Package ‘MplusAutomation’

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Type Package
Title Automating Mplus Model Estimation and Interpretation
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Description The MplusAutomation package leverages the flexibility of the R language to automate latent variable model estimation and interpretation using Mplus, a powerful latent variable modeling program developed by Muthen and Muthen (www.statmodel.com). Specifically, MplusAutomation provides routines for creating related groups of models, running batches of models, and extracting and tabulating model parameters and fit statistics.
License LGPL-3
URL https://github.com/michaelhallquist/MplusAutomation
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Change directory

The function takes a path and changes the current working directory to the path. If the directory specified in the path does not currently exist, it will be created.

Usage

\texttt{cd(base, pre, num)}
arguments

base: a character string with the base path to the directory. This is required.
pre: an optional character string with the prefix to add to the base path. Non character strings will be coerced to character class.
num: an optional character string, prefixed by pre. Non character strings will be coerced to character class.

Details

The function has been designed to be platform independent, although it has had limited testing. Path creation is done using file.path, the existence of the directory is checked using file.exists and the directory created with dir.create. Only the first argument, is required. The other optional arguments are handy when one wants to create many similar directories with a common base.

Value

NULL, changes the current working directory

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

Examples

## Not run:
# an example just using the base
cd("~/testdir")

# an example using the optional arguments
base <- "~/testdir"
pre <- "test_"

cd(base, pre, 1)
cd(base, pre, 2)

## End(Not run)

description

This is a method for returning the coefficients of an mplus.model object. It works directly on an object stored from readModels such as: object <- readModels("/path/to/model/model.out"). Method that calls coef.mplus.model. See further documentation there.
Usage

```r
## S3 method for class 'mplus.model'
coef(object, type = c("un", "std", "stdy", "stdyx"),
     params = c("regression", "loading", "undirected", "expectation",
               "variability", "new"), ..., raw = FALSE)

## S3 method for class 'mplusObject'
coef(object, ...)```

Arguments

- **object**: An object of class mplusObject
- **type**: A character vector indicating the type of coefficients to return. One of “un”, “std”, “stdy”, or “stdyx”.
- **params**: A character vector indicating what type of parameters to extract. Any combination of “regression”, “loading”, “undirected”, “expectation”, “variability”, and “new”. A single one can be passed or multiple. By default, all are used and all parameters are returned.
- **...**: Additional arguments to pass on (not currently used)
- **raw**: A logical defaulting to FALSE indicating whether to parse and return coefficients based on the type (regression, etc.) and relabel using an arrow notation, or to return the raw coefficients in a named vector.

Value

Either a data frame of class ‘mplus.model.coefs’, or in the case of multiple group models, a list of class ‘mplus.model.coefs’, where each element of the list is a data frame of class ‘mplus.model.coefs’, or a named vector of coefficients, if raw=TRUE.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

- `readModels`
- Other Mplus.Formatting: `extract, extract.mplus.model-method, extract.mplusObject-method, extract.mplus.model, extract.mplusObject; print.MplusRstructure; summary.mplusObject`

Examples

```r
## Not run:
# simple example of a model using builtin data
# demonstrates use
test <- mplusObject(
    TITLE = "test the MplusAutomation Package;",
    MODEL = "
    mpg ON wt hp;
```
compareModels

Compare the output of two Mplus models

Description

The compareModels function compares the output of two Mplus files and prints similarities and differences in the model summary statistics and parameter estimates. Options are provided for filtering out fixed parameters and nonsignificant parameters. When requested, compareModels will compute the chi-square difference test for nested models (does not apply to MLMV, WLSM, and WLSMV estimators, where DIFFTEST in Mplus is needed). Model outputs to be compared can be full summaries and parameters (generated by readModels), summary statistics only (extractModelSummaries), or parameters only (extractModelParameters).

Usage

compareModels(m1, m2, show = "all", equalityMargin = c(param = 1e-04, pvalue = 1e-04), compare = "unstandardized", sort = "none", showFixed = FALSE, showNS = TRUE, diffTest = FALSE)

Arguments

m1 The first Mplus model to be compared. Generated by readModels, extractModelSummaries, or extractModelParameters.
m2 The second Mplus model to be compared.
compareModels

show What aspects of the models should be compared. Options are "all", "summaries", "equal", "diff", "pdiff", and "unique". See below for details.

equalityMargin Defines the discrepancy between models that is considered equal. Different margins can be specified for p-value equality versus parameter equality. Defaults to .0001 for both.

compare Which parameter estimates should be compared. Options are "unstandardized", "stdyx.standardized" "stdy.standardized", and "std.standardized".

sort How to sort the output of parameter comparisons. Options are "none", "type", "alphabetical", and "maxDiff". See below for details.

showFixed Whether to display fixed parameters in the output (identified where the est/se = 999.000, per Mplus convention). Default to FALSE.

showNS Whether to display non-significant parameter estimates. Can be TRUE or FALSE, or a numeric value (e.g., .10) that defines what p-value is filtered as non-significant.

diffTest Whether to compute a chi-square difference test between the models. Assumes that the models are nested. Not available for MLMV, WLSMV, and ULSMV estimators. Use DIFFTEST in Mplus instead.

Details

The show parameter can be one or more of the following, which can be passed as a vector, such as c("equal", "pdiff").

show "all" Display all available model comparison. Equivalent to c("summaries", "equal", "diff", "pdiff", "unique").


"allsummaries" Prints a comparison of all summary statistics available in each model. May generate a lot of output.

"equal" Print parameter estimates that are equal between models (i.e., \( \leq \) equalityMargin["param"]).

"diff" Print parameter estimates that are different between models (i.e., \( > \) equalityMargin["param"]).

"pdiff" Print parameter estimates where the p-values differ between models (i.e., \( > \) equalityMargin["pvalue"]).

"unique" Print parameter estimates that are unique to each model.

The sort parameter determines the order in which parameter estimates are displayed. The following options are available:

sort "none" No sorting is performed, so parameters are output in the order presented in Mplus. (Default)

"type" Sort parameters by their role in the model. This groups output by regression coefficient (ON), factor loadings (BY), covariances (WITH), and so on. Within each type, output is alphabetical.

"alphabetical" Sort parameters in alphabetical order.

"maxDiff" Sort parameter output by the largest differences between models (high to low).
createModels

Value

No value is returned by this function. It is used to print model differences to the R console.

Author(s)

Michael Hallquist

Examples

# make me!!!

createModels  

Create Mplus Input Files from Template

Description

The createModels function processes a single Mplus template file and creates a group of related model input files. Definitions and examples for the template language are provided in the MplusAutomation vignette and are not duplicated here at the moment. See this PDF: http://cran.r-project.org/web/packages/MplusAutomation/vignettes/Vignette.pdf

Usage

createModels(templatefile)

Arguments

templatefile  
The filename (absolute or relative path) of an Mplus template file to be processed. Example “C:/MplusTemplate.txt”

Value

No value is returned by this function. It is solely used to process an Mplus template file.

Author(s)

Michael Hallquist

Examples

## Not run:
createModels("L2 Multimodel Template No iter.txt")

## End(Not run)
createSyntax  

Create the Mplus input text for an mplusObject

Description

This function takes an object of class mplusObject and creates the Mplus input text corresponding to it, including data link and variable names.

Usage

createSyntax(object, filename, check = TRUE, add = FALSE)

Arguments

- **object**: An object of class mplusObject
- **filename**: The name of the data file as a character vector
- **check**: A logical indicating whether or not to run parseMplus on the created input file. Checks for errors like lines that are too long, or for missing semi-colons and gives notes.
- **add**: A logical passed on to parseMplus whether to add semi colons to line ends. Defaults to FALSE.

Value

A character string containing all the text for the Mplus input file.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

prepareMplusData, mplusModeler

Examples

```r
# example mplusObject
eexample1 <- mplusObject(MODEL = "mpg ON wt;",
                         usevariables = c("mpg", "hp"), rdata = mtcars)

# create the Mplus input text
cat(createSyntax(example1, "example1.dat"), fill=TRUE)

# update the object, then create input text
cat(createSyntax(update(example1,
                  TITLE = "This is my title;",
                  MODEL = "mmpg ON hp;",
                  usevariables = c("mpg", "hp", "wt")), "example1.dat")
```

extract

fill=TRUE)
rm(example1)

extract

Extract function to make Mplus output work with the texreg package

Description

This is a method for extracting output in a format suitable for the texreg package. Uses coef for most the work.

Usage

extract.mplus.model(model, summaries = "none", ...)

extract.mplusObject(model, summaries = "none", ...)

## S4 method for signature 'mplus.model'
extract(model, summaries = "none", ...)

## S4 method for signature 'mplusObject'
extract(model, summaries = "none", ...)

Arguments

model A Mplus model object. This typically comes either from readModels directly, or indirectly via mplusModeler. The results will have different classes, but extract methods are defined for both.

summaries A character vector which summaries to include. Defaults to "none".

... Additional arguments passed to coef.mplus.model.

Value

A texreg object, or for multiple group models, a list of texreg objects.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

readModels

Other Mplus.Formatting: coef.mplus.model, coef.mplusObject; print.MplusRstructure; summary.mplusObject
Examples

```r
## Not run:
## simple example of a model using builtin data
## demonstrates use
test <- mplusObject(
  TITLE = "test the MplusAutomation Package;",
  MODEL = "
    mpg ON wt hp;
    wt WITH hp;",
  OUTPUT = "STANDARDIZED;",
  usevariables = c("mpg", "wt", "hp"),
  rdata = mtcars)

res <- mplusModeler(test, "mtcars.dat", modelout = "model1.inp", run = 1L)

extract(res$results)
## there is also a method for mplusObject class
extract(res)

screenreg(extract(res))
screenreg(extract(res, type = 'stdyx'))

screenreg(extract(res, type = 'un', params = 'regression'),
  single.row=TRUE)
screenreg(extract(res, type = 'un', params = 'regression', summaries = 'CFI'),
  single.row=TRUE)

## remove files
unlink("mtcars.dat")
unlink("model1.inp")
unlink("model1.out")
unlink("Mplus Run Models.log")
## End(Not run)
```

---

`extractModelParameters`

Extract model parameters from `MODEL RESULTS` section.

**Description**

Extracts the model parameters from the `MODEL RESULTS` section of one or more Mplus output files. If a particular output file has more than one results section (unstandardized, stdyx, stdy, and/or std), a list will be returned. If the target is a directory, all .out files therein will be parsed and a single list will be returned, where the list elements are named by the output file name. Returned parameters often include the parameter estimate, std. err, param/s.e., and two-tailed p-value.
extractModelParameters

Usage

extractModelParameters(target = getwd(), recursive = FALSE, filefilter, dropDimensions = FALSE, resultType)

Arguments

target the directory containing Mplus output files (.out) to parse OR the single output file to be parsed. May be a full path, relative path, or a filename within the working directory. Defaults to the current working directory. Example: “C:/Users/Michael/Mplus Runs”

recursive optional. If TRUE, parse all models nested in subdirectories within target. Defaults to FALSE.

filefilter a Perl regular expression (PCRE-compatible) specifying particular output files to be parsed within directory. See regex or http://www.pcre.org/pcre.txt for details about regular expression syntax.

dropDimensions Relevant only for multi-file parsing. If TRUE, then if only one output section (usually unstandardized) is present for all files in the parsed list, then eliminate the second-level list (which contains elements for each output section). The result is that the elements of the returned list are data.frame objects with the relevant parameters.

resultType N.B.: this parameter is deprecated and will be removed in a future version. The new default is to extract all results that are present and return a list (see below for details). resultType specified the results section to extract. If raw, the unstandardized estimates will be returned. “stdyx”, “stdy”, and “std” are the other options, which extract different standardized solutions. See the Mplus User’s Guide for additional details about the differences in these standardizations.

Value

If target is a single file, a list containing unstandardized and standardized results will be returned. If all standardized solutions are available, the list element will be named: unstandardized, stdyx.standardized, stdy.standardized, and std.standardized. If confidence intervals are output using OUTPUT:CINTERVAL, then a list element named ci.unstandardized will be included. Each of these list elements is a data.frame containing relevant model parameters.

If target is a directory, a list will be returned, where each element contains the results for a single file, and the top-level elements are named after the corresponding output file name. Each element within this list is itself a list, with elements as in the single file case above.

The core data.frame for each MODEL RESULTS section typically has the following structure:

paramHeader The header that begins a given parameter set. Example: "FACTOR1 BY"
param The particular parameter being measured (within paramHeader). Example: "ITEM1"
est Parameter estimate value.
se Standard error of the estimate
est_se Quotient of est/se, representing z-test/t-test in large samples
pval Two-tailed p-value for the est_se quotient.
In the case of output from Bayesian estimation (ESTIMATOR=BAYES), the data.frame will contain a different set of variables, including some of the above, as well as

- **posterior_sd** Posterior standard deviation of the estimate.
- **lower_2.5ci** Lower 2.5 percentile of the estimate.
- **upper_2.5ci** Upper 2.5 percentile (aka 97.5 percentile) of the estimate.

Also note that the pval column for Bayesian output represents a one-tailed estimate.

In the case of output from a Monte Carlo study (MONTECARLO: and MODEL POPULATION:), the data.frame will contain a different set of variables, including some of the above, as well as

- **population** Population parameter value.
- **average** Average parameter estimate across replications.
- **population_sd** Standard deviation of parameter value in population across replications.
- **average_se** Average standard error of estimated parameter value across replications.
- **mse** Mean squared error.
- **cover_95** Proportion of replications whose 95% confidence interval for the parameter includes the population value.
- **pct_sig_coef** Proportion of replications for which the two-tailed significance test of the parameter is significant (p < .05).

In the case of confidence interval output (OUTPUT:CINTERVAL), the list element ci.unstandardized will contain a different set of variables, including some of the above, as well as

- **low.5** Lower 0.5% CI estimate.
- **low2.5** Lower 2.5% CI estimate.
- **low5** Lower 5% CI estimate.
- **est** Parameter estimate value.
- **up5** Upper 5% (i.e., 95%) CI estimate.
- **up2.5** Upper 2.5% (i.e., 97.5%) CI estimate.
- **up.5** Upper 0.5% (i.e., 99.5%) CI estimate.

If the model contains multiple latent classes, an additional variable, LatentClass, will be included, specifying the latent class number. Also, the Categorical Latent Variables section will be included as LatentClass "Categorical.Latent.Variables."

If the model contains multiple groups, Group will be included.

If the model contains two-level output (between/within), BetweenWithin will be included.

**Author(s)**

Michael Hallquist

**See Also**

extractModelSummaries
**Examples**

```r
## Not run:
ex3.14 <- extractModelParameters(
## End(Not run)
```

**extractModelSummaries**  
*Extract summary statistics from a single output file or from a group of Mplus models within a directory*

**Description**

Parses a group of Mplus model output files (.out extension) for model fit statistics. At this time, the details extracted are fixed and include: Filename, InputInstructions, Title, Estimator, LL, BIC, aBIC, AIC, AICC. The infrastructure is in place to allow for user-specified selection of summary statistics in future versions.

**Usage**

`extractModelSummaries(target = getwd(), recursive = FALSE, filefilter)`

**Arguments**

- **target**: the directory containing Mplus output files (.out) to parse OR the single output file to be parsed. Defaults to the current working directory. Example: "C:/Users/Michael/Mplus Runs"
- **recursive**: optional. If TRUE, parse all models nested in subdirectories within directory. Defaults to FALSE.
- **filefilter**: a Perl regular expression (PCRE-compatible) specifying particular output files to be parsed within directory. See regex or [http://www.pcre.org/pcre.txt](http://www.pcre.org/pcre.txt) for details about regular expression syntax.

**Value**

Returns a data.frame containing model fit statistics for all output files within directory. The data.frame contains some of the following variables (depends on model type):

- **Title**: Title for the model, specified by the TITLE: command
- **Filename**: Filename of the output file
- **Estimator**: Estimator used for the model (e.g., ML, MLR, WLSMV, etc.)
- **LL**: Log-likelihood of the model
- **BIC**: Bayesian Information Criterion
- **aBIC**: Sample-Size-Adjusted BIC (Slove, 1987)
- **AIC**: Akaike’s Information Criterion
AICC  Corrected AIC, based on Sugiura (1978) and recommended by Burnham & Anderson (2002)
DIC  Deviance Information Criterion. Available in ESTIMATOR=BAYES output.
Parameters  Number of parameters estimated by the model
pD  Estimated number of parameters in Bayesian output
Observations  The number of observations for the model (does not support multiple-groups analysis at this time)
CFI  Confirmatory Fit Index
TLI  Tucker-Lewis Index
RMSEA_Estimate  Point estimate of root mean squared error of approximation
RMSEA_90CI_LB  Lower bound of the 90% Confidence Interval around the RMSEA estimate.
RMSEA_90CI_UB  Upper bound of the 90% Confidence Interval around the RMSEA estimate.
RMSEA_pLT05  Probability that the RMSEA estimate falls below .05, indicating good fit.
ChiSqM_Value  Model chi-squared value
ChiSqM_DF  Model chi-squared degrees of freedom
ChiSqM_PValue  Model chi-squared p-value
ObsRepChiSqDiff_95CI_LB  Lower bound of 95% confidence interval for the difference between observed and replicated chi-square values
ObsRepChiSqDiff_95CI_UB  Upper bound of 95% confidence interval for the difference between observed and replicated chi-square values
PostPred_PValue  Posterior predictive p-value
BLRT_RequestedDraws  Number of requested bootstrap draws for TECH14.
BLRT_KM1LL  Log-likelihood of the K-1 model (one less class) for the Bootstrapped Likelihood Ratio Test (TECH14).
BLRT_2xLLDiff  Two times the log-likelihood difference of the models with K and K-1 classes (TECH14).
BLRT_ParamDiff  Difference in the number of parameters for models with K and K-1 classes (TECH14).
BLRT_PValue  P-value of the Bootstrapped Likelihood Ratio Test (TECH14) testing whether the K class model is significantly better than K-1
BLRT_SuccessfulDraws  The number of successful bootstrapped samples used in the Bootstrapped Likelihood Ratio Test
SRMR  Standardized root mean square residual
SRMR_Between  For TYPE=TWOLEVEL output, standardized root mean square residual for between level
SRMR_Within  For TYPE=TWOLEVEL output, standardized root mean square residual for within level
extractModelSummaries

WMR
Weighted root mean square residual

ChiSqBaseline_Value
Baseline (unstructured) chi-squared value

ChiSqBaseline_DF
Baseline (unstructured) chi-squared degrees of freedom

ChiSqBaseline_PValue
Baseline (unstructured) chi-squared p value

NumFactors
For TYPE=EFA output, the number of factors

T11_KM1Starts
TECH11: Number of initial stage random starts for k-1 model

T11_KM1Final
TECH11: Number of final stage optimizations for k-1 model

T11_KM1LL
TECH11: Log-likelihood of the K-1 model used for the Vuong-Lo-Mendell-Rubin LRT

T11_VLMR_2xLLDiff
TECH11: 2 * Log-likelihood Difference of K-class vs. K-1-class model for the Vuong-Lo-Mendell-Rubin LRT

T11_VLMR_ParamDiff
TECH11: Difference in number of parameters between K-class and K-1-class model for the Vuong-Lo-Mendell-Rubin LRT

T11_VLMR_Mean
TECH11: Vuong-Lo-Mendell-Rubin LRT mean

T11_VLMR_SD
TECH11: Vuong-Lo-Mendell-Rubin LRT standard deviation

T11_VLMR_PValue
TECH11: Vuong-Lo-Mendell-Rubin LRT p-value

T11_LMR_Value
TECH11: Lo-Mendell-Rubin Adjusted LRT value

T11_LMR_PValue
TECH11: Lo-Mendell-Rubin Adjusted LRT p-value

Author(s)
Michael Hallquist

See Also
regex, runModels, readModels

Examples

```r
## Not run:
allExamples <- extractModelSummaries(
  "C:/Program Files/Mplus/Mplus Examples/User's Guide Examples")

## End(Not run)
```
**extractModIndices**

*Extract model modification indices.*

**Description**

Extracts the model modification indices from the MODEL MODIFICATION INDICES section of one or more Mplus output files. If the target is a directory, all .out files therein will be parsed and a single list will be returned, where the list elements are named by the output file name. Returned parameters typically include the pairwise relationships between variables to be freed, the change in model chi-square (M.I.), and the expected parameter change (E.P.C.).

**Usage**

```r
extractModIndices(target = getwd(), recursive = FALSE, filefilter)
```

**Arguments**

- `target`: the directory containing Mplus output files (.out) to parse OR the single output file to be parsed. May be a full path, relative path, or a filename within the working directory. Defaults to the current working directory. Example: “C:/Users/Michael/Mplus Runs”
- `recursive`: optional. If TRUE, parse all models nested in subdirectories within target. Defaults to FALSE.
- `filefilter`: a Perl regular expression (PCRE-compatible) specifying particular output files to be parsed within directory. See `regex` or [http://www.pcre.org/pcre.txt](http://www.pcre.org/pcre.txt) for details about regular expression syntax.

**Value**

If `target` is a single file, a data.frame containing modification results for the target output file will be returned. If `target` is a directory, a list will be returned, where each element contains a data.frame of the modification indices for a single file, and the top-level elements are named after the corresponding output file name. The basic data.frame containing the MODEL MODIFICATION INDICES section of outfile. Variables include:

- `modV1`: The first variable in the pair to be freed according to the M.I.
- `operator`: The suggested relationship between modV1 and modV2 (e.g., WITH for freeing the covariance between modV1 and modV2)
- `modV2`: The first variable in the pair to be freed according to the M.I.
- `MI`: The decrease in model chi-square if the specified relationship is freely estimated
- `EPC`: The expected parameter estimate between modV1 and modV2 if freed.
- `Std_EPC`: The EPC value standardized using the variances of the continuous latent variables.
- `StdYX_EPC`: The EPC value standardized using the variances of the continuous latent variables as well as the variances of the background and/or outcome variables.
getSavedata_Bparams

Author(s)

Michael Hallquist

See Also

readModels, extractModelSummaries, extractModelParameters

Examples

```r
## Not run:
ex3.14 <- extractModIndices(

## End(Not run)
```

getSavedata_Bparams Load the draws from the Bayesian model posterior distribution (SAVE-
DATA BPARAMETERS) command into an R data.frame

Description

This function reads a the BPARAMETERS output file from the Mplus SAVEDATA BPARAME-
TERS command and returns an R data.frame object.

Usage

getSavedata_Bparams(outfile, discardBurnin = TRUE)

Arguments

outfile Required. The name of the Mplus output file to read. Can be an absolute or
relative path. If outfile is a relative path or just the filename, then it is assumed
that the file resides in the working directory getwd().

discardBurnin Optional. Whether to discard the burn-in phase of each MCMC chain (i.e., the
first half).

Value

A list containing the draws from the MCMC chains for a Bayesian model that uses the SAVE-
DATA BPARAMETERS command. Each list element corresponds to a single MCMC chain, as
specified by the ANALYSIS: CHAINS syntax in Mplus. If discardBurnin is FALSE, then a superor-
dinate list is provided that divides output in terms of burn-in versus valid draw halves of the MCMC
chains. For documentation of how Mplus implements chain convergence checks and MCMC draws,
**Note**

Note that the outfile parameter should refer to the Mplus output file (.out extension), not the actual dataset generated by SAVEDATA. This function reads information about the dataset from the .out file and loads the dataset accordingly.

**Author(s)**

Michael Hallquist, Florian Boeing-Messing

**References**


**See Also**

`getSavedata_Fileinfo, getSavedata_Data`

**Examples**

```r
## Not run:
fileInfo <- getSavedata_Data("C:/Program Files/Mplus/Test Output.out")

## End(Not run)
```

---

**getSavedata_Data**

Load an analysis dataset from the SAVEDATA command into an R `data.frame`

**Description**

This function reads an analysis dataset generated by the Mplus SAVEDATA command and returns an R `data.frame` object.

**Usage**

`getSavedata_Data(outfile)`

**Arguments**

- `outfile` Required. The name of the Mplus output file to read. Can be an absolute or relative path. If `outfile` is a relative path or just the filename, then it is assumed that the file resides in the working directory `getwd()`.

**Value**

A `data.frame` containing the analysis dataset generated by the SAVEDATA command.
Note

Note that the outfile parameter should refer to the Mplus output file (.out extension), not the actual dataset generated by SAVEDATA. This function reads information about the dataset from the .out file and loads the dataset accordingly.

Author(s)

Michael Hallquist

See Also

getsavedata_fileinfo

Examples

```r
## Not run:
savedat <- getsavedata_data("C:/Program Files/Mplus/Test Output.out")

## End(Not run)
```

---

**getsavedata_fileinfo**  
Read Variable Names, Formats, and Widths from data generated by the SAVEDATA Command

**Description**

This function reads the SAVEDATA INFORMATION section from an Mplus output file that used the SAVEDATA command, and it returns a list with the filename, variable names, variable formats, and variable widths of the SAVEDATA file. If present, the function also parses information about the Bayesian Parameters (BPARAMETERS) file.

**Usage**

```r
getsavedata_Fileinfo(outfile)
```

**Arguments**

- **outfile**  
  required. The name of the Mplus output file to read. Can be an absolute or relative path. If outfile is a relative path or just the filename, then it is assumed that the file resides in the working directory getwd().

**Value**

Returns a list of SAVEDATA file information that includes:

- **fileName**  
  The name of the file containing the analysis dataset created by the Mplus SAVEDATA command.

- **fileVarNames**  
  A character vector containing the names of variables in the dataset.
fileVarFormats A character vector containing the Fortran-style formats of variables in the dataset.
fileVarWidths A numeric vector containing the widths of variables in the dataset (which is stored in fixed-width format).
bayesFile The name of the BPARAMETERS file containing draws from the posterior distribution created by the Mplus SAVEDATA BPARAMETERS command.
bayesVarNames A character vector containing the names of variables in the BPARAMETERS dataset.
techn3File A character vector of the tech 3 output.
techn4File A character vector of the tech 4 output.

Author(s)
Michael Hallquist

See Also
getSavedata_Data

Examples

```r
## Not run:
fileInfo <- getSavedata_Fileinfo("C:/Program Files/Mplus/Test Output.out")

## End(Not run)
```

---

**HTMLSummaryTable** Create an HTML file containing a summary table of Mplus model statistics

**Description**

Creates an HTML file containing a summary table of model fit statistics extracted using the extractModelSummaries function. By default, the following summary statistics are included: Title, LL, Parameters, AIC, AICC, BIC, RMSEA_Est, but these are customizable using the keepCols and dropCols parameters.

**Usage**

```r
HTMLSummaryTable(modelList, filename = file.path(getwd(), "Model Comparison.html"), keepCols, dropCols, sortBy, display = FALSE)
```
### Arguments

- **modellist**: A list of models (as a `data.frame`) returned from the `extractModelSummaries` function.
- **filename**: The name of the HTML file to be created. Can be an absolute or relative path. If `filename` is a relative path or just the filename, then it is assumed that the file resides in the working directory `getwd()`. Example: "Mplus Summary.html"
- **keepCols**: A vector of character strings indicating which columns/variables to display in the summary. Only columns included in this list will be displayed (all others excluded). By default, `keepCols` is: c("Title", "LL", "Parameters", "AIC", "AICC", "BIC", "RMSEA"). Example: c("Title", "LL", "AIC", "CFI")
- **dropCols**: A vector of character strings indicating which columns/variables to omit from the summary. Any column not included in this list will be displayed. By default, `dropCols` is `NULL`. Example: c("InputInstructions", "TLI")
- **sortBy**: optional. Field name (as character string) by which to sort the table. Typically an information criterion (e.g., "AIC" or "BIC") is used to sort the table. Defaults to "AICC".
- **display**: optional. This parameter specifies whether to display the table in a web browser upon creation (TRUE or FALSE).

### Value

No value is returned by this function. It is solely used to create an HTML file containing summary statistics.

### Note

You must choose between `keepCols` and `dropCols` because it is not sensible to use these together to include and exclude columns. The function will error if you include both parameters.

### Author(s)

Michael Hallquist

### See Also

- `extractModelSummaries`, `showSummaryTable`, `latexSummaryTable`

### Examples

# make me!!!
Description

Creates a LaTeX-formatted summary table of model fit statistics extracted using the `extractModelSummaries` function. The table syntax is returned by the function, which is useful for embedding LaTeX tables using Sweave. By default, the following summary statistics are included: Title, LL, Parameters, AIC, AICC, BIC, RMSEA, but these are customizable using the `keepCols` and `dropCols` parameters.

Usage

```r
latexsummarytable(modellist, keepCols, dropCols, sortby, label = NULL, caption = NULL)
```

Arguments

- **modellist**: A list of models (as a `data.frame`) returned from the `extractModelSummaries` function.
- **keepCols**: A vector of character strings indicating which columns/variables to display in the summary. Only columns included in this list will be displayed (all others excluded). By default, `keepCols` is: c("Title", "LL", "Parameters", "AIC", "AICC", "BIC", "RMSEA", "CFI"). Example: c("Title", "LL", "AIC", "CFI")
- **dropCols**: A vector of character strings indicating which columns/variables to omit from the summary. Any column not included in this list will be displayed. By default, `dropCols` is NULL. Example: c("InputInstructions", "TLI")
- **sortby**: optional. Field name (as character string) by which to sort the table. Typically an information criterion (e.g., "AIC" or "BIC") is used to sort the table. Defaults to "AICC"
- **label**: optional. A character string specifying the label for the LaTeX table, which can be used for referencing the table.
- **caption**: optional. A character string specifying the caption for the LaTeX table.

Value

A LaTeX-formatted table summarizing the `modellist` is returned (created by `xtable`).

Note

You must choose between `keepCols` and `dropCols` because it is not sensible to use these together to include and exclude columns. The function will error if you include both parameters.

Author(s)

Michael Hallquist
lookupTech1Parameter

**See Also**

extractModelSummaries, HTMLSummaryTable, showSummaryTable, Sweave

**Examples**

```r
# make me!!!
```

---

**lookupTech1Parameter** *Lookup the matrix element for a give parameter number*

**Description**

The `lookupTech1Parameter` function identifies the position in the Mplus model matrices corresponding to a given parameter defined in the TECHNICAL 1 PARAMETER SPECIFICATION OUTPUT. The goal of this function is to aid in identifying problematic parameters often printed in the warnings and errors section of Mplus output.

**Usage**

```r
lookupTech1Parameter(tech1output, paramNumber)
```

**Arguments**

- `tech1output` The object corresponding to the TECH1 parameter specification from readModels.
- `paramNumber` The parameter number to lookup

**Value**

A `data.frame` containing the row(s) and column(s) of TECH1 parameter specification matching the requested `paramNumber`.

**Author(s)**

Michael Hallquist

**See Also**

readModels

**Examples**

```r
## Not run:
models <- readModels("test1.out")
param <- lookupTech1Parameter(models$tech1, 16)

## End(Not run)
```
mplus.traceplot

Plot the samples for each MCMC chain as a function of iterations

Description
Displays a traceplot of the MCMC draws from the poster distribution of each parameter estimate for a Bayesian Mplus model. This function requires that 1) PLOT: TYPE=PLOT2; be included in the Mplus input file, 2) a gh5 file be present corresponding to the Mplus output file (and containing a bayesian_data section), and 3) that the rhdf5 package be installed to allow the gh5 file to be imported.

Usage
mplus.traceplot(mplus.model, rows = 4, cols = 4, parameters_only = TRUE)

Arguments
mplus.model An Mplus model extracted by the readModels function.
rows Number of rows to display per plot.
cols Optional. Number of columns to display per plot.
parameters_only Optional. If TRUE, only the unstandardized parameter estimates from the MCMC draws will be displayed (as opposed to standardized estimates, r-square estimates, etc.). The unstandardized estimates all begin with "Parameter" in the Mplus gh5 output.

Details
A multi-panel plot is drawn to the screen and the user is prompted to display the next plot if more than rows x columns estimates are in the model.

Value
No value is returned by this function. Called for the side effect of displaying an MCMC chains traceplot.

Note
Trace and density plots can also be obtained using the coda package and the bparameters element of the mplus.model object. This requires that the posterior draws be saved using SAVEDATA: BPARAMETERS syntax. See example below.

Author(s)
Joseph Glass, Michael Hallquist
See Also

plot.mcmc

Examples

## Not run:
myModel <- readModels("BayesModel_WithGHMCMC.out")
mplus.traceplot(myModel, rows=2, cols=3)

#alternative using the coda package
library(coda)
plot(myModel$bparameters$valid_draw)

## End(Not run)

Description

The MplusAutomation package leverages the flexibility of the R language to automate latent variable model estimation and interpretation using Mplus, a powerful latent variable modeling program developed by Muthen and Muthen (www.statmodel.com). Specifically, MplusAutomation provides routines for creating related groups of models, running batches of models, and extracting and tabulating model parameters and fit statistics.

Details

The MplusAutomation package has four primary purposes:

1. To automatically run groups/batches of models.
2. To provide routines to extract model fit statistics, parameter estimates, and raw data from Mplus output files.
3. To facilitate comparisons among models
4. To provide a template language that allows for the creation of related input files.

The core routine for running batches of models is runModels, with an easy-to-use GUI wrapper, runModels.Interactive.

To extract model summary statistics from one or more output files, see extractModelSummaries, which returns a data.frame of fit statistics for models located within a directory. Model fit results can be summarized in tabular form (for comparing among models) using showSummaryTable (displays table in separate window), HTMLSummaryTable (creates HTML file containing summary table), or LatexSummaryTable (returns a LaTeX-formatted table of summary statistics).

To extract raw data created by the SAVEDATA command (e.g., class membership probabilities or factor scores), see getSavedata_Data.

To extract unstandardized or standardized parameter estimates from a single output file, see extractModelParameters.
Summaries, parameters, modification indices, and SAVEDATA output can be extracted simultaneously using the `readModels` function, and this is the recommended way to extract output with this package.

Model fit and parameter comparisons between models can be obtained using `compareModels`.

To create a group of related models from a single template, see `createModels`. Please read the MplusAutomation vignette provided along with the package (and on the CRAN website) in order to understand the template language.

In addition to the major functions above, a function for converting an R data.frame for use with Mplus is provided: `prepareMplusData`. This converts the data.frame to a tab-delimited file and provides an Mplus syntax stub for variable names.

<table>
<thead>
<tr>
<th>Package</th>
<th>MplusAutomation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type</td>
<td>Package</td>
</tr>
<tr>
<td>Version</td>
<td>0.6-2</td>
</tr>
<tr>
<td>Date</td>
<td>2013-10-24</td>
</tr>
<tr>
<td>License</td>
<td>LGPL-3</td>
</tr>
<tr>
<td>LazyLoad</td>
<td>yes</td>
</tr>
</tbody>
</table>

**Author(s)**

Michael Hallquist <michael.hallquist@gmail.com>, Joshua F. Wiley <jwiley.psych@gmail.com>

Maintainer: Michael Hallquist <michael.hallquist@gmail.com>

**References**


**See Also**

See `runModels` for an example running a model.

```r
mplusModeler
```

**Create, run, and read Mplus models.**

**Description**

This is a convenience wrapper to automate many of the usual steps required to run an Mplus model. It relies in part on functions from the MplusAutomation package.

**Usage**

```r
mplusModeler(object, dataout, modelout, run = FALSE, check = FALSE, varwarnings = TRUE, ...)"
Arguments

- **object**
  - An object of class mplusObject

- **dataout**
  - the name of the file to output the data to for Mplus. If missing, defaults to modelout changing .inp to .dat.

- **modelout**
  - the name of the output file for the model. This is the file all the syntax is written to, which becomes the Mplus input file. It should end in .inp. If missing, defaults to dataout changing the extension to .inp.

- **run**
  - an integer indicating how many models should be run. Defaults to zero. If zero, the data and model input files are all created, but the model is not run. This can be useful for seeing how the function works and what setup is done. If one, a basic model is run. If greater than one, the model is bootstrapped with run replications as well as the basic model.

- **check**
  - logical whether the body of the Mplus syntax should be checked for missing semicolons using the parseMplus function. Defaults to FALSE.

- **varwarnings**
  - A logical whether warnings about variable length should be left, the default, or removed from the output file.

- **...**
  - additional arguments passed to the prepareMplusData function.

Details

Combined with functions from the MplusAutomation package, this function is designed to make it easy to fit Mplus models from R and to ease many of the usual frustrations with Mplus. For example, Mplus has very specific formats it accepts data in, but also very little data management facilities. Using R data management is easy. This function is designed to make using data from R in Mplus models easy. It is also common to want to fit many different models that are slight variants. This can be tedious in Mplus, but using R you can create one basic set of input, store it in a vector, and then just modify that (e.g., using regular expressions) and pass it to Mplus. You can even use loops or the *apply constructs to fit the same sort of model with little variants.

Value

An Mplus model object, with results. If run = 1, returns an invisible list of results from the run of the Mplus model (see readModels from the MplusAutomation package). If run = 0, the function returns a list with two elements, ‘model’ and ‘boot’ that are both NULL. if run >= 1, returns a list with two elements, ‘model’ and ‘boot’ containing the regular Mplus model output and the boot object, respectively. In all cases, the Mplus data file and input files are created.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

runModels and readModels
Examples

## Not run:
# minimal example of a model using builtin data, allowing R
# to automatically guess the correct variables to use
```
test <- mplusObject(MODEL = "mpg ON wt hp;
  wt WITH hp;", rdata = mtcars)
```
# estimate the model in Mplus and read results back into R
```
res <- mplusModeler(test, "mtcars.dat", modelout = "model1.inp", run = 1L)
```
# show summary
```
summary(res)
```
# remove files
```
unlink("mtcars.dat")
unlink("model1.inp")
unlink("model1.out")
```

## simple example of a model using builtin data
# demonstrates use with a few more sections
```
test2 <- mplusObject(
  TITLE = "test the MplusAutomation Package and mplusModeler wrapper;",
  MODEL = "
    mpg ON wt hp;
    wt WITH hp;",
  usevariables = c("mpg", "wt", "hp"),
  rdata = mtcars)
```
res2 <- mplusModeler(test2, "mtcars.dat", modelout = "model2.inp", run = 1L)
# remove files
```
unlink("mtcars.dat")
unlink("model2.inp")
unlink("model2.out")
```

## similar example using a robust estimator for standard errors
# and showing how an existing model can be easily updated and reused
```
test3 <- update(test2, ANALYSIS = ~ "ESTIMATOR = MLR;")
```
res3 <- mplusModeler(test3, "mtcars.dat", modelout = "model3.inp", run = 1L)
unlink("mtcars.dat")
unlink("model3.inp")
unlink("model3.out")

## now use the built in bootstrapping methods
# note that these work, even when Mplus will not bootstrap
# also note how categorical variables and weights are declared
# in particular, the usevariables for Mplus must be specified
# because more variables are included in the data than are in the
# model. Note the R usevariables includes all variables for both
# model and weights. The same is true for clustering.
```
test4 <- mplusObject(
  TITLE = "test MplusAutomation with bootstrapping;",
  MODEL = "
    mpg ON wt hp;
    wt WITH hp;",
  usevariables = c("mpg", "wt", "hp"),
  rdata = mtcars,
  estime = "MLR; ESTIMATOR = MLR;"
)
```
```
res4 <- mplusModeler(test4, "mtcars.dat", modelout = "model4.inp", run = 1L)
```
# remove files
```
unlink("mtcars.dat")
unlink("model4.inp")
unlink("model4.out")
```
mplusObject

Create an Mplus model object

Description

This is a function to create an Mplus model object in R. The object holds all the sections of an Mplus input file, plus some extra R ones. Once created, the model can be run using other functions such as mplusModeler or updated using methods defined for the update function.

Usage

mplusObject(TITLE = NULL, DATA = NULL, VARIABLE = NULL, DEFINE = NULL, ANALYSIS = NULL, MODEL = NULL, OUTPUT = NULL, SAVEDATA = NULL, PLOT = NULL, usevariables = NULL, rdata = NULL, autov = TRUE)

Arguments

TITLE A character string of the title for Mplus.
DATA A character string of the data section for Mplus (note, do not define the filename as this is generated automatically)
VARIABLE A character string of the variable section for Mplus (note, do not define the variable names from the dataset as this is generated automatically)
DEFINE A character string of the define section for Mplus (optional)
ANALYSIS A character string of the analysis section for Mplus (optional)
MODEL: A character string of the model section for Mplus (optional, although typically you want to define a model)
OUTPUT: A character string of the output section for Mplus (optional)
SAVEDATA: A character string of the savedata section for Mplus (optional)
PLOT: A character string of the plot section for Mplus (optional)
usevariables: A character vector of the variables from the R dataset to use in the model.
rdata: An R dataset to be used for the model.
auto: A logical (defaults to TRUE) argument indicating whether R should attempt to guess the correct variables to use from the R dataset, if usevariables is left NULL.

Details
Mplus model objects allow a base model to be defined, and then flexibly update the data, change the precise model, etc. If a section does not vary between models, you can leave it the same. For example, suppose you are fitting a number of models, but in all cases, wish to use maximum likelihood estimator. “ANALYSIS: ESTIMATOR = ML;” and would like standardized output, “OUTPUT: STDYX;”. Rather than retype those in every model, they can be defined in one Mplus model object, and then that can simply be updated with different models, leaving the analysis and output sections untouched. This also means that if a reviewer comes back and asks for all analyses to be re-run say using the robust maximum likelihood estimator, all you have to do is change it in the model object once, and re run all your code.

Value
A list of class mplusObject with elements

TITLE: The title in Mplus (if defined)
DATA: The data section in Mplus (if defined)
VARIABLE: The variable section in Mplus (if defined)
DEFINE: The define section in Mplus (if defined)
ANALYSIS: The analysis section in Mplus (if defined)
MODEL: The model section in Mplus (if defined)
OUTPUT: The output section in Mplus (if defined)
SAVEDATA: The savedata section in Mplus (if defined)
PLOT: The plot section in Mplus (if defined)
results: NULL by default, but can be later updated to include the results from the model run.
usevariables: A character vector of the variables from the R data set to be used.
rdata: The R data set to use for the model.

Author(s)
Joshua F. Wiley <jwiley.psych@gmail.com>
mplusRcov

See Also
mplusModeler

Examples

```r
example1 <- mplusObject(MODEL = "mpg ON wt;",
                         usevariables = c("mpg", "hp"), rdata = mtcars)
str(example1)
rm(example1)

# R figures out the variables automagically, with a message
example2 <- mplusObject(MODEL = "mpg ON wt;",
                         rdata = mtcars, autov = TRUE)
str(example2)
rm(example2)

# R warns if the first 8 characters of a (used) variable name are not unique
# as they will be indistinguishable in the Mplus output
example1 <- mplusObject(MODEL = "basename_01 ON basename_02;",
                         rdata = data.frame(basename_01 = 1:5, basename_02 = 5:1),
                         autov = TRUE)
rm(example1)
```

mplusRcov

Create Mplus code for various residual covariance structures.

Description

This function makes it easy to write the Mplus syntax for various residual covariance structure.

Usage

```r
mplusRcov(x, type = c("homogenous", "heterogenous", "cs", "toeplitz", "ar", "un"), r = "rho", e = "e", collapse = FALSE)
```

Arguments

- `x` input character vector of variable names, ordered by time
- `type` A character string indicating the type of residual covariance structure to be used. Defaults to ‘homogenous’. Current options include ‘homogenous’, ‘heterogenous’, ‘cs’ for compound symmetric, ‘toeplitz’ for banded toeplitz, ‘ar’ for autoregressive, and ‘un’ for unstructured.
- `r` a character vector of the base label to name covariance parameters. Defaults to ‘rho’.
- `e` a character vector of the error variance of the variable. Used to create constraints on the covariance parameters. Defaults to ‘e’.
- `collapse` whether to collapse the covariance code using ‘PWITH’. Note that at the time of writing, Mplus does not allow more than 80 characters per row. Defaults to FALSE.
Details

The homogenous residual covariance structure estimates one parameter: the residual variance, $\sigma^2_e$. The residual variance is assumed to be identical for all variables and all covariances are assumed to be zero. The structure is represented in this table.

\[
\begin{array}{cccccc}
t_1 & t_2 & t_3 & \ldots & t_n \\
t_1 & \sigma^2_e & & & \\
t_2 & 0 & \sigma^2_e & & \\
t_3 & 0 & 0 & \sigma^2_e & \\
\vdots & \ddots & \ddots & \ddots & \ddots \\
t_n & 0 & 0 & 0 & \ldots & \sigma^2_e \\
\end{array}
\]

The heterogenous residual covariance structure estimates $n$ parameters, where $n$ is the number of variables. A unique residual variance is estimated for every variable. All covariances are assumed to be zero. The structure is represented in this table.

\[
\begin{array}{cccccc}
t_1 & t_2 & t_3 & \ldots & t_n \\
t_1 & \sigma^2_{e1} & & & \\
t_2 & 0 & \sigma^2_{e2} & & \\
t_3 & 0 & 0 & \sigma^2_{e3} & \\
\vdots & \ddots & \ddots & \ddots & \ddots \\
t_n & 0 & 0 & 0 & \ldots & \sigma^2_{en} \\
\end{array}
\]

The compound symmetric residual covariance structure estimates two parameters: one for the residual variance, $\sigma^2_e$, and one for the covariance. The residual variance is assumed to be identical for all variables and all covariances are assumed to be identical. The structure is represented in this table.

\[
\begin{array}{cccccc}
t_1 & t_2 & t_3 & \ldots & t_n \\
t_1 & \sigma^2_e & & & \\
t_2 & \rho & \sigma^2_e & & \\
t_3 & \rho & \rho & \sigma^2_e & \\
\vdots & \ddots & \ddots & \ddots & \ddots \\
t_n & \rho & \rho & \rho & \ldots & \sigma^2_e \\
\end{array}
\]

The toeplitz residual covariance structure estimates $n$ parameters, one for every band of the matrix. The residual variance, $\sigma^2_e$, is assumed to be identical for all variables. The covariances one step removed are all assumed identical. Likewise for all further bands. The structure is represented in this table.

\[
\begin{array}{cccccc}
t_1 & t_2 & t_3 & \ldots & t_n \\
t_1 & \sigma^2_e & & & \\
t_2 & \rho & \sigma^2_e & & \\
t_3 & \rho_2 & \rho & \sigma^2_e & \\
\vdots & \ddots & \ddots & \ddots & \ddots \\
\end{array}
\]
The autoregressive residual covariance structure has two parameters: the residual variance, $\sigma^2_e$ and the correlation between adjacent time points, $\rho$. The variances are constrained to be equal for all time points. A single correlation parameter is estimated. The $\rho$ is the correlation between adjacent time points such as 1 and 2 or 2 and 3. More distant relationships are assumed to have smaller correlations, decreasing exponentially. Thus between 1 and 3, the estimate is $\rho^2$. The structure is represented in this table.

<table>
<thead>
<tr>
<th>t1</th>
<th>t2</th>
<th>t3</th>
<th>...</th>
<th>tn</th>
</tr>
</thead>
<tbody>
<tr>
<td>t1</td>
<td>$\sigma^2_e$</td>
<td>$\rho$</td>
<td>$\sigma^2_e$</td>
<td>...</td>
</tr>
<tr>
<td>t2</td>
<td>$\rho$</td>
<td>$\sigma^2_e$</td>
<td>$\rho$</td>
<td>$\sigma^2_e$</td>
</tr>
<tr>
<td>t3</td>
<td>$\rho^2$</td>
<td>$\rho$</td>
<td>$\rho$</td>
<td>$\sigma^2_e$</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>tn</td>
<td>$\rho^{n-1}$</td>
<td>$\rho^{n-2}$</td>
<td>$\rho^{n-3}$</td>
<td>...</td>
</tr>
</tbody>
</table>

Because structural equation models generally model covariance structures, the autoregressive residual structure must be parameterized in terms of covariances. This is done in two parts. First, the function returns syntax to estimate all the pairwise covariances, labelling the parameters $\rho$, $\rho^2$, etc. so that they are constrained to be equal. Next, it returns the syntax for the necessary model constraints to constrain the different covariances, to decrease exponentially in their correlations. This is done via:

$$\rho^2 = \left( \frac{\rho}{\sigma^2_e} \right)^2 \sigma^2_e$$

and likewise for all later time points.

The unstructured residual covariance structure estimates $\frac{n(n+1)}{2}$ parameters. It is unstructured in that every variance and covariance is freely estimated with no constraints. However, in most cases, this results in an overparameterized model and is unestimable. The structure is represented in this table.

<table>
<thead>
<tr>
<th>t1</th>
<th>t2</th>
<th>t3</th>
<th>...</th>
<th>tn</th>
</tr>
</thead>
<tbody>
<tr>
<td>t1</td>
<td>$\sigma^2_{e1}$</td>
<td>$\rho_1$</td>
<td>$\sigma^2_{e2}$</td>
<td>...</td>
</tr>
<tr>
<td>t2</td>
<td>$\rho_1$</td>
<td>$\sigma^2_{e2}$</td>
<td>$\rho_2$</td>
<td>$\sigma^2_{e3}$</td>
</tr>
<tr>
<td>t3</td>
<td>$\rho_2$</td>
<td>$\rho_3$</td>
<td>$\rho_3$</td>
<td>$\sigma^2_{e3}$</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>tn</td>
<td>$\rho_5$</td>
<td>$\rho_6$</td>
<td>$\rho_7$</td>
<td>...</td>
</tr>
</tbody>
</table>

**Value**

A named character vector of class ‘MplusRstructure’ with four elements:

- **all**: A character string collapsing all other sections.
- **Variances**: A character string containing all of the variances.
- **Covariances**: A character string containing all of the covariances, properly labelled to allow constraints and the autoregressive residual covariance structure.
Constraints

A character string containing the ‘MODEL CONSTRAINT’ section and code needed to parameterize the residual covariance structure as autoregressive.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

Examples

# all five structures collapsing
mplusRcov(letters[1:4], "homogenous", "rho", "e", TRUE)
mplusRcov(letters[1:4], "heterogenous", "rho", "e", TRUE)
mplusRcov(letters[1:4], "cs", "rho", "e", TRUE)
mplusRcov(letters[1:4], "toeplitz", "rho", "e", TRUE)
mplusRcov(letters[1:4], "ar", "rho", "e", TRUE)
mplusRcov(letters[1:4], "un", "rho", "e", TRUE)

# all five structures without collapsing
# useful for long names or many variables
# where a line may cross 80 characters
mplusRcov(letters[1:4], "homogenous", "rho", "e", FALSE)
mplusRcov(letters[1:4], "heterogenous", "rho", "e", FALSE)
mplusRcov(letters[1:4], "cs", "rho", "e", FALSE)
mplusRcov(letters[1:4], "toeplitz", "rho", "e", FALSE)
mplusRcov(letters[1:4], "ar", "rho", "e", FALSE)
mplusRcov(letters[1:4], "un", "rho", "e", FALSE)

paramExtract

Extract parameters from a data frame of Mplus estimates

Description

This is a simple convenience function designed to facilitate looking at specific parameter types by easily return a subset of a data frame with those types only. It is designed to follow up the results returned from the `readModels` function.

Usage

```r
paramExtract(x, params = c("regression", "loading", "undirected", "expectation", "variability", "new"))
```

Arguments

`x`  
A data frame (specifically the type returned by `readModels`) containing parameters. Should be specific such as unstandardized and the data frame must have a column called ‘paramHeader’.
paramExtract

params

A character string indicating the types of parameters to be returned. Options currently include ‘regression’, ‘loading’, ‘undirected’, ‘expectation’, ‘variability’, and ‘new’ for new/additional parameters. Regressions include regression of one variable ON another. ‘loading’ include indicator variables (which are assumed caused by the underlying latent variable) and variables in latent growth models (BY or |). Undirected paths currently only include covariances, indicated by the WITH syntax in Mplus. Expectation paths are the unconditional or conditional expectations of variables. In other words those parameters related to the first moments. For independent variables, these are the means, \( E(X) \) and the conditional means or intercepts, \( E(X|f(\theta)) \) where \( f(\theta) \) is the model, some function of the parameters, \( \theta \). Finally ‘variability’ refers to both variances and residual variances, corresponding to the second moments. As with the expectations, variances are unconditional for variables that are not predicted or conditioned on any other variable in the model whereas residual variances are conditional on the model. Note that R uses fuzzy matching so that each of these can be called via shorthand, ‘r’, ‘l’, ‘u’, ‘e’, and ‘v’.

Value

A subset data frame with the parameters of interest.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

readModels

Examples

```r
## Not run:
test <- mplusObject(
  TITLE = "test the MplusAutomation Package and my Wrapper;",
  MODEL = "
    mpg ON wt hp;
    wt WITH hp;
    usevariables = c("mpg", "wt", "hp"),
    rdata = mtcars)

res <- mplusModeler(test, "mtcars.dat", modelout = "model1.inp", run = 1L)

# store just the unstandardized parameters in 'd'
d <- res$results$parameters$unstandardized
# extract just regression parameters
paramExtract(d, "regression")
# extract other types of parameters using shorthand
paramExtract(d, "u")
paramExtract(d, "e")
paramExtract(d, "v")
```

parseMplus

Check Mplus code for missing semicolons or too long lines.

Description

The function parses a character string containing Mplus code and checks that every non blank line ends in either a colon or a semicolon. In addition, it checks that every line is less than 90 characters, because Mplus ignores everything after 90 characters on a line which can be a source of enigmatic errors.

Usage

parseMplus(x, add = FALSE)

Arguments

x a character string containing Mplus code.
add logical indicating whether or not to add semicolons to lines that do not have them. Defaults to FALSE.

Details

The function is fairly basic at the moment. It works by simply removing blank space (spaces, tabs, etc.) and then if a line does not terminate in a colon or semicolon, it returns a note and the line number. Optionally, it can add semicolons to any lines missing them and return the input with added semicolons. To check for lines that are too long, all trailing (but not before) white space is removed, and then the number of characters is checked.

Value

a character vector containing the input text and optionally added semicolons.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

mplusModeler
plot.mplusObject  

Examples

```r
# sample input
test <- "
MODEL:
  mpg ON wt hp;
  wt WITH hp
"
# check and return
cat(parseMplus(test), fill=TRUE)
# add missing semicolons and return
cat(parseMplus(test, TRUE), fill=TRUE)
test <- "
MODEL:
  mpg cyl disp hp drat wt qsec vs am gear PWITH cyl disp hp drat wt qsec vs am gear carb;
"
cat(parseMplus(test))
```

---

plot.mplusObject  Plot coefficients for an mplusObject

Description

This is a method for plotting the coefficients of an mplusObject.

Usage

```r
## S3 method for class 'mplusObject'
plot(x, y, type = c("stdyx", "un", "std", "stdy"), ...)
```

Arguments

- `x`  An object of class mplusObject
- `y`  Not currently used
- `type`  A character vector indicating the type of coefficients to return. One of “un”, “std”, “stdy”, or “stdyx”. Defaults to “stdyx”.
- `...`  Additional arguments to pass on (not currently used)

Value

Nothing. Called for its side effect of plotting the coefficients.

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>
Examples

```r
## Not run:
#
# simple example of a model using builtin data
demonstrates use

#titles of models

##test the MplusAutomation Package;
##MplusAutomation Package

#test mplusObject,

##mpg on wt hp

##wt WITH hp;

##STANDARDIZED;

##usevariables = c("mpg", "wt", "hp",
##rdata = mtcars)

##example of the coef method

plot(res)

##remove files

##unlink("mtcars.dat")

##unlink("model1.inp")

##unlink("model1.log")

##End(Not run)
```

---

prepareMplusData

Create tab-delimited file and Mplus input syntax from R data.frame

Description

The `prepareMplusData` function converts an R data.frame object into a tab-delimited file (without header) to be used in an Mplus input file. The corresponding Mplus syntax, including the data file definition and variable names, is printed to the console or optionally to an input file.

Usage

```
prepareMplusData(df, filename, keepCols, dropCols, inpfile = FALSE,
interactive = TRUE, overwrite = TRUE)
```

Arguments

- `df`: The R data.frame to be prepared for Mplus
- `filename`: The path and filename for the tab-delimited data file for use with Mplus. Example: "C:/Mplusdata/data1.dat"
- `keepCols`: A character vector specifying the variable names within `df` to be output to `filename` or a numeric vector of the column indices to be output or a logical vector corresponding to the same.
**Value**

Invisibly returns a character vector of the Mplus input syntax. Primarily called for its side effect of creating Mplus data files and optionally input files.

**Author(s)**

Michael Hallquist

**Examples**

```r
## Not run:
library(foreign)

study5 <- read.spss("reanalysis-study-5-mt-fall-08.sav", to.data.frame=TRUE)
ASData5 <- subset(study5, select=c("ppnum", paste("as", 1:33, sep="")))

prepareMplusData(ASData5, "study5.dat")

# basic example
test01 <- prepareMplusData(mtcars, "test01.dat")

# see that syntax was stored
test01

# tests for keeping and dropping variables
prepareMplusData(mtcars, "test02.dat", keepCols = c("mpg", "hp"))
prepareMplusData(mtcars, "test03.dat", keepCols = c(1, 2))
prepareMplusData(mtcars, "test04.dat",
    keepCols = c(TRUE, FALSE, FALSE, TRUE, FALSE,
    FALSE, FALSE, FALSE, FALSE, FALSE, FALSE))

prepareMplusData(mtcars, "test05.dat", dropCols = c("mpg", "hp"))
prepareMplusData(mtcars, "test06.dat", dropCols = c(1, 2))
prepareMplusData(mtcars, "test07.dat",
```
```r
dropCols = c(TRUE, FALSE, FALSE, TRUE, FALSE,
FALSE, FALSE, FALSE, FALSE, FALSE, FALSE))
```

```r
# interactive (test08.dat)
prepareMplusData(mtcars, interactive=TRUE)

# write syntax to input file, not stdout
prepareMplusData(mtcars, "test09.dat", inpfile=TRUE)

# write syntax to alternate input file, not stdout
prepareMplusData(mtcars, "test10.dat", inpfile="test10alt.inp")

# should be error, no file
prepareMplusData(mtcars, interactive=FALSE)

# new warnings if it is going to overwrite files
# (the default to be consistent with prior behavior)
prepareMplusData(mtcars, "test10.dat")

# new warnings if it is going to overwrite files
# (the default to be consistent with prior behavior)
prepareMplusData(mtcars, "test11.dat", inpfile="test10alt.inp")

# new errors if files exist and overwrite=FALSE
prepareMplusData(mtcars, "test10.dat",
inpfile="test10alt.inp", overwrite=FALSE)

## End(Not run)
```

---

**print.MplusRstructure**  
*Print an Mplus Residual Structure object*

---

**Description**

This is a method for printing an Mplus Residual Structure object.

**Usage**

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

**Arguments**

- `x`  
  An object of class MplusRstructure

- `...`  
  Additional arguments to pass on (not currently used)

**Value**

NULL Called for its side effect of printing the object to the console
**readModels**

*Read Parameters, Summary Statistics, and Savedata from Mplus Output*

**Description**

Extracts the model parameters, summary statistics, and savedata from the one or more Mplus output files. This function is essentially a wrapper around `extractModelParameters`, `extractModelSummaries`, and `getSavedata_Data`, respectively. The goal is to have a single function that parses all (supported) aspects of Mplus output and to combine these into a list object, with one element per output file identified.

**Usage**

```r
readModels(target = getwd(), recursive = FALSE, filefilter)
```

**Arguments**

- **target**: the directory containing Mplus output files (.out) to parse OR the single output file to be parsed. May be a full path, relative path, or a filename within the working directory. Defaults to the current working directory. Example: "C:/Users/Michael/Mplus Runs"
- **recursive**: optional. If TRUE, parse all models nested in subdirectories within target. Defaults to FALSE.
- **filefilter**: a Perl regular expression (PCRE-compatible) specifying particular output files to be parsed within directory. See regex or [http://www.pcre.org/pcre.txt](http://www.pcre.org/pcre.txt) for details about regular expression syntax.

**Examples**

```r
# default 'show' uses printing
mplusRcov(c("a", "b", "c"), type = "ar")

# also if calling print explicitly
print(mplusRcov(c("a", "b", "c"), type = "ar"))

# to see all aspects of the raw/original object
str(mplusRcov(c("a", "b", "c"), type = "ar"))
```
Value

A list with one element per file. Each element is composed of up to three subelements: summaries, parameters, and savedata. If target is a single file, then the top-level elements will be the summaries, parameters, and savedata, not a list of files.

summarьерs  Summary statistics from extractModelSummaries, having structure as specified by that function
parameters  Model parameters from extractModelParameters, having structure as specified by that function
class_counts  Latent class counts and proportions for models that include a categorical latent variable
mod_indices  Model modification indices from extractModIndices, having structure as specified by that function
savedata_info  File information about SAVEDATA files related to this output
savedata  SAVEDATA file as an R dataframe, as described in getSavedata_Data
bparameters  an mcmc_list object containing the draws from the MCMC chains for a Bayesian model that uses the SAVEDATA: BPARAMETERS command
residuals  a list containing relevant information from OUTPUT: RESIDUALS
tech1  a list containing parameter specification and starting values from OUTPUT: TECH1
tech3  a list containing parameter covariance and correlation matrices from OUTPUT: TECH3
tech4  a list containing means, covariances, and correlations for latent variables from OUTPUT: TECH4
tech7  a list containing sample statistics for each latent class from OUTPUT: TECH7
tech9  a list containing warnings/errors from replication runs for MONTECARLO analyses from OUTPUT: TECH9
tech12  a list containing observed versus estimated sample statistics for TYPE=MIXTURE analyses from OUTPUT: TECH12
lcCondMeans  conditional latent class means and pairwise comparisons, obtained using auxiliary(e) syntax in latent class models
gh5  a list containing data from the gh5 (graphics) file corresponding to this output. (Requires rhdf5 package)

Author(s)

Michael Hallquist

See Also

extractModelSummaries, extractModelParameters, extractModIndices, getSavedata_Fileinfo, getSavedata_Data, getSavedata_Bparams
runModels

Run Mplus Models

Description

This function runs a group of Mplus models (.inp files) located within a single directory or nested within subdirectories.

Usage

runModels(directory = getwd(), recursive = FALSE, filefilter = NULL, showOutput = FALSE, replaceOutfile = "always", logFile = "Mplus Run Models.log", Mplus_command = "Mplus", killOnFail = TRUE)

Arguments

directory the directory containing Mplus input files (.inp) to run. Defaults to the current working directory. Example: "C:/Users/Michael/Mplus Runs"

recursive optional. If TRUE, run all models nested in subdirectories within directory. Defaults to FALSE.

filefilter a Perl regular expression (PCRE-compatible) specifying particular input files to be run within directory. See regex or http://www.pcre.org/pcre.txt for details about regular expression syntax.

showOutput optional. If TRUE, show estimation output (TECH8) in the R console. Note that if run within Rgui, output will display within R, but if run via Rterm, a separate window will appear during estimation.

replaceOutfile optional. Currently supports three settings: “always”, which runs all models, regardless of whether an output file for the model exists; “never”, which does not run any model that has an existing output file; and “modifiedDate”, which only runs a model if the modified date for the input file is more recent than the output file modified date (implying there have been updates to the model).

logFile optional. If non-null, specifies a file (and optionally, directory) that records the settings passed into the function and the models run (or skipped) during the run.

Mplus_command optional. N.B.: No need to pass this parameter for most users (has intelligent defaults). Allows the user to specify the name/path of the Mplus executable to be used for running models. This covers situations where Mplus is not in the system’s path, or where one wants to test different versions of the Mplus program.
**runModels_Interactive**

**killOnFail**

optional. Windows only for now. If TRUE, kill all processes named mplus.exe when runModels does not terminate normally. Defaults to TRUE.

**Value**

None. Function is used for its side effects (running models).

**Author(s)**

Michael Hallquist

**See Also**

runModels_Interactive

**Examples**

```r
## Not run:
runModels("C:/Users/Michael/Mplus Runs", recursive=TRUE, showOutput=TRUE,
         replaceOutfile="modifiedDate", logFile="MPlusRunLog.txt",
         Mplus_command="C:\\Users\\Michael\\Mplus Install\\Mplus.exe")

## End(Not run)
```

---

**runModels_Interactive  Run Mplus Models Using Graphical Interface**

**Description**

This function is provides a graphical user interface to the runModels function. It uses Tcl/Tk to display a window in which the user can specify parameters for runModels, including the directory for runs, recursing through subdirectories, displaying output on the console, and replacing existing outfiles.

**Usage**

```r
runModels_Interactive(directory = getwd(), recursive = "0",
                      showOutput = "1", replaceOutfile = "1", checkDate = "0",
                      logFile = "1")
```

**Arguments**

- **directory**
  
  optional. The starting directory that will display in the dialog window. Defaults to the current working directory.

- **recursive**
  
  optional. Whether the recursive checkbox should be checked when the window opens. “0” for FALSE, “1” for TRUE.

- **showOutput**
  
  optional. Whether the show output checkbox should be checked when the window opens. “0” for FALSE, “1” for TRUE.
showSummaryTable

replaceOutfile  optional. Whether the replace outfile checkbox should be checked when the window opens. “0” for FALSE, “1” for TRUE.

checkDate  optional. Whether the check modified date checkbox should be checked when the window opens. “0” for FALSE, “1” for TRUE.

logFile  optional. Whether the log file checkbox should be checked when the window opens. “0” for FALSE, “1” for TRUE.

Details

This function exists as a GUI wrapper for runModels and does not provide any distinct functionality.

Value

None. Function is used to display user interface for running models.

Author(s)

Michael Hallquist

See Also

runModels

Examples

# interactive, none

showSummaryTable  Display summary table of Mplus model statistics in separate window

Description

Displays a summary table of model fit statistics extracted using the extractModelSummaries function. This function relies on the showData function from the relimp package, which displays data in a Tk-based window. By default, the following summary statistics are included: Title, LL, Parameters, AIC, AICC, BIC, RMSEA_Estimate, but these are customizible using the keepCols and dropCols parameters.

Usage

showSummaryTable(modelList, keepCols, dropCols, sortBy, font = "Courier 9")
Arguments

modelList A list of models (as a data.frame) returned from the extractModelSummaries function.

keepCols A vector of character strings indicating which columns/variables to display in the summary. Only columns included in this list will be displayed (all others excluded). By default, keepCols is: c("Title", "LL", "Parameters", "AIC", "AICC", "BIC", "RMSEA")

Example: c("Title", "LL", "AIC")

dropCols A vector of character strings indicating which columns/variables to omit from the summary. Any column not included in this list will be displayed. By default, dropCols is NULL. Example: c("InputInstructions", "TLI")

sortBy Optional. Field name (as character string) by which to sort the table. Typically an information criterion (e.g., "AIC" or "BIC") is used to sort the table. Defaults to "AICC".

font Optional. The font to be used to display the summary table. Defaults to Courier 9.

Value

No value is returned by this function. It is solely used to display the summary table in a separate window.

Note

You must choose between keepCols and dropCols because it is not sensible to use these together to include and exclude columns. The function will error if you include both parameters.

Author(s)

Michael Hallquist

See Also

extractModelSummaries HTMLSummaryTable LatexSummaryTable

Examples

# make me!!!

---

**summary.mplusObject**

*Summarize an mplusObject*

**Description**

This is a method for summarizing an mplusObject.
Usage

```r
## S3 method for class 'mplusObject'
summary(object, verbose = FALSE, ...)
```

Arguments

- `object`: An object of class `mplusObject`
- `verbose`: Logical whether to print verbose output. Defaults to `FALSE`.
- `...`: Additional arguments to pass on (not currently used)

Value

`NULL` Called for its side effect of printing a model summary to the console

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

See Also

Other Mplus.Formatting: `coef.mplus.model, coef.mplusObject; extract, extract,mplus.model-method, extract,mplusObject-method, extract.mplus.model, extract.mplusObject:print.MplusRstructure`

Examples

```r
## Not run:
# simple example of a model using builtin data
# demonstrates use
test <- mplusObject(
  TITLE = "test the MplusAutomation Package;",
  MODEL = "
    mpg ON wt hp;
    wt WITH hp;",
  usevariables = c("mpg", "wt", "hp"),
  rdata = mtcars)
res <- mplusModeler(test, "mtcars.dat", modelout = "model1.inp", run = 1L)

# example of the summary method
summary(res)

# example of verbose output
summary(res, verbose=TRUE)

# remove files
unlink("mtcars.dat")
unlink("model1.inp")
unlink("model1.out")
unlink("Mplus Run Models.log")

## End(Not run)
```
testBParamCompoundConstraint

Test inequality-constrained hypothesis for two or more parameters based on iterations of MCMC chains

Description

Tests an inequality-constrained hypothesis (van de Schoot, Hoijtink, Hallquist, & Boelen, in press) based on draws from the posterior distribution of the model parameters, which provides information about the proportion of the distribution that is in agreement with a given hypothesis. This function is used for more complex hypotheses about three or more parameters, whereas testBParamConstraint tests a simple two-parameter hypothesis.

Usage

testBParamCompoundConstraint(bparams, test)

Arguments

bparams: An object containing draws from the posterior distribution (class mplus.model or mplus.bparameters). Obtained by SAVEDATA:BPARAMETERS in Mplus and getSavedata_Bparams or readModels in MplusAutomation.

test: The R code defining the parameter test of three or more parameters. Example: "(STAITOT.ON.CG > STAITOT.ON.UG) & (BDIM.ON.CG > BDIM.ON.UG)."

Details

This function accepts a bparameters object containing iterations of the MCMC chains (rows) for each model parameter (columns) and prints out the number and proportion of draws that are consistent with the requested hypothesis test.

The test argument is evaluated directly as R code, with the bparams object attached so that variable names are available directly in the environment. Because the goal is to evaluate the test for each draw from the posterior distribution, remember to use vector-based logic operators, not boolean operators. That is, stick to & or | for joining tests of parameters, rather than && or || since the latter will return a single TRUE/FALSE, which is irrelevant.

An example test in R logic would be "(STAITOT.ON.CG > STAITOT.ON.UG) & (BDIM.ON.CG > BDIM.ON.UG)."

Value

No value is returned by this function. Instead, two summary tables are printed to the screen containing the number and proportion of draws consistent with the hypothesis.

Author(s)

Michael Hallquist
**testBParamConstraint**

Test inequality-constrained hypothesis for two parameters based on iterations of MCMC chains

**Description**

Tests a simple inequality-constrained hypothesis (van de Schoot, Hoijtink, Hallquist, & Boelen, in press) based on draws from the posterior distribution of the model parameters, which provides information about the proportion of the distribution that is in agreement with a given hypothesis. This function is used for simple hypothesis for two parameters, whereas testBParamCompoundConstraint gives full access to multiple parameters and R's logic syntax. This function accepts a bparameters object containing iterations of the MCMC chains (rows) for each model parameter (columns) and prints out the number and proportion of draws that are consistent with the requested hypothesis test. The coef1, operator, and coef2 arguments are appended in sequence, so that the hypothesis test is constructed from left-to-right. e.g., `testBParamConstraint(bparamsDF, "MGM.TRT1", ">", "MGM.EX2")`.

**Usage**

```r
testBParamConstraint(bparams, coef1, operator, coef2)
```

**Arguments**

- **bparams**: An object containing draws from the posterior distribution (class `mplus.model` or `mplus.bparameters`). Obtained by `SAVEDATA:BPARAMETERS` in Mplus and `getSavedata_Bparams` or `readModels` in `MplusAutomation`.
- **coef1**: The name of the first parameter to be compared. Example: "MGM.TRT1"
- **operator**: A logical operator to compare the two parameters. Should be one of `>=`, `>`, `<`, or `<=`. Example: ">="
- **coef2**: The name of the first parameter to be compared. Example: "MGM.EX2"
update.mplusObject

Value

No value is returned by this function. Instead, two summary tables are printed to the screen containing the number and proportion of draws consistent with the hypothesis.

Author(s)

Michael Hallquist

See Also

testBParamCompoundConstraint

Examples

```r
## Not run:
# using bparameters directly
btest <- getSavedata_Bparams("model vbl_simpel_b.out")
testBParametersConstraint(btest, "STDYX_STATTOT.ON.CG", ">", "STDYX_STATTOT.ON.UCG")

# or using readModels
btest <- readModels("model vbl_simpel_b.out")
testBParametersConstraint(btest, "STDYX_STATTOT.ON.CG", ">", "STDYX_STATTOT.ON.UCG")
## End(Not run)
```

---

update.mplusObject  
Update an Mplus model object

Description

This is a method for updating an Mplus model object. It takes an Mplus model object as the first argument, and then optionally any sections to update. There are two ways to update a section using a formula interface. ~ "new stuff" will replace a given section with the new text. Alternately, you can add additional text using ~ + "additional stuff". Combined these let you replace or add to a section.

Usage

```r
## S3 method for class 'mplusObject'
update(object, ...)
```

Arguments

- **object**  
  An object of class mplusObject
- **...**  
  Additional arguments to pass on
Value

An (updated) Mplus model object

Author(s)

Joshua F. Wiley <jwiley.psych@gmail.com>

Examples

```r
example1 <- mplusObject(MODEL = "mpg ON wt;",
  usevariables = c("mpg", "hp"), rdata = mtcars)
x <- ~ "ESTIMATOR = ML;"
str(update(example1, rdata = iris))
str(update(example1, ANALYSIS = x))
str(update(example1, MODEL = ~ "wt ON hp;"))
str(update(example1, MODEL = ~ . + "wt ON hp;"))
str(update(example1, ANALYSIS = x, MODEL = ~ . + "wt ON hp;"))

# test to make sure . in Mplus code does not cause problems
str(update(example1, ANALYSIS = x, MODEL = ~ . + "wt ON hp*.5;"))
r(x)
```
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