

Package ‘tidyLPA’

October 14, 2022

Type Package

Title Easily Carry Out Latent Profile Analysis (LPA) Using Open-Source
or Commercial Software

Version 1.1.0

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Description An interface to the 'mclust' package to easily
carry out latent profile analysis ("LPA"). Provides functionality to
estimate commonly-specified models. Follows a tidy approach, in that
output is in the form of a data frame that can subsequently be
computed on. Also has functions to interface to the commercial 'MPlus'
software via the 'MplusAutomation' package.

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URL <https://data-edu.github.io/tidyLPA/>

BugReports <https://github.com/data-edu/tidyLPA/issues>

Depends R (>= 2.10)

Imports dplyr, ggplot2, gtable, grid, mclust, methods, mix,
MplusAutomation, tibble

Suggests knitr, lme4, missForest, parallel, pillar, rmarkdown,
testthat

VignetteBuilder knitr

Encoding UTF-8

LazyData true

RoxygenNote 7.1.2

NeedsCompilation no

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Repository CRAN

Date/Publication 2021-11-17 11:40:02 UTC

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AHP	<i>Select best model using analytic hierarchy process</i>
-----	---

Description

Integrates information from several fit indices, and selects the best model.

Usage

```
AHP(
  fitindices,
  relative_importance = c(AIC = 0.2323, AWE = 0.1129, BIC = 0.2525, CLC = 0.0922, KIC =
  0.3101)
)
```

Arguments

fitindices	A matrix or data.frame of fit indices, with colnames corresponding to the indices named in <code>relative_importance</code> .
------------	---

relative_importance

A named numeric vector. Names should correspond to columns in `fitindices`, and values represent the relative weight assigned to the corresponding fit index. The default value corresponds to the fit indices and weights assigned by Akogul and Erisoglu. To assign uniform weights (i.e., each index is weighted equally), assign an equal value to all.

Details

Many fit indices are available for model selection. Following the procedure developed by Akogul and Erisoglu (2017), this function integrates information from several fit indices, and selects the best model, using Saaty's (1990) Analytic Hierarchy Process (AHP). Conceptually, the process consists of the following steps:

1. For each fit index, calculate the amount of support provided for each model, relative to the other models.
2. From these comparisons, obtain a "priority vector" of the amount of support for each model.
3. Compute a weighted average of the priority vectors for all fit indeces, with weights based on a simulation study examining each fit index' ability to recover the correct number of clusters (Akogul & Erisoglu, 2016).
4. Select the model with the highest weighted average priority.

Value

Numeric.

Author(s)

Caspar J. van Lissa

Examples

```
iris[,1:4] %>%
  estimate_profiles(1:4) %>%
  get_fit() %>%
  AHP()
```

calc_lrt

Lo-Mendell-Rubin likelihood ratio test

Description

Implements the ad-hoc adjusted likelihood ratio test (LRT) described in Formula 15 of Lo, Mendell, & Rubin (2001), or LMR LRT.

Usage

```
calc_lrt(n, null_ll, null_param, null_classes, alt_ll, alt_param, alt_classes)
```

Arguments

n	Integer. Sample size
null_ll	Numeric. Log-likelihood of the null model.
null_param	Integer. Number of parameters of the null model.
null_classes	Integer. Number of classes of the null model.
alt_ll	Numeric. Log-likelihood of the alternative model.
alt_param	Integer. Number of parameters of the alternative model.
alt_classes	Integer. Number of classes of the alternative model.

Value

A numeric vector containing the likelihood ratio LR, the ad-hoc corrected LMR, degrees of freedom, and the LMR p-value.

References

Lo Y, Mendell NR, Rubin DB. Testing the number of components in a normal mixture. Biometrika. 2001;88(3):767–778. doi:10.1093/biomet/88.3.767

Examples

```
calc_lrt(150L, -741.02, 8, 1, -488.91, 13, 2)
```

compare_solutions *Compare latent profile models*

Description

Takes an object of class 'tidyLPA', containing multiple latent profile models with different number of classes or model specifications, and helps select the optimal number of classes and model specification.

Usage

```
compare_solutions(x, statistics = "BIC")
```

Arguments

x	An object of class 'tidyLPA'.
statistics	Character vector. Which statistics to examine for determining the optimal model. Defaults to 'BIC'.

Value

An object of class 'bestLPA' and 'list', containing a tibble of fits 'fits', a named vector 'best', indicating which model fit best according to each fit index, a numeric vector 'AHP' indicating the best model according to the AHP, an object 'plot' of class 'ggplot', and a numeric vector 'statistics' corresponding to argument of the same name.

Author(s)

Caspar J. van Lissa

Examples

```
iris_subset <- sample(nrow(iris), 20) # so examples execute quickly
results <- iris %>%
  subset(select = c("Sepal.Length", "Sepal.Width",
    "Petal.Length", "Petal.Width")) %>%
  estimate_profiles(1:3) %>%
  compare_solutions()
```

curry_mac

Simulated MAC data

Description

This simulated dataset, based on Curry et al., 2019, contains data on moral relevance and judgment across the seven domains of the Morality As Cooperation scale.

Usage

```
data(curry_mac)
```

Format

A data.frame with 1392 rows and 42 variables.

Details

sex	factor	Self-identified sex of participants, Male, Female, or Transgendered.
age_years	numeric	Participants' age in years.
KinshipR	numeric	Mean score of moral relevance, kinship subscale.
MutualismR	numeric	Mean score of moral relevance, mutualism subscale.
ExchangeR	numeric	Mean score of moral relevance, exchange subscale.
HawkR	numeric	Mean score of moral relevance, hawk subscale.
DoveR	numeric	Mean score of moral relevance, dove subscale.
DivisionR	numeric	Mean score of moral relevance, division subscale.
PossessionR	numeric	Mean score of moral relevance, possession subscale.
KinshipJ	numeric	Mean score of moral judgment, kinship subscale.

MutualismJ	numeric	Mean score of moral judgment, mutualism subscale.
ExchangeJ	numeric	Mean score of moral judgment, exchange subscale.
HawkJ	numeric	Mean score of moral judgment, hawk subscale.
DoveJ	numeric	Mean score of moral judgment, dove subscale.
DivisionJ	numeric	Mean score of moral judgment, division subscale.
PossessionJ	numeric	Mean score of moral judgment, possession subscale.

References

Curry, O. S., Jones Chesters, M., & Van Lissa, C. J. (2019). Mapping morality with a compass: Testing the theory of ‘morality-as-cooperation’ with a new questionnaire. *Journal of Research in Personality*, 78, 106–124. doi: [10.1016/j.jrp.2018.10.008](https://doi.org/10.1016/j.jrp.2018.10.008)

<i>empathy</i>	<i>Simulated empathy data</i>
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Description

This simulated dataset, based on Van Lissa et al., 2014, contains six annual assessments of adolescents’ mean scores on the empathic concern and perspective taking subscales of the Interpersonal Reactivity Index (Davis, 1983). The first measurement wave occurred when adolescents were, on average, 13 years old, and the last one when they were 18 years old.

Usage

```
data(empathy)
```

Format

A data frame with 467 rows and 13 variables.

Details

ec1	numeric	Mean score of empathic concern in wave 1
ec2	numeric	Mean score of empathic concern in wave 2
ec3	numeric	Mean score of empathic concern in wave 3
ec4	numeric	Mean score of empathic concern in wave 4
ec5	numeric	Mean score of empathic concern in wave 5
ec6	numeric	Mean score of empathic concern in wave 6
pt1	numeric	Mean score of perspective taking in wave 1
pt2	numeric	Mean score of perspective taking in wave 2
pt3	numeric	Mean score of perspective taking in wave 3
pt4	numeric	Mean score of perspective taking in wave 4
pt5	numeric	Mean score of perspective taking in wave 5
pt6	numeric	Mean score of perspective taking in wave 6
sex	factor	Adolescent sex; M = male, F = female.

References

Van Lissa, C. J., Hawk, S. T., Branje, S. J., Koot, H. M., Van Lier, P. A., & Meeus, W. H. (2014). Divergence Between Adolescent and Parental Perceptions of Conflict in Relationship to Adolescent Empathy Development. *Journal of Youth and Adolescence*, (Journal Article), 1–14. doi: [10.1007/s1096401401525](https://doi.org/10.1007/s1096401401525)

`estimate_profiles` *Estimate latent profiles*

Description

Estimates latent profiles (finite mixture models) using the open source package [mclust](#), or the commercial program Mplus (using the R-interface of [MplusAutomation](#)).

Usage

```
estimate_profiles(
  df,
  n_profiles,
  models = NULL,
  variances = "equal",
  covariances = "zero",
  package = "mclust",
  select_vars = NULL,
  ...
)
```

Arguments

<code>df</code>	data.frame of numeric data; continuous indicators are required for mixture modeling.
<code>n_profiles</code>	Integer vector of the number of profiles (or mixture components) to be estimated.
<code>models</code>	Integer vector. Set to <code>NULL</code> by default, and models are constructed from the <code>variances</code> and <code>covariances</code> arguments. See Details for the six models available in tidyLPA.
<code>variances</code>	Character vector. Specifies which variance components to estimate. Defaults to "equal" (constrain variances across profiles); the other option is "varying" (estimate variances freely across profiles). Each element of this vector refers to one of the models you wish to run.
<code>covariances</code>	Character vector. Specifies which covariance components to estimate. Defaults to "zero" (do not estimate covariances; this corresponds to an assumption of conditional independence of the indicators); other options are "equal" (estimate covariances between items, constrained across profiles), and "varying" (free covariances across profiles).

<code>package</code>	Character. Which package to use; 'mclust' or 'MplusAutomation' (requires Mplus to be installed). Default: 'mclust'.
<code>select_vars</code>	Character. Optional vector of variable names in <code>df</code> , to be used for model estimation. Defaults to <code>NULL</code> , which means all variables in <code>df</code> are used.
<code>...</code>	Additional arguments are passed to the estimating function; i.e., <code>Mclust</code> , or <code>mplusModeler</code> .

Details

Six models are currently available in tidyLPA, corresponding to the most common requirements. These are:

1. Equal variances and covariances fixed to 0
2. Varying variances and covariances fixed to 0
3. Equal variances and equal covariances
4. Varying variances and equal covariances (not able to be fit w/ mclust)
5. Equal variances and varying covariances (not able to be fit w/ mclust)
6. Varying variances and varying covariances

Two interfaces are available to estimate these models; specify their numbers in the `models` argument (e.g., `models = 1`, or `models = c(1, 2, 3)`), or specify the variances/covariances to be estimated (e.g.: `variances = c("equal", "varying")`, `covariances = c("zero", "equal")`). Note that when mclust is used, `models = c(1, 2, 3, 6)` are the only models available.

Value

A list of class 'tidyLPA'.

Examples

```
iris_sample <- iris[c(1:4, 51:54, 101:104), ] # to make example run more quickly

# Example 1:
iris_sample %>%
  subset(select = c("Sepal.Length", "Sepal.Width",
    "Petal.Length")) %>%
  estimate_profiles(3)

# Example 2:
iris %>%
  subset(select = c("Sepal.Length", "Sepal.Width",
    "Petal.Length")) %>%
  estimate_profiles(n_profiles = 1:4, models = 1:3)

# Example 3:
iris_sample %>%
  subset(select = c("Sepal.Length", "Sepal.Width",
```

```
"Petal.Length")) %>%  
estimate_profiles(n_profiles = 1:4, variances = c("equal", "varying"),  
covariances = c("zero", "zero"))
```

estimate_profiles_mclust

Estimate latent profiles using mclust

Description

Estimates latent profiles (finite mixture models) using the open source package [mclust](#).

Usage

```
estimate_profiles_mclust(df, n_profiles, model_numbers, select_vars, ...)
```

Arguments

<code>df</code>	data.frame with two or more columns with continuous variables
<code>n_profiles</code>	Numeric vector. The number of profiles (or mixture components) to be estimated. Each number in the vector corresponds to an analysis with that many mixture components.
<code>model_numbers</code>	Numeric vector. Numbers of the models to be estimated. See estimate_profiles for a description of the models available in tidyLPA.
<code>select_vars</code>	Character. Optional vector of variable names in <code>df</code> , to be used for model estimation. Defaults to <code>NULL</code> , which means all variables in <code>df</code> are used.
<code>...</code>	Parameters passed directly to Mclust . See the documentation of Mclust .

Value

An object of class 'tidyLPA' and 'list'

Author(s)

Casper J. van Lissa

estimate_profiles_mplus2*Estimate latent profiles using Mplus*

Description

Estimates latent profiles (finite mixture models) using the commercial program Mplus, through the R-interface of [MplusAutomation](#).

Usage

```
estimate_profiles_mplus2(
  df,
  n_profiles,
  model_numbers,
  select_vars,
  ...,
  keepfiles = FALSE
)
```

Arguments

<code>df</code>	data.frame with two or more columns with continuous variables
<code>n_profiles</code>	Numeric vector. The number of profiles (or mixture components) to be estimated. Each number in the vector corresponds to an analysis with that many mixture components.
<code>model_numbers</code>	Numeric vector. Numbers of the models to be estimated. See estimate_profiles for a description of the models available in tidyLPA.
<code>select_vars</code>	Character. Optional vector of variable names in <code>df</code> , to be used for model estimation. Defaults to <code>NULL</code> , which means all variables in <code>df</code> are used.
<code>...</code>	Parameters passed directly to mplusModeler . See the documentation of mplusModeler .
<code>keepfiles</code>	Logical. Whether to retain the files created by <code>mplusModeler</code> (e.g., for future reference, or to manually edit them).

Value

An object of class 'tidyLPA' and 'list'

Author(s)

Caspar J. van Lissa

get_data	<i>Get data from objects generated by tidyLPA</i>
----------	---

Description

Get data from objects generated by tidyLPA.

Usage

```
get_data(x, ...)

## S3 method for class 'tidyLPA'
get_data(x, ...)

## S3 method for class 'tidyProfile'
get_data(x, ...)
```

Arguments

- x An object generated by tidyLPA.
... further arguments to be passed to or from other methods. They are ignored in this function.

Value

If one model is fit, the data is returned in wide format as a tibble. If more than one model is fit, the data is returned in long form. See the examples.

Methods (by class)

- tidyLPA: Get data for a latent profile analysis with multiple numbers of classes and models, of class 'tidyLPA'.
- tidyProfile: Get data for a single latent profile analysis object, of class 'tidyProfile'.

Author(s)

Caspar J. van Lissa

Examples

```
## Not run:
if(interactive()){
  library(dplyr)
  # the data is returned in wide form
  results <- iris %>%
    select(Sepal.Length, Sepal.Width, Petal.Length, Petal.Width) %>%
    estimate_profiles(3)
  get_data(results)
```

```
# note that if more than one model is fit, the data is returned in long form
results1 <- iris %>%
  select(Sepal.Length, Sepal.Width, Petal.Length, Petal.Width) %>%
  estimate_profiles(c(3, 4))
get_data(results1)
}

## End(Not run)
```

get_estimates*Get estimates from objects generated by tidyLPA***Description**

Get estimates from objects generated by tidyLPA.

Usage

```
get_estimates(x, ...)

## S3 method for class 'tidyLPA'
get_estimates(x, ...)

## S3 method for class 'tidyProfile'
get_estimates(x, ...)
```

Arguments

- x An object generated by tidyLPA.
- ... further arguments to be passed to or from other methods. They are ignored in this function.

Value

A tibble.

Methods (by class)

- **tidyLPA:** Get estimates for a latent profile analysis with multiple numbers of classes and models, of class 'tidyLPA'.
- **tidyProfile:** Get estimates for a single latent profile analysis object, of class 'tidyProfile'.

Author(s)

Caspar J. van Lissa

Examples

```
## Not run:
if(interactive()){
  results <- iris %>%
    select(Sepal.Length, Sepal.Width, Petal.Length, Petal.Width) %>%
    estimate_profiles(3)
  get_estimates(results)
  get_estimates(results[[1]])
}

## End(Not run)
```

get_fit

Get fit indices from objects generated by tidyLPA

Description

Get fit indices from objects generated by tidyLPA.

Usage

```
get_fit(x, ...)

## S3 method for class 'tidyLPA'
get_fit(x, ...)

## S3 method for class 'tidyProfile'
get_fit(x, ...)
```

Arguments

- x An object generated by tidyLPA.
- ... further arguments to be passed to or from other methods. They are ignored in this function.

Value

A tibble. Learn more at https://data-edu.github.io/tidyLPA/articles/Introduction_to_tidyLPA.html#getting-fit-statistics

Methods (by class)

- **tidyLPA:** Get fit indices for a latent profile analysis with multiple numbers of classes and models, of class 'tidyLPA'.
- **tidyProfile:** Get fit indices for a single latent profile analysis object, of class 'tidyProfile' .

Author(s)

Casper J. van Lissa

Examples

```
## Not run:
if(interactive()){
  results <- iris %>%
    select(Sepal.Length, Sepal.Width, Petal.Length, Petal.Width) %>%
    estimate_profiles(3)
  get_fit(results)
  get_fit(results[[1]])
}

## End(Not run)
```

id_edu

Simulated identity data

Description

This simulated dataset, based on Crochetti et al., 2014, contains five annual assessments of adolescents' mean scores on the commitment, exploration (in depth), and reconsideration subscales of the Utrecht-Management of Identity Commitments Scale (Crocetti et al., 2008). The scores reported here reflect the educational identity subscales of this instrument. The first measurement wave occurred when adolescents were, on average, 14 years old, and the last one when they were 18 years old.

Usage

```
data(id_edu)
```

Format

A data frame with 443 rows and 16 variables.

Details

com1	numeric	Mean score of educational commitment in wave 1
exp1	numeric	Mean score of educational exploration in wave 1
rec1	numeric	Mean score of educational reconsideration in wave 1
com2	numeric	Mean score of educational commitment in wave 2
exp2	numeric	Mean score of educational exploration in wave 2
rec2	numeric	Mean score of educational reconsideration in wave 2
com3	numeric	Mean score of educational commitment in wave 3
exp3	numeric	Mean score of educational exploration in wave 3
rec3	numeric	Mean score of educational reconsideration in wave 3

com4	numeric	Mean score of educational commitment in wave 4
exp4	numeric	Mean score of educational exploration in wave 4
rec4	numeric	Mean score of educational reconsideration in wave 4
com5	numeric	Mean score of educational commitment in wave 5
exp5	numeric	Mean score of educational exploration in wave 5
rec5	numeric	Mean score of educational reconsideration in wave 5
sex	factor	Adolescent sex; M = male, F = female.

References

Crocetti, E., Klimstra, T. A., Hale, W. W., Koot, H. M., & Meeus, W. (2013). Impact of early adolescent externalizing problem behaviors on identity development in middle to late adolescence: A prospective 7-year longitudinal study. *Journal of Youth and Adolescence*, 42(11), 1745-1758.
doi: [10.1007/s1096401399246](https://doi.org/10.1007/s1096401399246)

pisaUSA15

student questionnaire data with four variables from the 2015 PISA for students in the United States

Description

student questionnaire data with four variables from the 2015 PISA for students in the United States

Usage

`pisaUSA15`

Format

Data frame with columns #'

broad_interest composite measure of students' self reported broad interest
enjoyment composite measure of students' self reported enjoyment
instrumental_mot composite measure of students' self reported instrumental motivation
self_efficacy composite measure of students' self reported self efficacy ...

Source

<http://www.oecd.org/pisa/data/>

plot_bivariate	<i>Create correlation plots for a mixture model</i>
----------------	---

Description

Creates a faceted plot of two-dimensional correlation plots and unidimensional density plots for an object of class 'tidyProfile'.

Usage

```
plot_bivariate(
  x,
  variables = NULL,
  sd = TRUE,
  cors = TRUE,
  rawdata = TRUE,
  bw = FALSE,
  alpha_range = c(0, 0.1),
  return_list = FALSE
)
```

Arguments

<code>x</code>	tidyProfile object to plot. A tidyProfile is one element of a tidyLPA analysis.
<code>variables</code>	Which variables to plot. If NULL, plots all variables that are present in all models.
<code>sd</code>	Logical. Whether to show the estimated standard deviations as lines emanating from the cluster centroid.
<code>cors</code>	Logical. Whether to show the estimated correlation (standardized covariance) as ellipses surrounding the cluster centroid.
<code>rawdata</code>	Logical. Whether to plot raw data, weighted by posterior class probability.
<code>bw</code>	Logical. Whether to make a black and white plot (for print) or a color plot. Defaults to FALSE, because these density plots are hard to read in black and white.
<code>alpha_range</code>	Numeric vector (0-1). Sets the transparency of geom_density and geom_point.
<code>return_list</code>	Logical. Whether to return a list of ggplot objects, or just the final plot. Defaults to FALSE.

Value

An object of class 'ggplot'.

Author(s)

Caspar J. van Lissa

Examples

```
# Example 1
iris_sample <- iris[c(1:10, 51:60, 101:110), ] # to make example run more quickly
## Not run:
iris_sample %>%
  subset(select = c("Sepal.Length", "Sepal.Width")) %>%
  estimate_profiles(n_profiles = 2, models = 1) %>%
  plot_bivariate()

## End(Not run)
# Example 2
## Not run:
mtcars %>%
  subset(select = c("wt", "qsec", "drat")) %>%
  poms() %>%
  estimate_profiles(3) %>%
  plot_bivariate()

## End(Not run)
```

plot_density

Create density plots for mixture models

Description

Creates a faceted plot of density plots for an object of class 'tidyLPA'. For each variable, a Total density plot will be shown, along with separate density plots for each latent class, where cases are weighted by the posterior probability of being assigned to that class.

Usage

```
plot_density(
  x,
  variables = NULL,
  bw = FALSE,
  conditional = FALSE,
  alpha = 0.2,
  facet_labels = NULL
)
```

Arguments

- | | |
|------------------|--|
| x | Object to plot. |
| variables | Which variables to plot. If NULL, plots all variables that are present in all models. |
| bw | Logical. Whether to make a black and white plot (for print) or a color plot. Defaults to FALSE, because these density plots are hard to read in black and white. |

<code>conditional</code>	Logical. Whether to show a conditional density plot (surface area is divided amongst the latent classes), or a classic density plot (surface area of the total density plot is equal to one, and is subdivided amongst the classes).
<code>alpha</code>	Numeric (0-1). Only used when bw and conditional are FALSE. Sets the transparency of geom_density, so that classes with a small number of cases remain visible.
<code>facet_labels</code>	Named character vector, the names of which should correspond to the facet labels one wishes to rename, and the values of which provide new names for these facets. For example, to rename variables, in the example with the 'iris' data below, one could specify: <code>facet_labels = c("Pet_leng" = "Petal length")</code> .

Value

An object of class 'ggplot'.

Author(s)

Caspar J. van Lissa

Examples

```
## Not run:
results <- iris %>%
  subset(select = c("Sepal.Length", "Sepal.Width",
    "Petal.Length", "Petal.Width")) %>%
  estimate_profiles(1:3)

## End(Not run)
## Not run:
plot_density(results, variables = "Petal.Length")

## End(Not run)
## Not run:
plot_density(results, bw = TRUE)

## End(Not run)
## Not run:
plot_density(results, bw = FALSE, conditional = TRUE)

## End(Not run)
## Not run:
plot_density(results[[2]], variables = "Petal.Length")

## End(Not run)
```

plot_profiles *Create latent profile plots*

Description

Creates a profile plot according to best practices, focusing on the visualization of classification uncertainty by showing:

1. Bars reflecting a confidence interval for the class centroids
2. Boxes reflecting the standard deviations within each class; a box encompasses +/- 64% of the observations in a normal distribution
3. Raw data, whose transparency is weighted by the posterior class probability, such that each datapoint is most clearly visible for the class it is most likely to be a member of.

Usage

```
plot_profiles(  
  x,  
  variables = NULL,  
  ci = 0.95,  
  sd = TRUE,  
  add_line = TRUE,  
  rawdata = TRUE,  
  bw = FALSE,  
  alpha_range = c(0, 0.1),  
  ...  
)  
  
## Default S3 method:  
plot_profiles(  
  x,  
  variables = NULL,  
  ci = 0.95,  
  sd = TRUE,  
  add_line = FALSE,  
  rawdata = TRUE,  
  bw = FALSE,  
  alpha_range = c(0, 0.1),  
  ...  
)
```

Arguments

x	An object containing the results of a mixture model analysis.
variables	A character vectors with the names of the variables to be plotted (optional).

<code>ci</code>	Numeric. What confidence interval should the errorbars span? Defaults to a 95% confidence interval. Set to NULL to remove errorbars.
<code>sd</code>	Logical. Whether to display a box encompassing +/- 1SD Defaults to TRUE.
<code>add_line</code>	Logical. Whether to display a line, connecting cluster centroids belonging to the same latent class. Defaults to TRUE. Note that the additional information conveyed by such a line is limited.
<code>rawdata</code>	Should raw data be plotted in the background? Setting this to TRUE might result in long plotting times.
<code>bw</code>	Logical. Should the plot be black and white (for print), or color?
<code>alpha_range</code>	The minimum and maximum values of alpha (transparency) for the raw data. Minimum should be 0; lower maximum values of alpha can help reduce overplotting.
<code>...</code>	Arguments passed to and from other functions.

Value

An object of class 'ggplot'.

Author(s)

Caspar J. van Lissa

Examples

```
# Example 1
iris_sample <- iris[c(1:10, 51:60, 101:110), ] # to make example run more quickly

iris_sample %>%
  subset(select = c("Sepal.Length", "Sepal.Width")) %>%
  estimate_profiles(n_profiles = 1:2, models = 1:2) %>%
  plot_profiles()

# Example 2

mtcars %>%
  subset(select = c("wt", "qsec", "drat")) %>%
  poms() %>%
  estimate_profiles(1:4) %>%
  plot_profiles(add_line = F)
```

poms	<i>Apply POMS-coding to data</i>
------	----------------------------------

Description

Takes in a data.frame, and applies POMS (proportion of maximum)-coding to the numeric columns.

Usage

```
poms(data)
```

Arguments

data	A data.frame.
------	---------------

Value

A data.frame.

Author(s)

Caspar J. van Lissa

Examples

```
data <- data.frame(a = c(1, 2, 2, 4, 1, 6),
                    b = c(6, 6, 3, 5, 3, 4),
                    c = c("a", "b", "b", "t", "f", "g"))
poms(data)
```

print.tidyLPA	<i>Print tidyLPA</i>
---------------	----------------------

Description

S3 method 'print' for class 'tidyLPA'.

Usage

```
## S3 method for class 'tidyLPA'
print(
  x,
  stats = c("AIC", "BIC", "Entropy", "prob_min", "prob_max", "n_min", "n_max",
            "BLRT_p"),
  digits = 2,
  na.print = "",
  ...
)
```

Arguments

- x An object of class 'tidyLPA'.
- stats Character vector. Statistics to be printed. Default: c("AIC", "BIC", "Entropy", "prob_min", "prob_max", "n_min", "n_max", "BLRT_p").
- digits minimal number of significant digits, see [print.default](#).
- na.print a character string which is used to indicate NA values in printed output, or NULL. See [print.default](#).
- ... further arguments to be passed to or from other methods. They are ignored in this function.

Author(s)

Caspar J. van Lissa

Examples

```
## Not run:
if(interactive()){
  iris %>%
    select(Sepal.Length, Sepal.Width, Petal.Length, Petal.Width) %>%
    estimate_profiles(3)
}

## End(Not run)
```

Description

S3 method 'print' for class 'tidyProfile'.

Usage

```
## S3 method for class 'tidyProfile'
print(x, digits = 2, na.print = "", ...)
```

Arguments

- x An object of class 'tidyProfile'.
- digits minimal number of significant digits, see [print.default](#).
- na.print a character string which is used to indicate NA values in printed output, or NULL. See [print.default](#).
- ... further arguments to be passed to or from other methods. They are ignored in this function.

Author(s)

Caspar J. van Lissa

Examples

```
## Not run:
if(interactive()){
  tmp <- iris %>%
    select(Sepal.Length, Sepal.Width, Petal.Length, Petal.Width) %>%
    estimate_profiles(3)
  tmp[[2]]
}

## End(Not run)
```

single_imputation *Apply single imputation to data*

Description

This function accommodates several methods for single imputation of data. Currently, the following methods are defined:

- "imputeData"Applies the mclust native imputation function [imputeData](#)
- "missForest"Applies non-parameteric, random-forest based data imputation using [missForest](#). Radom forests can accommodate any complex interactions and non-linear relations in the data. My simulation studies indicate that this method is preferable to mclust's imputeData (see examples).

Usage

```
single_imputation(x, method = "imputeData")
```

Arguments

x	A data.frame or matrix.
method	Character. Imputation method to apply, Default: 'imputeData'

Value

A data.frame

Author(s)

Caspar J. van Lissa

Examples

```

## Not run:
library(ggplot2)
library(missForest)
library(mclust)

dm <- 2
k <- 3
n <- 100
V <- 4

# Example of one simulation
class <- sample.int(k, n, replace = TRUE)
dat <- matrix(rnorm(n*V, mean = (rep(class, each = V)-1)*dm), nrow = n,
              ncol = V, byrow = TRUE)
results <- estimate_profiles(data.frame(dat), 1:5)
plot_profiles(results)
compare_solutions(results)

# Simulation for parametric data (i.e., all assumptions of latent profile
# analysis met)
simulation <- replicate(100, {
  class <- sample.int(k, n, replace = TRUE)
  dat <- matrix(rnorm(n*V, mean = (rep(class, each = V)-1)*dm), nrow = n,
                ncol = V, byrow = TRUE)

  d <- prodNA(dat)

  d_mf <- missForest(d)$ximp
  m_mf <- Mclust(d_mf, G = 3, "EEI")
  d_im <- imputeData(d, verbose = FALSE)
  m_im <- Mclust(d_im, G = 3, "EEI")

  class_tabl_mf <- sort(prop.table(table(class, m_mf$classification)),
                         decreasing = TRUE)[1:3]
  class_tabl_im <- sort(prop.table(table(class, m_im$classification)),
                         decreasing = TRUE)[1:3]
  c(sum(class_tabl_mf), sum(class_tabl_im))
})

# Performance on average
rowMeans(simulation)
# Performance SD
colSD(t(simulation))
# Plot shows slight advantage for missForest
plotdat <- data.frame(accuracy = as.vector(simulation), model =
  rep(c("mf", "im"), n))
ggplot(plotdat, aes(x = accuracy, colour = model))+geom_density()

# Simulation for real data (i.e., unknown whether assumptions are met)
simulation <- replicate(100, {
  d <- prodNA(iris[,1:4])
})

```

```

d_mf <- missForest(d)$ximp
m_mf <- Mclust(d_mf, G = 3, "EEI")
d_im <- imputeData(d, verbose = FALSE)
m_im <- Mclust(d_im, G = 3, "EEI")

class_tabl_mf <- sort(prop.table(table(iris$Species,
                                      m_mf$classification)), decreasing = TRUE)[1:3]
class_tabl_im <- sort(prop.table(table(iris$Species,
                                      m_im$classification)), decreasing = TRUE)[1:3]
c(sum(class_tabl_mf), sum(class_tabl_im))

# Performance on average
rowMeans(simulation)
# Performance SD
colSD(t(simulation))
# Plot shows slight advantage for missForest
plotdat <- data.frame(accuracy = as.vector(tmp),
                       model = rep(c("mf", "im"), n))
ggplot(plotdat, aes(x = accuracy, colour = model))+geom_density()

## End(Not run)

```

Description

Latent Profile Analysis (LPA) is a statistical modeling approach for estimating distinct profiles, or groups, of variables. In the social sciences and in educational research, these profiles could represent, for example, how different youth experience dimensions of being engaged (i.e., cognitively, behaviorally, and affectively) at the same time.

Details

tidyLPA provides the functionality to carry out LPA in R. In particular, tidyLPA provides functionality to specify different models that determine whether and how different parameters (i.e., means, variances, and covariances) are estimated and to specify (and compare solutions for) the number of profiles to estimate.

Description

tidyLPA suggests using the pipe operator, `%>%`, from the magrittr package (imported here from the dplyr package).

Arguments

lhs, rhs An object and a function to apply to it

Examples

```
# Instead of
subset(iris, select = c("Sepal.Length", "Sepal.Width", "Petal.Length", "Petal.Width"))
# you can write
iris %>%
  subset(select = c("Sepal.Length", "Sepal.Width", "Petal.Length", "Petal.Width"))
```

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