

Package ‘miclust’

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Type Package

Title Multiple Imputation in Cluster Analysis

Version 1.2.8

Description Implementation of a framework for cluster analysis with selection of the final number of clusters and an optional variable selection procedure. The package is designed to integrate the results of multiple imputed datasets while accounting for the uncertainty that the imputations introduce in the final results. In addition, the package can also be used for a cluster analysis of the complete cases of a single dataset. The package also includes specific methods to summarize and plot the results. The methods are described in Basagana et al. (2013) <[doi:10.1093/aje/kws289](https://doi.org/10.1093/aje/kws289)>.

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Contents

miclust-package	2
getdata	3
getvariablesfrequency	4

miclust	5
minhanes	9
plot.miclust	10
print.miclust	10
print.summary.miclust	11
summary.miclust	11
Index	14

miclust-package	<i>miclust-package: integrating multiple imputation with cluster analysis</i>
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Description

Cluster analysis with selection of the final number of clusters and an optional variable selection procedure. The package is designed to integrate the results of multiply imputed data sets while accounting for the uncertainty that the imputations introduce in the final results. See ‘Procedure’ below for further details on how the tool works.

Procedure

The tool consists of a two-step procedure. In the first step, the user provides the data to be analyzed. They can be a single data.frame or a list of data.frames including the raw data and the imputed data sets. In the latter case, getdata needs to be used first to get data prepared. In the second step, the miclust performs k-means clustering with selection of the final number of clusters and an optional (backward or forward) variable selection procedure. Specific summary and plot methods are provided to summarize and visualize the impact of the imputations on the results.

Authors

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References

The methodology used in the package is described in

Basagana X, Barrera-Gomez J, Benet M, Anto JM, Garcia-Aymerich J. A Framework for Multiple Imputation in Cluster Analysis. *American Journal of Epidemiology*. 2013;177(7):718-725.

getdata	<i>Creates a midata object.</i>
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Description

Creates an object of class `miData` to be clustered by the function `miclust`.

Usage

```
getdata(data)
```

Arguments

<code>data</code>	a list or <code>data.frame</code> object. If it is a data frame, it is assumed to contain just the raw data, with or without missing data. If it is a list of data frames, it is assumed that the first element contains the raw data and the remaining ones correspond to multiple imputed data sets. Since all variables are considered in the clustering procedure, no identifier variables must be present in the data. In addition, all variables need to be treated as numeric (i.e. categorical variables must be coded with numeric values). See Details below.
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Details

All variables in data frames in `impdata` are standardized by `getdata`, so categorical variables need to be coded with numeric values. Standardization is performed by centering all variables at the mean and then dividing by the standard deviation (or the difference between the maximum and the minimum values for binary variables). Such a standardization is applied only to the imputed data sets. The standardization of the raw data is internally applied by the `miclust` if needed (which is the case of analyzing just the raw data, i.e. complete cases analysis).

Value

An object of classes `c("list", "midata")` including the following items:

rawdata a data frame containing the raw data.

impdata if data is an object of class `list`, `impdata` is a list containing the standardized imputed data sets.

See Also

`miclust`.

Examples

```
### data minhanes:
data(minhanes)
class(minhanes)

### number of imputed datasets:
length(minhanes) - 1

### raw data with missing values:
summary(minhanes[[1]])

### first imputed data set:
minhanes[[2]]
summary(minhanes[[2]])

### data preparation for a complete case cluster analysis:
data1 <- getdata(minhanes[[1]])
class(data1)
names(data1)

### there are no imputed data sets:
data1$impdata

### data preparation for a multiple imputation cluster analysis:
data2 <- getdata(minhanes)
class(data2)
names(data2)

### number of imputed data sets:
length(data2$impdata)

### imputed data sets are standardized:
summary(data2$rawdata)
summary(data2$impdata[[1]])
```

getvariablesfrequency *Calculates the ranked selection frequency of the variables.*

Description

Creates a ranked selection frequency for all the variables that have been selected at least once along the analyzed imputed data sets. `getvariablesfrequency` can be useful for customizing the plot of these frequencies as it is shown in Examples below.

Usage

```
getvariablesfrequency(x, k = NULL)
```

Arguments

- x** an object of class `miclust` obtained with the function `miclust`.
- k** the number of clusters. The default value is the optimal number of clusters obtained by the function `miclust`.

Value

A list including the following items:

percfreq vector of the selection frequencies (percentage of times) of the variables in decreasing order.

varnames names of the variables.

See Also

`miclust`.

Examples

```
### see examples in miclust.
```

<code>miclust</code>	<i>Cluster analysis in multiple imputed data sets with optional variable selection.</i>
----------------------	---

Description

Performs cluster analysis in multiple imputed data sets with optional variable selection. Results can be summarized and visualized with the `summary` and `plot` methods.

Usage

```
miclust(
  data,
  method = "kmeans",
  search = c("none", "backward", "forward"),
  ks = 2:3,
  maxvars = NULL,
  usedimp = NULL,
  distance = c("manhattan", "euclidean"),
  centpos = c("means", "medians"),
  initcl = c("hc", "rand"),
  verbose = TRUE,
  seed = NULL
)
```

Arguments

<code>data</code>	object of class <code>mi</code> data obtained with the function <code>getdata</code> .
<code>method</code>	clustering method. Currently, only "kmeans" is accepted.
<code>search</code>	search algorithm for the selection variable procedure: "backward", "forward" or "none". If "none" (default), no variable selection is performed.
<code>ks</code>	the values of the explored number of clusters. Default is exploring 2 and 3 clusters.
<code>maxvars</code>	if <code>method = "forward"</code> , the maximum number of variables to be selected.
<code>usedimp</code>	numeric. Which imputed data sets must be included in the cluster analysis. If <code>NULL</code> (default), all available imputed data sets are included. If <code>usedimp</code> is numeric (or a numeric vector), its values indicate which imputed data sets are included.
<code>distance</code>	two metrics are allowed to compute distances: "manhattan" (default) and "euclidean".
<code>centpos</code>	position computation of the cluster centroid. If "means" (default) the position of the centroid is computed by the mean. If "medians", by the median.
<code>initcl</code>	starting values for the clustering algorithm. If "rand", they are randomly selected; if "hc", they are computed via hierarchical clustering. See Details below.
<code>verbose</code>	a logical value indicating output status messages. Default is <code>TRUE</code> .
<code>seed</code>	a number. Seed for reproducibility of results. Default is <code>NULL</code> (no seed).

Details

The optimal number of clusters and the final set of variables are selected according to CritCF. CritCF is defined as

$$CritCF = \left(\frac{2m}{2m+1} \cdot \frac{1}{1+W/B} \right)^{\frac{1+\log_2(k+1)}{1+\log_2(m+1)}},$$

where m is the number of variables, k is the number of clusters, and W and B are the within- and between-cluster inertias. Higher values of CritCF are preferred (Breaban, 2011). See References below for further details about the clustering algorithm.

For computational reasons, option "rand" is suggested instead of "hc" for high dimensional data.

Value

A list with class "miclust" including the following items:

clustering a list of lists containing the results of the clustering algorithm for each analyzed data set and for each analyzed number of clusters. Includes information about selected variables and the cluster vector.

completecasesperc if data contains a single data frame, percentage of complete cases in data.

data input data.

ks the values of the explored number of clusters.

usedimp indicator of the imputed data sets used.

kfin optimal number of clusters.

critcf if data contains a single data frame, critcf contains the optimal (maximum) value of CritCF (see Details) and the number of selected variables in the reduction procedure for each explored number of clusters. If data is a list, critcf contains the optimal value of CritCF for each imputed data set and for each explored value of the number of clusters.

numberofselectedvars number of selected variables.

selectedkdistribution if data is a list, frequency of selection of each analyzed number of clusters.

method input method.

search input search.

maxvars input maxvars.

distance input distance.

centpos input centpos.

selmetriccent an object of class kccaFamily needed by the specific summary method.

initcl input initcl.

References

- Basagana X, Barrera-Gomez J, Benet M, Anto JM, Garcia-Aymerich J. A framework for multiple imputation in cluster analysis. American Journal of Epidemiology. 2013;177(7):718-25.
- Breaban M, Luchian H. A unifying criterion for unsupervised clustering and feature selection. Pattern Recognition 2001;44(4):854-65.

See Also

[getdata](#) for data preparation before using miclust.

Examples

```
### data preparation:
minhanes1 <- getdata(data = minhanes)

#####
###
### Example 1:
###
### Multiple imputation clustering process with backward variable selection
###
#####

### using only the imputations 1 to 10 for the clustering process and exploring
### 2 vs. 3 clusters:
minhanes1clust <- miclust(data = minhanes1, search = "backward", ks = 2:3,
                        usedimp = 1:10, seed = 4321)

minhanes1clust
minhanes1clust$ksfin ### optimal number of clusters

### graphical summary:
plot(minhanes1clust)
```

```

### selection frequency of the variables for the optimal number of clusters:
y <- getvariablesfrequency(minhanes1clust)
y
plot(y$percfreq, type = "h", main = "", xlab = "Variable",
      ylab = "Percentage of times selected", xlim = 0.5 + c(0, length(y$varnames)),
      lwd = 15, col = "blue", xaxt = "n")
axis(1, at = 1:length(y$varnames), labels = y$varnames)

### default summary for the optimal number of clusters:
summary(minhanes1clust)

## summary forcing 3 clusters:
summary(minhanes1clust, k = 3)

#####
###
### Example 2:
###
### Same analysis but without variable selection
###
#####

minhanes2clust <- miclust(data = minhanes1, ks = 2:3, usedimp = 1:10, seed = 4321)
minhanes2clust
plot(minhanes2clust)
summary(minhanes2clust)

#####
###
### Example 3:
###
### Complete case clustering process with backward variable selection
###
#####

nhanes0 <- getdata(data = minhanes[[1]])
nhanes2clust <- miclust(data = nhanes0, search = "backward", ks = 2:3, seed = 4321)
nhanes2clust

summary(nhanes2clust)

### nothing to plot for a single data set analysis
# plot(nhanes2clust)

#####
###
### Example 4:
###
### Complete case clustering process without variable selection
###
#####

```



```
nhanes3clust <- miclust(data = nhanes0, ks = 2:3, seed = 4321)
nhanes3clust
summary(nhanes3clust)
```

minhanes

Multiple imputation for nhanes data.

Description

A list with 101 data sets. The first data set contains nhanes data from mice package. The remaining data sets were obtained by applying the multiple imputation function mice from package mice.

Usage

```
minhanes
```

Format

A list of 101 data.frames each of them with 25 observations of the following 4 variables:

age age group (1 = 20-39, 2 = 40-59, 3 = 60+). Treated as numerical.

bmi body mass index (kg/m²)

hyp hypertensive (1 = no, 2 = yes). Treated as numerical.

chl total serum cholesterol (mg/dL)

Source

<https://CRAN.R-project.org/package=mice>

Examples

```
data(minhanes)
### raw data:
minhanes[[1]]
summary(minhanes[[1]])

### number of imputed data sets:
length(minhanes) - 1

### first imputed data set:
minhanes[[2]]
summary(minhanes[[2]])
```

plot.miclust	<i>Shows a graphical representation of the results.</i>
--------------	---

Description

Creates a graphical representation of the results of [miclust](#).

Usage

```
## S3 method for class 'miclust'  
plot(x, k = NULL, ...)
```

Arguments

x	object of class miclust obtained with the function miclust .
k	number of clusters. The default value is the optimal number of clusters obtained by miclust .
...	further arguments for the plot function.

Value

a plot to visualize the clustering results.

See Also

[miclust](#), [summary.miclust](#).

print.miclust	<i>Prints the results.</i>
---------------	----------------------------

Description

Creates a summary print of the results of [miclust](#).

Usage

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

Arguments

x	object of class miclust obtained with the function miclust .
...	further arguments for the print method.

Value

prints a description of the clustering main results.

print.summary.miclust *Prints the summary of results.*

Description

Prints the summary of the results of [summary.miclust](#).

Usage

```
## S3 method for class 'summary.miclust'  
print(x, digits = 2, ...)
```

Arguments

x	object of class <code>summary.miclust</code> obtained with the method summary.miclust .
digits	digits for the print method. Default is 2.
...	further arguments for the print method.

Value

a print of the summary of the results generated by [summary.miclust](#).

See Also

[miclust](#), [summary.miclust](#).

`summary.miclust` *Summarizes the results.*

Description

Performs a within-cluster descriptive analysis of the variables after the clustering process performed by the function [miclust](#).

Usage

```
## S3 method for class 'miclust'  
summary(object, k = NULL, quantilevars = NULL, ...)
```

Arguments

<code>object</code>	object of class <code>miclust</code> obtained with the function <code>miclust</code> .
<code>k</code>	number of clusters. The default value is the optimal number of clusters obtained by <code>miclust</code> .
<code>quantilevars</code>	numeric. If a variable selection procedure was used, the cut-off percentile in order to decide the number of selected variables in the variable reduction procedure by decreasing order of presence along the imputations results. The default value is <code>quantilevars = 0.5</code> , i.e., the number of selected variables is the median number of selected variables along the imputations.
<code>...</code>	further arguments for the plot function.

Value

An object with classes `c("list", "summary.miclust")` including the following items:

allocationprobabilities if imputations were analyzed, descriptive summary of the probability of cluster assignment.

classmatrix if imputations were analyzed, the individual probabilities of cluster assignment.

cluster if imputations were analyzed, the final individual cluster assignment.

clusterssize if imputations were analyzed, size of the imputed cluster and between-imputations summary of the cluster size.

clustervector if a single data set (raw data set) has been clustered, a vector containing the individuals cluster assignments.

clustervectors if imputed data sets have been clustered, the individual cluster assignment in each imputation.

completecasesperc if a single data set (raw data set) has been clustered, the percentage of complete cases in the data set.

k number of clusters.

kappas if imputations were analyzed, the Cohen's kappa values after comparing the cluster vector in the first imputation with the cluster vector in each of the remaining imputations.

kappadistribution a summary of kappas.

m number of imputations used in the descriptive analysis which is the total number of imputations provided.

quantilevars if variable selection was performed, the input value of `quantilevars`.

search search algorithm for the selection variable procedure.

selectedvariables if variable selection was performed, the selected variables obtained considering `quantilevars`.

selectedvarspresence if imputations were analyzed and variable selection was performed, the presence of the selected variables along imputations.

summarybycluster within-cluster descriptive analysis of the selected variables.

usedimp indicator of imputations used in the clustering procedure.

summary.miclust

13

See Also

[miclust](#), [plot.miclust](#).

Examples

```
### see examples in miclust.
```

Index

* datasets

minhanes, [9](#)

getdata, [3](#), [6](#), [7](#)

getvariablesfrequency, [4](#)

miclust, [3](#), [5](#), [5](#), [10–13](#)

miclust-package, [2](#)

minhanes, [9](#)

plot.miclust, [10](#), [13](#)

print.miclust, [10](#)

print.summary.miclust, [11](#)

summary.miclust, [10](#), [11](#), [11](#)