPowerFitNonNested(dat2Mod1, dat2Mod2, dat1Mod1=NULL, dat1Mod2=NULL,
cutoff = NULL, usedFit = NULL, alpha = 0.05, contN = TRUE, contMCAR = TRUE,
contMAR = TRUE, useContour = TRUE, logistic = TRUE, onetailed = FALSE)
```

Arguments

dat2Mod1	SimResult that saves the simulation of analyzing Model 1 by datasets created from Model 2
dat2Mod2	${\tt SimResult}$ that saves the simulation of analyzing Model 2 by datasets created from Model 2
dat1Mod1	${\tt SimResult}$ that saves the simulation of analyzing Model 1 by datasets created from Model 1
dat1Mod2	${\tt SimResult}$ that saves the simulation of analyzing Model 2 by datasets created from Model 1
cutoff	A vector of priori cutoffs for the differences in fit indices.
usedFit	Vector of names of fit indices that researchers wish to plot.
alpha	A priori alpha level
contN	Include the varying sample size in the power plot if available
contMCAR	Include the varying MCAR (missing completely at random percentage) in the power plot if available
contMAR	Include the varying MAR (missing at random percentage) in the power plot if available

useContour	If there	e are two of	sample size	e, percen	t completely	y at randor	n, and percent miss-

ing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective

plot is used.

logistic If logistic is TRUE and the varying parameter exists (e.g., sample size or per-

cent missing), the plot based on logistic regression predicting the significance by the varying parameters is preferred. If FALSE, the overlaying scatterplot with

a line of cutoff is plotted.

onetailed If TRUE, the function will use the cutoff from one-tail test. If FALSE, the function

will use the cutoff from two-tailed test.

Value

NONE. Only plot the fit indices distributions.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

- SimResult for simResult that used in this function.
- getCutoffNonNested to find the cutoffs of the differences in fit indices for non-nested model comparison
- plotCutoffNonNested to visualize the cutoffs of the differences in fit indices for non-nested model comparison
- getPowerFitNonNested to find the power in rejecting the non-nested model by the difference in fit indices cutoffs

Examples

```
## Not run:
loading.A <- matrix(0, 8, 2)</pre>
loading.A[1:3, 1] <- NA
loading.A[4:8, 2] <- NA
LX.A <- bind(loading.A, 0.7)
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, "runif(1, 0.7, 0.9)")
RTD <- binds(diag(8))</pre>
CFA.Model.A <- model(LY = LX.A, RPS = RPH, RTE = RTD, modelType="CFA")
loading.B <- matrix(0, 8, 2)</pre>
loading.B[1:4, 1] <- NA
loading.B[5:8, 2] <- NA
LX.B <- bind(loading.B, 0.7)
CFA.Model.B <- model(LY = LX.B, RPS = RPH, RTE = RTD, modelType="CFA")
# The actual number of replications should be greater than 10.
Output.A.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.A)</pre>
Output.A.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.A)
Output.B.A <- sim(10, n=500, model=CFA.Model.A, generate=CFA.Model.B)
Output.B.B <- sim(10, n=500, model=CFA.Model.B, generate=CFA.Model.B)
```

82 plotPowerSig

```
plotPowerFitNonNested(Output.B.A, Output.B.B, dat1Mod1=Output.A.A, dat1Mod2=Output.A.B)
plotPowerFitNonNested(Output.B.A, Output.B.B, cutoff=c(AIC=0, BIC=0))
## End(Not run)
```

plotPowerSig

Plot multiple logistic curves given a significance result matrix

Description

This function will plot the significance results given the value of predictors.

Usage

```
plotPowerSig(sig, x = NULL, xval=NULL, mainName = NULL, useContour = TRUE)
```

Arguments

sig	The data.frame of a significance result, which contains only TRUE for significance and FALSE for not significance.
X	The data.frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the object.
xval	The values of predictor that researchers would like to find the fit indices cutoffs from.
mainName	A vector of the titles of the graphs
useContour	If there are two of sample size, percent completely at random, and percent missing at random are varying, the plotCutoff function will provide 3D graph. Contour graph is a default. However, if this is specified as FALSE, perspective

Value

NONE. Only plot the fit indices distributions.

plot is used.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• plotPower

Examples

No example

plotQtile 83

plotQtile	Build a scatterplot with overlaying line of quantiles of predicted values

Description

Build a scatterplot with overlaying line of quantiles of predicted values

Usage

```
plotQtile(x, y, df=0, qtile=NULL, ...)
```

Arguments

Х	The values of the independent variable (e.g., a vector of sample size)
У	The values of the dependent variable
df	The degree of freedom in spline method
qtile	The quantile values used to plot a graph
	Other arguments in the plot command

Value

None. This function will plot only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

```
# No example
```

plotScatter	Plot overlaying scatter plots visualizing the power of rejecting mis-
	specified models

Description

This function will find the fit indices cutoff values if not specified and then plot the fit indices value against the value of predictors. The plot will include the fit indices value of the alternative models, the fit indices value of the null model (if specified), and the fit indices cutoffs.

Usage

```
plotScatter(altObject, nullObject=NULL, cutoff=NULL, usedFit = NULL, x, alpha=0.05, df=5)
```

84 popDiscrepancy

Arguments

altObject The result object (data.frame) saves the simulation result of fitting the hypoth-

esized model when the hypothesized model is FALSE.

nullObject The result object (data.frame) saves the simulation result of fitting the hypoth-

esized model when the hypothesized model is TRUE. This argument may be not

specified if the cutoff is specified.

cutoff A vector of priori cutoffs for fit indices.

usedFit Vector of names of fit indices that researchers wish to plot.

x The data. frame of the predictor values. The number of rows of the x argument

should be equal to the number of rows in the object.

alpha A priori alpha level

df The degree of freedom used in spline method in predicting the fit indices by the

predictors. If df is 0, the spline method will not be applied.

Value

NONE. Only plot the fit indices distributions.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• plotPowerFit

Examples

No example

popDiscrepancy	Find the discrepancy value between two means and covariance matri-
	Ces

Description

Find the discrepancy value between two means and covariance matrices

Usage

```
popDiscrepancy(paramM, paramCM, misspecM, misspecCM)
```

Arguments

paramM	The model-	implied mea	n from the rea	al parameters
--------	------------	-------------	----------------	---------------

paramCM The model-implied covariance matrix from the real parameters

misspecM The model-implied mean from the real and misspecified parameters

The model implied mean from the real and impospermed parameters

misspecCM The model-implied covariance matrix from the real and misspecified parameters

popMisfitMACS 85

Details

The discrepancy value (F_0 ; Browne & Cudeck, 1992) is calculated by

$$F_0 = tr\left(\tilde{\Sigma}\Sigma^{-1}\right) - \log\left|\tilde{\Sigma}\Sigma^{-1}\right| - p + \left(\tilde{\mu} - \mu\right)'\Sigma^{-1}\left(\tilde{\mu} - \mu\right).$$

where μ is the model-implied mean from the real parameters, Σ is the model-implied covariance matrix from the real parameters, $\tilde{\mu}$ is the model-implied mean from the real and misspecified parameters, $\tilde{\Sigma}$ is the model-implied covariance matrix from the real and misspecified parameter, p is the number of indicators.

Value

The discrepancy between two means and covariance matrices

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

References

Browne, M. W., & Cudeck, R. (1992). Alternative ways of assessing model fit. *Sociological Methods & Research*, 21, 230-258.

Examples

```
\begin{array}{l} \text{m1} <-\text{ rep}(0,\ 3)\\ \text{m2} <-\text{ c}(0.1,\ -0.1,\ 0.05)\\ \text{S1} <-\text{ matrix}(\text{c}(1,\ 0.6,\ 0.5,\ 0.6,\ 1,\ 0.4,\ 0.5,\ 0.4,\ 1),\ 3,\ 3)\\ \text{S2} <-\text{ matrix}(\text{c}(1,\ 0.55,\ 0.55,\ 0.55,\ 1,\ 0.55,\ 0.55,\ 0.55,\ 1),\ 3,\ 3)\\ \text{popDiscrepancy}(\text{m1},\ \text{S1},\ \text{m2},\ \text{S2}) \end{array}
```

popMisfitMACS

Find population misfit by sufficient statistics

Description

Find the value quantifying the amount of population misfit: F_0 , RMSEA, and SRMR.

Usage

```
popMisfitMACS(paramM, paramCM, misspecM, misspecCM, dfParam=NULL, fit.measures="all")
```

Arguments

paramM	The model-implied mean from the real parameters
paramCM	The model-implied covariance matrix from the real parameters
misspecM	The model-implied mean from the real and misspecified parameters
misspecCM	The model-implied covariance matrix from the real and misspecified parameters
dfParam	The degree of freedom of the real model
fit.measures	The names of indices used to calculate population misfit. There are three types of misfit: 1) discrepancy function ("f0"; see popDiscrepancy), 2) root mean squared error of approximation ("rmsea"; Equation 12 in Browne & Cudeck, 1992), and 3) standardized root mean squared residual ("srmr")

86 predProb

Details

The root mean squared error of approximation (RMSEA) is calculated by

$$RMSEA = \sqrt{\frac{F_0}{df}}$$

where F_0 is the discrepancy value between two means vectors and covariance matrices (see popDiscrepancy) and df is the degree of freedom in the real model.

The standardized root mean squared residual can be calculated by

$$SRMR = \sqrt{\frac{2\sum_{i}\sum_{j\leq i}\left(\frac{s_{ij}}{\sqrt{s_{ii}}\sqrt{s_{jj}}} - \frac{\hat{\sigma}_{ij}}{\sqrt{\hat{\sigma}_{ii}}\sqrt{\hat{\sigma}_{jj}}}\right)}{p(p+1)}}$$

where s_{ij} is the observed covariance between indicators i and j, $\hat{\sigma}_{ij}$ is the model-implied covariance between indicators i and j, p is the number of indicators.

Value

The vector of the misfit indices

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

References

Browne, M. W., & Cudeck, R. (1992). Alternative ways of assessing model fit. *Sociological Methods & Research*, 21, 230-258.

Examples

```
\begin{array}{l} \text{m1} <-\text{ rep}(0,\ 3)\\ \text{m2} <-\text{ c}(0.1,\ -0.1,\ 0.05)\\ \text{S1} <-\text{ matrix}(\text{c}(1,\ 0.6,\ 0.5,\ 0.6,\ 1,\ 0.4,\ 0.5,\ 0.4,\ 1),\ 3,\ 3)\\ \text{S2} <-\text{ matrix}(\text{c}(1,\ 0.55,\ 0.55,\ 0.55,\ 1,\ 0.55,\ 0.55,\ 0.55,\ 1),\ 3,\ 3)\\ \text{popMisfitMACS}(\text{m1},\ \text{S1},\ \text{m2},\ \text{S2}) \end{array}
```

predProb

Function to get predicted probabilities from logistic regression

Description

Function to get predicted probabilities from logistic regression

Usage

```
predProb(newdat, glmObj)
```

printIfNotNull 87

Arguments

newdat A vector of values for all predictors, including the intercept

glmObj An object from a fitted glm run with a logit link

Value

Predictive probability of success given the values in the newdat argument.

Author(s)

Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>)

See Also

- continuousPower
- getPower

Examples

No example

printIfNotNull

Provide basic summary of each object if that object is not NULL.

Description

Provide basic summary of each object if that object is not NULL. This function is mainly used in the summary function from the linkS4class{SimSet} object.

Usage

```
printIfNotNull(object, name=NULL)
```

Arguments

object The target object to be printed, which can be linkS4class{SimMatrix}, linkS4class{SymMatrix}

or linkS4class{SimVector}.

name The name of the target object

Value

None. This function will print only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

```
# This function is not public
# AL <- simVector(rep(NA, 5), "0")
# printIfNotNull(AL, "Factor mean")</pre>
```

88 pValue

pValue	Find p-values (1 - percentile)	

Description

This function will provide *p* value from comparing number and vector or the analytic result to the observed data (in lavaan) and the simulation result (in SimResult).

Usage

```
pValue(target, dist, ...)
```

Arguments

A value, multiple values, or a model output object used to find p values. This argument could be a cutoff of a fit index.

dist

The comparison distribution, which can be a vector of numbers, a data frame, or a result object.

Other values that will be explained specifically for each class

Details

In comparing fit indices, the *p* value is the proportion of the number of replications that provide poorer fit (e.g., less CFI value or greater RMSEA value) than the analysis result from the observed data. If the target is a critical value (e.g., fit index cutoff) and the dist is the sampling distribution underlying the alternative hypothesis, this function can provide a statistical power.

Value

Mostly, this function provides a vector of p values based on the comparison. If the target is a model output object and dist is a result object, the p values of fit indices are provided, as well as two additional values: andRule and orRule. The andRule is based on the principle that the model is retained only when all fit indices provide good fit. The proportion is calculated from the number of replications that have all fit indices indicating a better model than the observed data. The proportion from the andRule is the most stringent rule in retaining a hypothesized model. The orRule is based on the principle that the model is retained only when at least one fit index provides good fit. The proportion is calculated from the number of replications that have at least one fit index indicating a better model than the observed data. The proportion from the orRule is the most lenient rule in retaining a hypothesized model.

Methods

signature(target="numeric", dist="vector") This method will find the *p* value (quantile rank) of the target value on the dist vector. The additional arguments are revDirec, x, xval, condCutoff, and df. The revDirec is a logical argument whether to reverse the direction of comparison. If TRUE, the proportion of the dist that is lower than target value is reported. If FALSE, the proportion of the dist that is higher than the target value is reported. The x is the data. frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the dist. The xval is the values of predictor that researchers would like to find the fit indices cutoffs from. The condCutoff is a logical argument. If TRUE, the cutoff is applicable only a given value of xval. If FALSE, the cutoff is applicable in any

pValue 89

values of predictor. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

signature(target="numeric", dist="data.frame") This method will find the p value of each columns in the dist based on the value specified in the target. The additional arguments are revDirec, x, xval, df, and asLogical. The revDirec is a logical vector whether to reverse the direction of comparison. If TRUE, the proportion of the dist that is lower than target value is reported. If FALSE, the proportion of the dist that is higher than the target value is reported. The x is the data.frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the dist. The xval is the values of predictor that researchers would like to find the fit indices cutoffs from. The df is the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied. The asLogical is to provide the result as the matrix of significance result (TRUE) or just the proportion of significance result (FALSE).

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimResult to run a simulation study

Examples

```
## Not run:
# Compare number with a vector
pValue(0.5, rnorm(1000, 0, 1))
# Compare numbers with a data frame
pValue(c(0.5, 0.2), data.frame(rnorm(1000, 0, 1), runif(1000, 0, 1)))
# Compare an analysis result with a result of simulation study
library(lavaan)
loading <- matrix(0, 9, 3)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loading[7:9, 3] <- NA
targetmodel <- estmodel(LY=loading, modelType="CFA", indLab=paste("x", 1:9, sep=""))</pre>
out <- analyze(targetmodel, HolzingerSwineford1939)</pre>
loading.trivial <- matrix("runif(1, -0.2, 0.2)", 9, 3)</pre>
loading.trivial[is.na(loading)] <- 0</pre>
mismodel <- model.lavaan(out, std=TRUE, LY=loading.trivial)</pre>
simout <- sim(20, n=nrow(HolzingerSwineford1939), mismodel)</pre>
pValue(out, simout)
## End(Not run)
```

90 pValueCondCutoff

pValueCondCutoff	Find a p value when the target is conditional (valid) on a specific value of a predictor	
------------------	--	--

Description

Find a p value when the target is conditional (valid) on a specific value of a predictor. That is, the target value is applicable only a given value of a predictor.

Usage

```
pValueCondCutoff(target, dist, revDirec = FALSE, x = NULL, xval = NULL, df = 0)
```

Arguments

target	A target value used to find p values.
dist	The comparison distribution, which can be a vector of numbers, a data frame, or a result object.
revDirec	A logical argument whether to reverse the direction of comparison. If TRUE, the proportion of the dist that is lower than target value is reported. If FALSE, the proportion of the dist that is higher than the target value is reported.
x	the data.frame of the predictor values. The number of rows of the x argument should be equal to the number of rows in the dist
xval	the values of predictor that researchers would like to find the fit indices cutoffs from.
df	the degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

Value

A vector of p values based on the comparison.

Author(s)

 $Sunthud\ Pornprasert manit\ (University\ of\ Kansas; < psunthud@ku.edu>)$

See Also

• pValue

Examples

No example

pValueNested 91

pValueNested	Find p-values (1 - percentile) for a nested model comparison

Description

This function will provide p value from comparing the differences in fit indices between nested models with the simulation results of both parent and nested models when the nested model is true.

Usage

```
pValueNested(outNested, outParent, simNested, simParent, usedFit = NULL, nVal = NULL, pmMCARval = NULL
```

Arguments

outNested	lavaan that saves the analysis result of the nested model from the target dataset
outParent	lavaan that saves the analysis result of the parent model from the target dataset
simNested	SimResult that saves the analysis results of nested model from multiple replications
simParent	SimResult that saves the analysis results of parent model from multiple replications
usedFit	Vector of names of fit indices that researchers wish to getCutoffs from. The default is to getCutoffs of all fit indices.
nVal	The sample size value that researchers wish to find the p value from.
pmMCARval	The percent missing completely at random value that researchers wish to find the p value from.
pmMARval	The percent missing at random value that researchers wish to find the the p value from.
df	The degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.

Details

In comparing fit indices, the *p* value is the proportion of the number of replications that provide less preference for nested model (e.g., larger negative difference in CFI values or larger positive difference in RMSEA values) than the analysis result from the observed data.

Value

This function provides a vector of *p* values based on the comparison of the difference in fit indices from the real data with the simulation result. The *p* values of fit indices are provided, as well as two additional values: andRule and orRule. The andRule is based on the principle that the model is retained only when all fit indices provide good fit. The proportion is calculated from the number of replications that have all fit indices indicating a better model than the observed data. The proportion from the andRule is the most stringent rule in retaining a hypothesized model. The orRule is based on the principle that the model is retained only when at least one fit index provides good fit. The proportion is calculated from the number of replications that have at least one fit index indicating a better model than the observed data. The proportion from the orRule is the most lenient rule in retaining a hypothesized model.

92 pValueNonNested

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimResult to run a simulation study

Examples

```
## Not run:
library(lavaan)
LY <- matrix(1, 4, 2)
LY[,2] <- 0:3
PS <- matrix(NA, 2, 2)
TY <- rep(0, 4)
AL \leftarrow rep(NA, 2)
TE <- diag(NA, 4)
nested <- estmodel(LY=LY, PS=PS, TY=TY, AL=AL, TE=TE, modelType="CFA", indLab=paste("t", 1:4, sep=""))</pre>
LY2 <- matrix(1, 4, 2)
LY2[,2] \leftarrow c(0, NA, NA, 3)
parent <- estmodel(LY=LY2, PS=PS, TY=TY, AL=AL, TE=TE, modelType="CFA", indLab=paste("t", 1:4, sep=""))</pre>
outNested <- analyze(nested, Demo.growth)</pre>
outParent <- analyze(parent, Demo.growth)</pre>
loadingMis <- matrix(0, 4, 2)</pre>
loadingMis[2:3, 2] <- "runif(1, -0.1, 0.1)"
datamodel <- model.lavaan(outNested, LY=loadingMis)</pre>
n <- nrow(Demo.growth)</pre>
simNestedNested <- sim(30, n=n, nested, generate=datamodel)</pre>
simNestedParent <- sim(30, n=n, parent, generate=datamodel)</pre>
pValueNested(outNested, outParent, simNestedNested, simNestedParent)
## End(Not run)
```

pValueNonNested

Find p-values (1 - percentile) for a non-nested model comparison

Description

This function will provide p value from comparing the results of fitting real data into two models against the simulation from fitting the simulated data from both models into both models. The p values from both sampling distribution under the datasets from the first and the second models are reported.

Usage

```
\label{eq:pvalueNonNested} $$ pValueNonNested(outMod1, outMod2, dat1Mod1, dat1Mod2, dat2Mod1, dat2Mod2, usedFit = NULL, nVal = NULL, pmMCARval = NULL, pmMARval = NULL, df = 0, onetailed=FALSE)
```

pValueNonNested 93

Arguments

outMod1	lavaan that saves the analysis result of the first model from the target dataset
outMod2	lavaan that saves the analysis result of the second model from the target dataset
dat1Mod1	${\tt SimResult}$ that saves the simulation of analyzing Model 1 by datasets created from Model 1
dat1Mod2	${\tt SimResult}$ that saves the simulation of analyzing Model 2 by datasets created from Model 1
dat2Mod1	${\tt SimResult}$ that saves the simulation of analyzing Model 1 by datasets created from Model 2
dat2Mod2	${\tt SimResult}$ that saves the simulation of analyzing Model 2 by datasets created from Model 2
usedFit	Vector of names of fit indices that researchers wish to getCutoffs from. The default is to getCutoffs of all fit indices.
nVal	The sample size value that researchers wish to find the p value from.
pmMCARval	The percent missing completely at random value that researchers wish to find the p value from.
pmMARval	The percent missing at random value that researchers wish to find the the p value from.
df	The degree of freedom used in spline method in predicting the fit indices by the predictors. If df is 0, the spline method will not be applied.
onetailed	If TRUE, the function will convert the p value based on two-tailed test.

Details

In comparing fit indices, the p value is the proportion of the number of replications that provide less preference for either model 1 or model 2 than the analysis result from the observed data. In two-tailed test, the function will report the proportion of values under the sampling distribution that are more extreme that one obtained from real data. If the resulting p value is high (> .05) on one model and low (< .05) in the other model, the model with high p value is preferred. If the p values are both high or both low, the decision is undetermined.

Value

This function provides a vector of p values based on the comparison of the difference in fit indices from the real data with the simulation results. The p values of fit indices are provided, as well as two additional values: andRule and orRule. The andRule is based on the principle that the model is retained only when all fit indices provide good fit. The proportion is calculated from the number of replications that have all fit indices indicating a better model than the observed data. The proportion from the andRule is the most stringent rule in retaining a hypothesized model. The orRule is based on the principle that the model is retained only when at least one fit index provides good fit. The proportion is calculated from the number of replications that have at least one fit index indicating a better model than the observed data. The proportion from the orRule is the most lenient rule in retaining a hypothesized model.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

94 pValueVariedCutoff

See Also

• SimResult to run a simulation study

Examples

```
## Not run:
 library(lavaan)
 loading <- matrix(0, 11, 3)</pre>
 loading[1:3, 1] <- NA
 loading[4:7, 2] <- NA
 loading[8:11, 3] <- NA
 path.A <- matrix(0, 3, 3)</pre>
 path.A[2:3, 1] <- NA
 path.A[3, 2] <- NA
 model.A <- estmodel(LY=loading, BE=path.A, modelType="SEM", indLab=c(paste("x", 1:3, sep=""), paste("y", 1</pre>
 out.A <- analyze(model.A, PoliticalDemocracy)</pre>
 path.B <- matrix(0, 3, 3)</pre>
 path.B[1:2, 3] <- NA
 path.B[1, 2] <- NA
 model.B <- estmodel(LY=loading, BE=path.B, modelType="SEM", indLab=c(paste("x", 1:3, sep=""), paste("y", 1</pre>
 out.B <- analyze(model.B, PoliticalDemocracy)</pre>
 loading.mis <- matrix("runif(1, -0.2, 0.2)", 11, 3)</pre>
 loading.mis[is.na(loading)] <- 0</pre>
 datamodel.A <- model.lavaan(out.A, std=TRUE, LY=loading.mis)</pre>
 datamodel.B <- model.lavaan(out.B, std=TRUE, LY=loading.mis)</pre>
 n <- nrow(PoliticalDemocracy)</pre>
 output.A.A <- sim(5, n=n, model.A, generate=datamodel.A)
 output.A.B <- sim(5, n=n, model.B, generate=datamodel.A)
 output.B.A <- sim(5, n=n, model.A, generate=datamodel.B)
 output.B.B <- sim(5, n=n, model.B, generate=datamodel.B)</pre>
 # The output may contain some warnings here. When the number of replications increases (e.g., 1000), the w.
 pValueNonNested(out.A, out.B, output.A.A, output.A.B, output.B.A, output.B.B)
 ## End(Not run)
pValueVariedCutoff
                           Find a p value when the cutoff is specified as a vector given the values
```

Description

Find a p value when the cutoff is specified as a vector given the values of predictors.

of predictors

Usage

```
pValueVariedCutoff(cutoff, obtainedValue, revDirec = FALSE, x = NULL, xval = NULL)
```

revText 95

Arguments

cutoff A vector of values used to find p values. Each value in the vector should be the

target value conditional (applicable) to each value of the predictors (x) respec-

tively.

obtainedValue The comparison distribution, which can be a vector of numbers, a data frame, or

a result object.

revDirec A logical argument whether to reverse the direction of comparison. If TRUE, the

proportion of the dist that is lower than target value is reported. If FALSE, the

proportion of the dist that is higher than the target value is reported.

x the data.frame of the predictor values. The number of rows of the x argument

should be equal to the number of rows in the dist

xval the values of predictor that researchers would like to find the fit indices cutoffs

from.

Value

A vector of *p* values based on the comparison.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• pValue

Examples

No example

revText

Reverse the proportion value by subtracting it from 1

Description

Reverse the proportion value by subtracting it from 1. This function can reverse a value reported in text, such as from "> .98" to "< .02"

Usage

revText(val)

Arguments

val The value to be reversed

Value

The reversed value or text

96 setPopulation

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
# This is a private function.
# revText(.96)
# revText("> .60")
```

setPopulation

Set the data generation population model underlying an object

Description

This function will set the data generation population model to be an appropriate one. If the appropriate data generation model is put (the same model as the analysis model), the additional features can be seen when we run a summary function on the target object, such as bias in parameter estimates or percentage coverage.

Usage

```
setPopulation(target, population)
```

Arguments

target The result object that you wish to set the data generation population model

(linkS4class{SimResult}).

population The population parameters specified in the linkS4class{SimSem} object

Value

The target object that is changed the parameter.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimResult for result object

Examples

```
# See each class for an example.
## Not run:
loading <- matrix(0, 7, 3)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loading[1:7, 3] <- NA
loadingVal <- matrix(0, 7, 3)</pre>
```

setPopulation 97

```
loadingVal[1:3, 1] <- "runif(1, 0.5, 0.7)"</pre>
loadingVal[4:6, 2] <- "runif(1, 0.5, 0.7)"
loadingVal[1:6, 3] <- "runif(1, 0.3, 0.5)"</pre>
loadingVal[7, 3] <- 1</pre>
loading.mis <- matrix("runif(1, -0.2, 0.2)", 7, 3)</pre>
loading.mis[is.na(loading)] <- 0</pre>
loading.mis[,3] <- 0</pre>
loading.mis[7,] <- 0</pre>
LY <- bind(loading, loadingVal, misspec=loading.mis)
RPS <- binds(diag(3))</pre>
path <- matrix(0, 3, 3)</pre>
path[2, 1] <- NA
BE <- bind(path, "runif(1, 0.3, 0.5)")
RTE <- binds(diag(7))</pre>
VY <- bind(c(rep(NA, 6), 0), c(rep(1, 6), ""))</pre>
datamodel <- model(LY=LY, RPS=RPS, BE=BE, RTE=RTE, VY=VY, modelType="SEM")</pre>
loading <- matrix(0, 7, 3)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loading[7, 3] <- NA</pre>
path <- matrix(0, 3, 3)</pre>
path[2, 1] <- NA
path[1, 3] <- NA
path[2, 3] <- NA
errorCov <- diag(NA, 7)</pre>
errorCov[7, 7] <- 0
facCov <- diag(3)</pre>
analysis <- estmodel(LY=loading, BE=path, TE=errorCov, PS=facCov, modelType="SEM", indLab=paste("y", 1:7, september 1:7, september 2:10 | 1:7, september 2
Output <- sim(100, n=200, analysis, generate=datamodel)
loadingVal <- matrix(0, 7, 3)</pre>
loadingVal[1:3, 1] \leftarrow 0.6
loadingVal[4:6, 2] <- 0.6</pre>
loadingVal[7, 3] <- 1</pre>
LY <- bind(loading, loadingVal)
pathVal <- matrix(0, 3, 3)</pre>
pathVal[2, 1] <- 0.4</pre>
pathVal[1, 3] <- 0.4
pathVal[2, 3] <- 0.4</pre>
BE <- bind(path, pathVal)</pre>
PS <- binds(facCov)
errorCovVal <- diag(0.64, 7)</pre>
errorCovVal[7, 7] <- 0
TE <- binds(errorCov, errorCovVal)</pre>
population <- model(LY=LY, PS=PS, BE=BE, TE=TE, modelType="SEM")</pre>
Output <- setPopulation(Output, population)</pre>
summary(Output)
## End(Not run)
```

98 sim

sim

Run a monte carlo simulation with a structural equation model.

Description

TBA

Usage

```
sim(nRep, model, n, generate = NULL, rawData = NULL, miss = NULL, datafun=NULL, outfun=NULL,
pmMCAR = NULL, pmMAR = NULL, facDist = NULL, indDist = NULL, errorDist = NULL, sequential = FALSE
modelBoot = FALSE, realData = NULL, maxDraw = 50, misfitType = "f0",
misfitBounds = NULL, averageNumMisspec = NULL, optMisfit=NULL, optDraws = 50,
aux = NULL, seed = 123321, silent = FALSE, multicore = FALSE, cluster = FALSE, numProc = NULL,
paramOnly = FALSE, dataOnly=FALSE, ...)
```

Arguments

nRep	Number of replications. Users can specify as NULL and specify n, pmMCAR, and pmMAR $$
model	SimSem object created by model. Will be used to generate data and analyze it.
n	Sample size. This argument is not necessary except the user wish to vary sample size across replications. The sample size here is a vector of sample size in integers. For the random distribution object, if the resulting value has decimal, the value will be rounded.
generate	SimSem object created by model. If included, this will be used to generate data instead of
rawData	If included, this data is used for simulations instead of being generated from the SimSem template. Should be a list of length nRep.
miss	Missing data handling template, created by the function miss.
datafun	Function to be applied to generated data set at each replication.
outfun	Function to be applied to the lavaan output at each replication. The output of this function in each replication will be saved in the simulation output (SimResult). The extra outputs can be obtained by the getExtraOutput function.
pmMCAR	The percent completely missing at random. This argument is not necessary except the user wish to vary percent missing completely at random across replications. The pmMCAR here is a vector of percent missing, which the values can be in between 0 and 1 only. The specification of objMissing is not needed (but is needed if users wish to specify complex missing value data generation or wish to use multiple imputation).
pmMAR	The percent missing at random. This argument is not necessary except the user wish to vary percent missing at random across replications. The pmMAR here is a vector of percent missing, which the values can be in between 0 and 1 only. The specification of objMissing is not needed (but is needed if users wish to specify complex missing value data generation or wish to use multiple imputation).
facDist	A SimDataDist object or list of objects for the distribution of factors. If one ob-

ject is passed, all factors will have the same distribution. Use when sequential

is TRUE.

sim 99

indDist A SimDataDist object or list of objects for a distribution of indicators. If one object is passed, each indicator will have the same distribution. Use when sequential is FALSE. errorDist An object or list of objects of type SimDataDist indicating the distribution of errors. If a single SimDataDist is specified, each error will be genrated with that distribution. sequential If TRUE, use a sequential method to create data such that the data from factor are generated first and apply to a set of equations to obtain the data of indicators. If FALSE, create data directly from model-implied mean and covariance of indicators. modelBoot When specified, a model-based bootstrap is used for data generation. See draw for further information. This argument requires real data to be passed to realData. A data frame containing real data. The data generated will follow the distriburealData tion of this data set. Integer specifying the maximum number of attempts to draw a valid set of pamaxDraw rameters (no negative error variance, standardized coefficients over 1). Character vector indicating the fit measure used to assess the misfit of a set of misfitType parameters. Can be "f0", "rmsea", "srmr", or "all". misfitBounds Vector that contains upper and lower bounds of the misfit measure. Sets of parameters drawn that are not within these bounds are rejected. averageNumMisspec TRUE or FALSE. ?? optMisfit Character vector of either "min" or "max" indicating either maximum or minimum optimized misfit. If not null, the set of parameters out of the number of draws in "optDraws" that has either the maximum or minimum misfit of the given misfit type will be returned. optDraws Number of parameter sets to draw if optMisfit is not null. The set of parameters with the maximum or minimum misfit will be returned. The names of auxiliary variables saved in a vector aux Random number seed. Reproducibility across multiple cores or clusters is enseed sured using R'Lecuyer package. silent If TRUE, suppress warnings. Use multiple processors within a computer. Specify as TRUE to use it. multicore cluster Not applicable now. Use for specify nodes in hpc in order to be parallelizable.

Value

TBA

. . .

numProc

paramOnly

dataOnly

Author(s)

will find the maximum number of processors.

Additional arguments to be passed to lavaan.

Number of processors for using multiple processors. If it is NULL, the package

If TRUE, only the parameters from each replication will be returned.

If TRUE, only the data generated from each replication will be returned.

100 SimDataDist-class

Examples

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
LY <- bind(loading, 0.7)

latent.cor <- matrix(NA, 2, 2)
diag(latent.cor) <- 1
RPS <- binds(latent.cor, 0.5)

RTE <- binds(diag(6))

VY <- bind(rep(NA,6),2)

CFA.Model <- model(LY = LY, RPS = RPS, RTE = RTE, modelType = "CFA")

Output <- sim(20, CFA.Model,n=200)
summary(Output)</pre>
```

SimDataDist-class

Class "SimDataDist"

Description

This class will provide the distribution of a dataset.

Objects from the Class

Objects can be created by bindDist function. It can also be called from the form new("SimDataDist", ...).

Slots

p: Number of variables

margins: A character vector specifying all the marginal distributions

paramMargins: A list whose each component is a list of named components, giving the parameter values of the marginal distributions.

keepScale: Transform back to retain the mean and standard deviation of a variable equal to the model implied mean and standard deviation (with sampling error)

reverse: To mirror each variable or not. If TRUE, reverse the distribution of a variable (e.g., from positive skewed to negative skewed).

Methods

- summaryTo summarize the object
- plotDistTo plot a density distribution (for one variable) or a contour plot (for two variables). If the object has more than two variables, the var argument can be used to select the index of plotting variables. For two variables, the default is to have correlation of 0. To change a correlation, the r argument can be used. The xlim and ylim can be specified to set the ranges of variables.
- extractExtract elements from an object. The next argument is the position of the object to be extracted.

SimMatrix-class 101

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• bindDist The constructor of this class.

Examples

```
# Need to be fixed
showClass("SimDataDist")

d1 <- list(df=2)
d2 <- list(df=3)
d3 <- list(df=4)
d4 <- list(df=5)
d5 <- list(df=3)
d6 <- list(df=3)
d6 <- list(df=4)
d7 <- list(df=5)
d8 <- list(df=6)</pre>
dist <- bindDist(c(rep("t", 4), rep("chisq", 8)), d1, d2, d3, d4, d5, d6, d7, d8, d5, d6, d7, d8)
```

SimMatrix-class

Matrix object: Random parameters matrix

Description

This object can be used to represent a matrix in SEM model. It contains free parameters, fixed values, and starting values. This object can be represented factor loading matrix or regreesion coefficient matrix.

Objects from the Class

This object is created by bind function. Objects can be also created by calls of the form new("SimMatrix", ...).

Slots

```
free: TBA
popParam: TBA
misspec: TBA
symmetric: TBA
```

Methods

summaryShort Provides a short summary of all information in the object summary Provides a thorough description of all information in the object

102 SimMissing-class

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimVector for random parameter vector.

Examples

```
showClass("SimMatrix")

loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loadingValues <- matrix(0, 6, 2)
loadingValues[1:3, 1] <- 0.7
loadingValues[4:6, 2] <- 0.7
LX <- bind(loading, loadingValues)
summary(LX)
# run(LX)

LY <- bind(loading, "rnorm(1, 0.6, 0.05)")
summary(LY)
# run(LY)</pre>
```

SimMissing-class

Class "SimMissing"

Description

Missing information imposing on the complete dataset

Objects from the Class

Objects can be created by simMissing function. It can also be called from the form new("SimMissing", ...).

Slots

cov: Column indices of any normally distributed covariates used in the data set.

pmMCAR: Decimal percent of missingness to introduce completely at random on all variables.

pmMAR: Decimal percent of missingness to introduce using the listed covariates as predictors.

nforms: The number of forms for planned missing data designs, not including the shared form.

itemGroups: List of lists of item groupings for planned missing data forms. Without this, items will be divided into groups sequentially (e.g. 1-3,4-6,7-9,10-12)

twoMethod: Vector of (percent missing, column index). Will put a given percent missing on that column in the matrix to simulate a two method planned missing data research design.

prAttr: Probability (or vector of probabilities) of an entire case being removed due to attrition at a given time point. See imposeMissing for further details.

package: The package used in multiple imputation. If "default", the full-information maximum likelihood is used.

SimResult-class 103

timePoints: Number of timepoints items were measured over. For longitudinal data, planned missing designs will be implemented within each timepoint.

ignoreCols: The columns not imposed any missing values for any missing data patterns

threshold: The threshold of covariates that divide between the area to impose missing and the area not to impose missing. The default threshold is the mean of the covariate.

covAsAux: If TRUE, the covariate listed in the object will be used as auxiliary variables when putting in the model object. If FALSE, the covariate will be included in the analysis.

logical: A matrix of logical values (TRUE/FALSE). If a value in the dataset is corresponding to the TRUE in the logical matrix, the value will be missing.

args: A list of additional options to be passed to the multiple imputation function in each package.

Methods

• summary To summarize the object

Author(s)

Patrick Miller(University of Kansas; <patr1ckm@ku.edu>) Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>) Kyle Lang (University of Kansas; <kylelang@ku.edu>) Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• imposeMissing for directly imposing missingness into a dataset.

Examples

No Example

SimResult-class

Class "SimResult"

Description

This class will save data analysis results from multiple replications and ready to find some useful statistics, such as fit indices cutoffs or power.

Objects from the Class

Objects can be created by sim. It can also be called from the form new("SimResult", ...).

Slots

modelType: Analysis model type (CFA, Path, or SEM)

nRep: Number of replications have been created and run simulated data.

coef: Parameter estimates from each replication

se: Standard errors of parameter estimates from each replication

fit: Fit Indices values from each replication

converged: Number of convergence replications

104 SimResult-class

seed: Seed number.

paramValue: Population model underlying each simulated dataset.

misspecValue: Misspecified-parameter values that are imposed on the population model in each replication.

popFit: The amount of population misfit. See details at popMisfitMACS

FMI1: Fraction Missing Method 1.

FMI2: Fraction Missing Method 2.

stdCoef: Standardized coefficients from each replication

n: Sample size of the analyzed data.

pmMCAR: Percent missing completely at random.

pmMAR: Percent missing at random.

extraOut: Extra outputs obtained from running the function specified in outfun argument.

timing: Time elapsed in each phase of the simulation.

Methods

- getCutoff to getCutoff of fit indices based on a priori alpha level.
- getPowerFit to getPowerFit of rejection when the simResult is the alternative hypothesis and users specify cutoffs of the fit indices.
- plotCutoff to plot null hypothesis sampling distributions of fit indices with an option to draw fit indices cutoffs by specifying a priori alpha level.
- plotPowerFit to plot alternative hypothesis (and null hypothesis) with a priori cutoffs or alpha level.
- summary to summarize the result output
- summaryParam to summarize all parameter estimates
- anova find the averages of model fit statistics and indices for nested models, as well as the differences of model fit indices among models. This function requires at least two SimResult objects. See anova for further details.
- summaryPopulation to summarize the data generation population underlying the simulation study.
- getPopulation to extract the data generation population underlying the simulation study. This method will return a data frame of the population underlying each replication.
- setPopulation to put the appropriate data generation model into the result object. If the appropriate data generation model is put (the same model as the analysis model), the bias in parameter estimates and standard errors will be able to be calculated by the summary function. The first argument is the result object. The second argument can be either data.frame of the population or SimSet of the population. See the 'modeling with covariate' in the manual for an example.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• sim for the constructor of this class

SimSem-class 105

Examples

```
showClass("SimResult")
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")

# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output <- sim(5, n=500, CFA.Model)
summary(Output)
getCutoff(Output, 0.05)
summaryParam(Output)
summaryPopulation(Output)</pre>
```

SimSem-class

Class "SimSem"

Description

TBA

Objects from the Class

TBA

Slots

```
pt: TBA
dgen: TBA
modelType: TBA
groupLab: TBA
```

Methods

summary Get the summary of model specification

Author(s)

See Also

• Create an object this class by CFA, Path Analysis, or SEM model by model.

106 SimVector-class

Examples

```
showClass("SimSem")
loading <- matrix(0, 6, 2)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loadingValues <- matrix(0, 6, 2)</pre>
loadingValues[1:3, 1] <- 0.7</pre>
loadingValues[4:6, 2] <- 0.7</pre>
LX <- bind(loading, loadingValues)
summary(LX)
latent.cor <- matrix(NA, 2, 2)</pre>
diag(latent.cor) <- 1</pre>
RPH <- binds(latent.cor, 0.5)</pre>
# Error Correlation Object
error.cor <- matrix(0, 6, 6)
diag(error.cor) <- 1</pre>
RTD <- binds(error.cor)
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
summary(CFA.Model)
#run(CFA.Model)
#CFA.Model2 <- extract(CFA.Model, y=1:3, e=1)</pre>
#summary(CFA.Model2)
```

SimVector-class

Vector object: Random parameters vector

Description

This object can be used to represent a vector in SEM model. It contains free parameters, fixed values, and starting values. This object can be represented mean, intercept, or variance vectors.

Objects from the Class

This object is created by bind function. Objects can be created by calls of the form new("SimVector", ...).

Slots

```
free: TBA
popParam: TBA
misspec: TBA
```

Methods

summaryShort Provides a short summary of all information in the object summary Provides a thorough description of all information in the object

sortList 107

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

SimMatrix for random parameter matrix

Examples

```
showClass("SimVector")

factor.mean <- rep(NA, 2)
factor.mean.starting <- c(5, 2)
AL <- bind(factor.mean, factor.mean.starting)
#run(AL)
summary(AL)
summaryShort(AL)</pre>
```

sortList

Sort two objects in a list

Description

Sort two objects in a list by swapping the values of both objects so that the first object contains the lower value and the second object contains the larger value

Usage

```
sortList(object)
```

Arguments

object

The list with two objects (e.g., vector, matrix)

Value

The sorted list

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
# No example
```

108 summaryConverge

summaryConverge	Provide a comparison between the characteristics of convergent repli-
	cations and nonconvergent replications

Description

This function provides a comparison between the characteristics of convergent replications and non-convergent replications. The comparison includes sample size (if varying), percent missing completely at random (if varying), percent missing at random (if varying), parameter values, misspecified-parameter values (if applicable), and population misfit (if applicable).

Usage

```
summaryConverge(object)
```

Arguments

```
object SimResult object being described
```

Value

A list with the following elements:

- Converged The number of convergent and nonconvergent replications
- n Sample size
- pmMCAR Percent missing completely at random
- pmMAR Percent missing at random
- paramValue Parameter values
- misspecValue Misspecified-parameter values
- popFit Population misfit

Each element will provide the head-to-head comparison between convergent and nonconvergent replications properties.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
## Not run:
path.BE <- matrix(0, 4, 4)
path.BE[3, 1:2] <- NA
path.BE[4, 3] <- NA
starting.BE <- matrix("", 4, 4)
starting.BE[3, 1:2] <- "runif(1, 0.3, 0.5)"
starting.BE[4, 3] <- "runif(1, 0.5, 0.7)"
mis.path.BE <- matrix(0, 4, 4)
mis.path.BE[4, 1:2] <- "runif(1, -0.1, 0.1)"
BE <- bind(path.BE, starting.BE, misspec=mis.path.BE)</pre>
```

summaryFit 109

```
residual.error <- diag(4)</pre>
residual.error[1,2] <- residual.error[2,1] <- NA</pre>
RPS <- binds(residual.error, "rnorm(1, 0.3, 0.1)")
loading <- matrix(0, 12, 4)</pre>
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loading[7:9, 3] <- NA
loading[10:12, 4] <- NA
mis.loading <- matrix("runif(1, -0.3, 0.3)", 12, 4)
mis.loading[is.na(loading)] <- 0</pre>
LY <- bind(loading, "runif(1, 0.7, 0.9)", misspec=mis.loading)
mis.error.cor <- matrix("rnorm(1, 0, 0.1)", 12, 12)
diag(mis.error.cor) <- 0</pre>
RTE <- binds(diag(12), misspec=mis.error.cor)</pre>
SEM.Model <- model(RPS = RPS, BE = BE, LY=LY, RTE=RTE, modelType="SEM")</pre>
n1 <- list(mean = 0, sd = 0.1)
chi5 <- list(df = 5)
facDist <- bindDist(c("chisq", "chisq", "norm", "norm"), chi5, chi5, n1, n1)</pre>
dat <- generate(SEM.Model, n=500, sequential=TRUE, facDist=facDist)</pre>
out <- analyze(SEM.Model, dat, estimator="mlr")</pre>
simOut <- sim(50, n=500, SEM.Model, sequential=TRUE, facDist=facDist, estimator="mlr")\\
summaryConverge(simOut)
## End(Not run)
```

summaryFit

Provide summary of model fit across replications

Description

This function will provide fit index cutoffs for values of alpha, and mean fit index values across all replications.

Usage

```
summaryFit(object, alpha = NULL)
```

Arguments

object SimResult to be summarized

alpha The alpha level used to find the fit indices cutoff. If there is no varying condition,

a vector of different alpha levels can be provided.

110 summaryMisspec

Value

A data frame that provides fit statistics cutoffs and means

When linkS4class{SimResult} has fixed simulation parameters the first colmns are fit index cutoffs for values of alpha and the last column is the mean fit across all replications. Rows are

- Chi Chi-square fit statistic
- AIC Akaike Information Criterion
- BIC Baysian Information Criterion
- · RMSEA Root Mean Square Error of Approximation
- CFI Comparative Fit Index
- TLI Tucker-Lewis Index
- · SRMR Standardized Root Mean Residual

When linkS4class{SimResult} has random simulation parameters (sample size or percent missing), columns are the fit indices listed above and rows are values of the random parameter.

See details in popDiscrepancy and popMisfitMACS

Author(s)

Alexander M. Schoemann (University of Kansas; <schoemann@ku.edu>) Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

SimResult for the result object input

Examples

```
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")
# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output <- sim(5, n=500, CFA.Model)
summaryFit(Output)</pre>
```

summaryMisspec

Provide summary of the population misfit and misspecified-parameter values across replications

Description

This function provides the summary of the population misfit and misspecified-parameter values across replications. The summary will be provided for the convergent replications only.

summaryParam 111

Usage

```
summaryMisspec(object)
```

Arguments

object SimResult object being described

Value

A data frame that provides the summary of population misfit and misspecified-parameter values imposed on the real parameters

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

SimResult for the object input

Examples

```
## Not run:
path <- matrix(0, 4, 4)
path[3, 1:2] <- NA
path[4, 3] <- NA
pathVal <- matrix("", 4, 4)</pre>
pathVal[3, 1:2] <- "runif(1, 0.3, 0.5)"</pre>
pathVal[4, 3] <- "runif(1, 0.5, 0.7)"</pre>
pathMis <- matrix(0, 4, 4)</pre>
pathMis[4, 1:2] <- "runif(1, -0.1, 0.1)"</pre>
BE <- bind(path, pathVal, pathMis)</pre>
residual.error <- diag(4)</pre>
residual.error[1,2] <- residual.error[2,1] <- NA</pre>
RPS <- binds(residual.error, "rnorm(1, 0.3, 0.1)")
Path.Model <- model(RPS = RPS, BE = BE, modelType="Path")
# The number of replications in actual analysis should be much more than 5
ParamObject <- sim(5, n=200, Path.Model)</pre>
summaryMisspec(ParamObject)
## End(Not run)
```

summaryParam

Provide summary of parameter estimates and standard error across replications

112 summaryParam

Description

This function will provide averages of parameter estimates, standard deviations of parameter estimates, averages of standard errors, and power of rejection with a priori alpha level for the null hypothesis of parameters equal 0.

Usage

```
summaryParam(object, alpha = 0.05, detail = FALSE)
```

Arguments

object SimResult object being described

alpha The alpha level used to find the statistical power of each parameter estimate detail If TRUE, more details about each parameter estimate are provided, such as rel-

ative bias, standardized bias, or relative standard error bias.

Value

A data frame that provides the statistics described above from all parameters. For using with linkS4class{SimResult}, each column means

- Estimate. Average: Average of parameter estimates across all replications
- Estimate. SD: Standard Deviation of parameter estimates across all replications
- Average . SE: Average of standard errors across all replications
- Power (Not equal 0): Proportion of significant replications when testing whether the parameters are different from zero
- Average.Param: Parameter values or average values of parameters if random parameters are specified
- SD.Param: Standard Deviations of parameters. Appeared only when random parameters are specified.
- Average.Bias: The difference between parameter estimates and parameter underlying data
- SD.Bias: Standard Deviations of bias across all replications. Appeared only when random parameters are specified. This value is the expected value of average standard error when random parameter are specified.
- Coverage: The percentage of (1-alpha)% confidence interval covers parameters underlying the data.
- Rel.Bias: Relative Bias, which is (Estimate.Average Average.Param)/Average.Param. Hoogland and Boomsma (1998) proposed that the cutoff of .05 may be used for acceptable relative bias. This option will be available when detail=TRUE. This value will not be available when parameter values are very close to 0.
- Std.Bias: Standardized Bias, which is (Estimate.Average-Average.Param)/Estimate.SD for fixed parameters and (Estimate.Average-Average.Param)/SD.Bias for random parameters. Collins, Schafer, and Kam (2001) recommended that biases will be only noticeable when standardized bias is greater than 0.4 in magnitude. This option will be available when detail=TRUE
- Rel.SE.Bias: Relative Bias in standard error, which is (Average.SE-Estimate.SD)/Estimate.SD for fixed parameters and (Average.SE-SD.Bias)/SD.Bias for random parameters. Hoogland and Boomsma (1998) proposed that 0.10 is the acceptable level. This option will be available when detail=TRUE

summaryPopulation 113

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

References

Collins, L. M., Schafer, J. L., & Kam, C. M. (2001). A comparison of inclusive and restrictive strategies in modern missing data procedures. *Psychological Methods*, *6*, 330-351.

Hoogland, J. J., & Boomsma, A. (1998). Robustness studies in covariance structure modeling. *Sociological Methods & Research*, 26, 329-367.

See Also

SimResult for the object input

Examples

```
showClass("SimResult")
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, 0.7)
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")

# We make the examples running only 5 replications to save time.
# In reality, more replications are needed.
Output <- sim(5, n=500, CFA.Model)
summaryParam(Output)
summaryParam(Output, detail=TRUE)</pre>
```

summary Population

Summarize the data generation population model underlying a result object

Description

Summarize the data generation population model underlying a result object

Usage

```
summaryPopulation(object)
```

Arguments

object

The result object that you wish to extract the data generation population model from (linkS4class{SimResult}).

Value

None except using for linkS4class{SimResult} which the return value is a data.frame of the summary of population model across replications.

114 summaryShort

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

• SimResult for result object

Examples

```
## Not run:
loading <- matrix(0, 6, 1)
loading[1:6, 1] <- NA
LX <- bind(loading, "runif(1, 0.4, 0.9)")
RPH <- binds(diag(1))
RTD <- binds(diag(6))
CFA.Model <- model(LY = LX, RPS = RPH, RTE = RTD, modelType="CFA")

# We will use only 10 replications to save time.
# In reality, more replications are needed.
Output <- sim(10, n=200, model=CFA.Model)
summaryPopulation(Output)

## End(Not run)</pre>
```

summaryShort

Provide short summary of an object.

Description

Provide short summary if it is available. Otherwise, it is an alias for summary.

Usage

```
summaryShort(object, ...)
```

Arguments

```
object Desired object being described ... any additional arguments
```

Value

NONE. This function will print on screen only.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

See Also

This is the list of classes that can use summaryShort method.

- SimMatrix
- SimVector

twoTailedPValue 115

Examples

```
loading <- matrix(0, 6, 2)
loading[1:3, 1] <- NA
loading[4:6, 2] <- NA
loadingValues <- matrix(0, 6, 2)
LX <- bind(loading, "runif(1, 0.8, 0.9)")
summaryShort(LX)</pre>
```

twoTailedPValue

Find two-tailed p value from one-tailed p value

Description

Find two-tailed p value from one-tailed p value

Usage

```
twoTailedPValue(vec)
```

Arguments

vec

A vector of one-tailed *p* value.

Value

A vector of two-tailed *p* value.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
# No example
```

validateCovariance

Validate whether all elements provides a good covariance matrix

Description

Validate whether all elements provides a good covariance matrix

Usage

```
validateCovariance(resVar, correlation, totalVar = NULL)
```

Arguments

resVar A vector of residual variances

correlation A correlation matrix

totalVar A vector of total variances

116 validateObject

Value

Return TRUE if the covariance matrix is good

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

No example

validateObject

Validate whether the drawn parameters are good.

Description

Validate whether the drawn parameters are good (providing an identified model).

Usage

```
validateObject(paramSet)
```

Arguments

paramSet

A target set of parameters

Value

Return TRUE if the target parameters are good.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

No example

validatePath 117

validatePath	Validate whether the regression coefficient (or loading) matrix is good
	0 10

Description

Validate whether the regression coefficient (or loading) matrix is good

Usage

```
validatePath(path, var.iv, var.dv)
```

Arguments

path A regression coefficient or loading matrix

var.iv The variances of variables corresponding to the columns var.dv The variances of variables corresponding to the rows

Value

Return TRUE if the target regression coefficient matrix is good.

Author(s)

Sunthud Pornprasertmanit (University of Kansas; psunthud@ku.edu)

Examples

No example

whichMonotonic	Extract a part of a vector that is monotonically increasing or decreas-
	ing

Description

Extract a part of a vector that is monotonically increasing or decreasing. This function will go to the anchor value and extract the neighbor values that are monotonically increasing or decreasing.

Usage

```
whichMonotonic(vec, ord=NULL, anchor=NULL)
```

Arguments

vec The target vector to be extracted

ord The names of each element of the vector to be attached

anchor The position of the element to be anchored. The default value is the middle

position.

118 whichMonotonic

Value

The monotonic part of a vector

Author(s)

Sunthud Pornprasertmanit (University of Kansas; <psunthud@ku.edu>)

Examples

```
# This is a private function.
# whichMonotonic(c(3, 4, 1, 2, 3, 5, 2, 1))
```

Index

*Topic classes	generate, 7, 13, 17, 33, 60, 62
SimDataDist-class, 100	getCondQtile, 36
SimMatrix-class, 101	getCutoff, 36, 37, 39, 41, 45, 47, 48, 65, 76,
SimMissing-class, 102	104
SimResult-class, 103	getCutoff,data.frame-method
SimVector-class, 106	(getCutoff), 37
	<pre>getCutoff, matrix-method (getCutoff), 37</pre>
all.equal, <u>62</u>	<pre>getCutoff,SimResult-method(getCutoff),</pre>
analyze, 3, 7, 17, 60, 62	37
anova, 4, <i>104</i>	<pre>getCutoff-methods (getCutoff), 37</pre>
anova, SimResult-method (anova), 4	getCutoffNested, 38, 41, 67, 79
	getCutoffNonNested, 40, 50, 68, 81
bind, 5, 101, 106	getExtraOutput, 41, 98
bindDist, 8, 100, 101	getKeywords, 42
binds (bind), 5	getPopulation, 43, 104
alaan O	getPopulation,SimResult-method
<pre>clean, 9 cleanSimResult, 9</pre>	(SimResult-class), 103
	getPower, 29, 30, 32, 44, 75, 87
continuousPower, 10, 29, 30, 32, 87	getPowerFit, 45, 76, 104
createData, 11, <i>12</i> , <i>14</i> , <i>35</i>	<pre>getPowerFit,data.frame,vector-method</pre>
draw, 12, 13, 14, 17, 35, 60, 62, 99	(getPowerFit), 45
,,,,,,,,	<pre>getPowerFit,matrix,vector-method</pre>
estmodel, 15	(getPowerFit), 45
extract,SimVector-method	<pre>getPowerFit,SimResult,missing-method</pre>
(SimVector-class), 106	(getPowerFit), 45
extractLavaanFit, 17	<pre>getPowerFit,SimResult,vector-method</pre>
	(getPowerFit), 45
find2Dhist, 18	<pre>getPowerFit-methods (getPowerFit), 45</pre>
findFactorIntercept, 18, 20-27	getPowerFitNested, 47, 79
findFactorMean, 19, 19, 21-27	<pre>getPowerFitNested,SimResult,SimResult,missing-method</pre>
findFactorResidualVar, 19, 20, 20, 22-27	(getPowerFitNested), 47
findFactorTotalCov, 19–21, 21, 23–27	<pre>getPowerFitNested,SimResult,SimResult,vector-method</pre>
findFactorTotalVar, 19-22, 23, 24-27	(getPowerFitNested), 47
findIndIntercept, 19–23, 24, 25–27	getPowerFitNested-methods
findIndMean, 19–24, 25, 26, 27	(getPowerFitNested), 47
findIndResidualVar, 19-25, 26, 27	getPowerFitNonNested, 49, 81
findIndTotalVar, 19-26, 27	${\tt getPowerFitNonNested, SimResult, SimResult, missing-method}$
findphist, 28	(getPowerFitNonNested), 49
findPossibleFactorCor, 28, 31	${\tt getPowerFitNonNested}, {\tt SimResult}, {\tt SimResult}, {\tt vector-method}$
findPower, 29, 32	(getPowerFitNonNested), 49
findRecursiveSet, 29, 30, 31	getPowerFitNonNested-methods
findRowZero, 31	(getPowerFitNonNested), 49
findTargetPower, 32	
fitMeasuresChi, 33	impose (imposeMissing), 51

120 INDEX

imposeMissing, 51, 57, 102, 103	pValueNested, 55, 91
interpolate, 53	pValueNonNested, 55, 92
	pValueVariedCutoff,94
lavaan, 41, 54, 61, 88, 91, 93, 98	
likRatioFit, 54	revText, 95
loadingFromAlpha, 56	
	setPopulation, 96, 104
miss, 56, 98	setPopulation,SimResult,data.frame-method
model, 7, 17, 58, 62, 98, 105	(SimResult-class), 103
model.lavaan, 61	setPopulation,SimResult,SimSet-method
multipleAllEqual, 62	(SimResult-class), 103
Mvdc, 8	setPopulation,SimResult,VirtualRSet-method
	(SimResult-class), 103
overlapHist, 63	sim, 17, 42, 53, 57, 60, 62, 98, 103, 104
nlo+200+;lo 64	SimDataDist, 8, 12, 34, 35, 69, 70, 98, 99
plot3DQtile, 64 plotCutoff, 65, 104	SimDataDist-class, 100
plotCutoff, data.frame-method	SimMatrix, 5, 7, 58, 59, 107, 114
(plotCutoff), 65	SimMatrix-class, 101
plotCutoff, SimResult-method	SimMissing, <i>51</i> , <i>53</i> , <i>57</i>
(plotCutoff), 65	SimMissing-class, 102
plotCutoff-methods (plotCutoff), 65	SimResult, 4, 5, 8–11, 37–41, 43–50, 54, 55,
plotCutoffNested, 66, 79	65-68, 72, 74-76, 78-81, 88, 89,
plotCutoffNonNested, 41, 68, 81	91–94, 96, 98, 108–114
plotDist, 69	SimResult-class, 103
plotDist, SimDataDist-method	SimSem, 7, 13, 17, 33-35, 60, 62, 98
(SimDataDist-class), 100	SimSem-class, 105
plotIndividualScatter, 70	SimVector, 5, 7, 58, 59, 102, 114
plotLogisticFit, 71	SimVector-class, 106
plotMisfit, 72	sortList, 107
plotOverHist, 73	summary, 96, 103, 104, 106
plotPower, 74, 82	summary,SimDataDist-method
plotPowerFit, 71–73, 75, 77, 78, 84, 104	(SimDataDist-class), 100
plotPowerFitDf, 77	summary,SimMatrix-method
plotPowerFitNested, 78	(SimMatrix-class), 101
plotPowerFitNonNested, 80	summary, SimMissing-method
plotPowerSig, 82	(SimMissing-class), 102
plotQtile, 83	summary, SimResult-method
plotScatter, 83	(SimResult-class), 103
popDiscrepancy, 84, 85, 86, 110	<pre>summary,SimSem-method(SimSem-class),</pre>
popMisfitMACS, 85, 104, 110	105
predProb, 86	summary, SimVector-method
printIfNotNull, 87	(SimVector-class), 106
pValue, 88, 90, 95	summaryConverge, 108
pValue, ANY-method (pValue), 88	summaryFit, 109
pValue,lavaan,SimResult-method	summaryMisspec, 110
(pValue), 88	summaryParam, <i>104</i> , 111
pValue, numeric, data. frame-method	summaryPopulation, <i>104</i> , 113
(pValue), 88	summaryPopulation,SimResult-method
pValue, numeric, vector-method (pValue),	(SimResult-class), 103
88	summaryShort, <i>101</i> , <i>106</i> , 114
pValue-methods (pValue), 88	summaryShort, ANY-method (summaryShort),
pValueCondCutoff, 90	114

INDEX 121