Package ‘fsdaR’

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**Title**  Robust Data Analysis Through Monitoring and Dynamic Visualization

**Version**  0.9-0

**VersionNote**  Released 0.8-1 on 2023-03-09 on CRAN

**Description**  Provides interface to the 'MATLAB' toolbox 'Flexible Statistical Data Analysis (FSDA)' which is comprehensive and computationally efficient software package for robust statistics in regression, multivariate and categorical data analysis. The current R version implements tools for regression: (forward search, S- and MM-estimation, least trimmed squares (LTS) and least median of squares (LMS)), for multivariate analysis (forward search, S- and MM-estimation), for cluster analysis and cluster-wise regression. The distinctive feature of our package is the possibility of monitoring the statistics of interest as a function of breakdown point, efficiency or subset size, depending on the estimator. This is accompanied by a rich set of graphical features, such as dynamic brushing, linking, particularly useful for exploratory data analysis.

**Depends**  R (>= 3.5.0)

**Imports**  rJava, methods, stats4, ggplot2

**Suggests**  robustbase, rrcov, MASS

**SystemRequirements**  (license-free) MATLAB Runtime (MCR) V 9.12, Java (>=8)

**LazyLoad**  yes

**LazyData**  yes

**License**  GPL (>= 3)

**URL**  https://github.com/UniprJRC/fsdaR

**BugReports**  https://github.com/UniprJRC/fsdaR/issues

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Description

There are 60 observations on a response $y$ with the values of three explanatory variables. The scatter plot matrix of the data shows $y$ increasing with each of $x1$, $x2$ and $x3$. The plot of residuals against fitted values shows no obvious pattern. However the FS finds that there are 6 masked outliers.

Usage

data(bank_data)

Format

A data frame with 1949 rows and 14 variables. The variables are as follows:

- $x1$: Personal loans
- $x2$: Financing and hire-purchase
- $x3$: Mortgages
- $x4$: Life insurance
- $x5$: Share amount
- $x6$: Bond account
- $x7$: Current account
- $x8$: Salary deposits
- $x9$: Debit cards
- $x10$: Credit cards
- $x11$: Telephone banking
- $x12$: Domestic direct debits
- $x13$: Money transfers
- $y$: Profit/loss

Source


References

Description

Takes as input the output of function `tclustICsol` (that is a structure containing the best relevant solutions) and produces the car-bike plot. This plot provides a concise summary of the best relevant solutions. This plot shows on the horizontal axis the value of $c$ and on the vertical axis the value of $k$. For each solution we draw a rectangle for the interval of values for which the solution is best and stable and a horizontal line which departs from the rectangle for the values of $c$ in which the solution is only stable. Finally, for the best value of $c$ associated to the solution, we show a circle with two numbers, the first number indicates the ranked solution among those which are not spurious and the second one the ranked number including the spurious solutions. This plot has been baptized 'car-bike', because the first best solutions (in general 2 or 3) are generally best and stable for a large number of values of $c$ and therefore will have large rectangles. In addition, these solutions are likely to be stable for additional values of $c$ and therefore are likely to have horizontal lines departing from the rectangles (from here the name ’cars’). Finally, local minor solutions (which are associated with particular values of $c$ and $k$) do not generally present rectangles or lines and are shown with circles (from here the name ’bikes’).

Usage

```r
carbikeplot(out, SpuriousSolutions = FALSE, trace = FALSE, ...)
```

Arguments

- **out**: An S3 object of class `tclusticsol.object`, (output of `tclustICsol`) containing the relevant solutions.
- **SpuriousSolutions**: Whether to include or not spurious solutions. By default spurious solutions are not included into the plot.
- **trace**: Whether to print intermediate results. Default is `trace=FALSE`.
- **...**: Potential further arguments passed to lower level functions.

Author(s)

FSDA team, <valentin.todorov@chello.at>

References

Examples

## Not run:

## Car-bike plot for the geyser data ==========================

```r
data(geyser2)
out <- tclustIC(geyser2, whichIC="MIXMIX", plot=FALSE, alpha=0.1)

## Find the best solutions using as Information criterion MIXMIX
print("Best solutions using MIXMIX")
outMIXMIX <- tclustICsol(out, whichIC="MIXMIX", plot=FALSE, NumberOfBestSolutions=6)
print(outMIXMIX$MIXMIXbs)
carbikeplot(outMIXMIX)

## Car-bike plot for the flea data ==========================

```r
data(flea)
Y <- as.matrix(flea[, 1:(ncol(flea)-1)]) # select only the numeric variablesownames(Y) <- 1:nrow(Y)
head(Y)
out <- tclustIC(Y, whichIC="CLACLA", plot=FALSE, alpha=0.1, nsamp=100)

## Find the best solutions using as Information criterion CLACLA
print("Best solutions using CLACLA")
outCLACLA <- tclustICsol(out, whichIC="CLACLA", plot=FALSE, NumberOfBestSolutions=66)
## Produce the car-bike plot
carbikeplot(outCLACLA)

## End(Not run)
```

corfwdplot

**Monitoring the correlations between consecutive distances or residuals**

Description

Provides a method for obtaining the maximum empirical efficiency (in case of MM estimates) or maximum empirical breakdownplot (in case of S estimates) or maximum subset size (in case of forward search), using various measures of correlation between the \( n \) Mahalanobis distances or residuals at adjacent values of efficiency, breakdown point or subset size.

Usage

corfwdplot(out, trace = FALSE, ...)

Arguments

out  An object of S3 class returned by one of the estimation functions with the monitoring option selected (monitoring=TRUE): fsreda.object, sregeda.object, mmregeda.object, fsmeda.object, smulteda.object or mmmulteda.object. This is a list containing the monitoring of minimum Mahalanobis distance in case of multivariate analysis or the monitoring of residuals in case of regression. The needed elements of out are

1. MAL: matrix containing the squared Mahalanobis distances monitored in each step of the forward search. Every row is associated with a unit (row of data matrix Y). This matrix can be created using one of the functions fsmult, smult or mmmult with the monitoring option selected, i.e. monitoring=TRUE.

2. RES: matrix containing the residuals monitored in each step of the forward search. Every row is associated with a unit (row of data matrix Y). This matrix can be created using the function fsreg with the monitoring option selected, i.e. monitoring=TRUE.

3. bdp: a vector containing breakdown point values that have been used, in case of S estimates.

4. eff: a vector containing efficiency values that have been used, in case of MM estimates.

trace  Whether to print intermediate results. Default is trace=FALSE.

...  potential further arguments passed to lower level functions.

Value

A ggplot plot object which can be printed on screen or to a file.

Author(s)

FSDA team, <valentin.todorov@chello.at>

Examples

## Not run:

data(hbk, package="robustbase")
(out <- fsmult(hbk[,1:3], monitoring=TRUE))
corfwdplot(out)

(out1 <- smult(hbk, monitoring=TRUE, trace=TRUE))
corfwdplot(out1)

(out2 <- mmmult(hbk[,1:3], monitoring=TRUE, trace=TRUE))
corfwdplot(out2)

(out3 <- fsreg(hbk[,1:3], hbk[,4], monitoring=TRUE, trace=TRUE, method="FS"))
corfwdplot(out3)

(out4 <- fsreg(hbk[,1:3], hbk[,4], monitoring=TRUE, trace=TRUE, method="S"))
covplot(out4)

(out5 <- fsreg(hbk[,1:3], hbk[,4], monitoring=TRUE, trace=TRUE, method="MM"))
corfwdplot(out5)

## End(Not run)

covplot Monitoring of the covariance matrix

Description

Plots the trajectories of the elements of the covariance (correlation) matrix monitored

Usage

covplot(
  out,
  xlim,
  ylim,
  xlab,
  ylab,
  main,
  lwd,
  lty,
  col,
  cex.lab,
  cex.axis,
  subsize,
  fg.thresh,
  fg.unit,
  fg.labstep,
  fg.lwd,
  fg.lty,
  fg.col,
  fg.mark,
  fg.cex,
  standard,
  fground,
  tag,
  datatooltip,
  trace = FALSE,
  ...
)
covplot

Arguments

out
An object of S3 class `fsmeda.object` returned by `fsmult` with monitoring=TRUE
- a list containing the monitoring of minimum Mahalanobis distance.
The needed elements of `out` are:
1. S2cov: matrix containing the monitoring of the elements of the covariance
   matrix in each step of the forward search:
2. Un: matrix containing the order of entry of each unit (necessary if data-
tooltip or databrush is selected).
3. X: The data matrix.

xlim
Controls the x scale in the plot. `xlim` is a vector with two elements controlling
minimum and maximum on the x-axis. Default is to use automatic scale.

ylim
Controls the y scale in the plot. `ylim` is a vector with two elements controlling
minimum and maximum on the y-axis. Default is to use automatic scale.

xlab
A title for the x axis

ylab
A title for the y axis

main
An overall title for the plot

lwd
The line width, a positive number, defaulting to 1

lty
The line type. Line types can either be specified as an integer (1=solid (default),
2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the char-
acter strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash".
The latter two are not supported by Matlab.

col
Colors to be used for the highlighted units

cex.lab
The magnification to be used for x and y labels relative to the current setting of
cex

cex.axis
The magnification to be used for axis annotation relative to the current setting
of cex

subsize
Numeric vector containing the subset size with length equal to the number of
columns of matrix of mahalanobis distances. The default value of `subsize` is
`(nrow(MAL) - ncol(MAL) + 1):nrow(MAL)`

fg.thresh
(automatic to `fg.unit`) numeric vector of length 1 or 2 which specifies the high-
lighted trajectories. If `length(fg.thresh) == 1` the highlighted trajectories
are those of units that throughout the search had at least once a mahalanobis
distance greater than `fg.thresh`. The default value is `fg.thresh=2.5`. If
`length(fg.thresh) == 2` the highlighted trajectories are those of units that
throughout the search had a mahalanobis distance at least once bigger than

fg.unit
(automatic to `fg.thresh`), vector containing the list of the units to be highlighted.
If `fg.unit` is supplied, `fg.thresh` is ignored.

fg.labstep
numeric vector which specifies the steps of the search where to put labels for the
highlighted trajectories (units). The default is to put the labels at the initial and
final steps of the search. `fg.labstep=''` means no label.

fg.lwd
The line width for the highlighted trajectories (units). Default is 1.
fg.lty
The line type for the highlighted trajectories (units). Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.

gf.col
colors to be used for the highlighted units.

gf.mark
Controls whether to plot highlighted trajectories as symbols. if gf.mark==TRUE each line is plotted using a different symbol else no marker is used (default).

gf.cex
Controls the font size of the labels of the trajectories in foreground. If gf.cex=0 no labels will be shown - equivalent to gf.labstop="".

standard
MATLAB-style arguments - appearance of the plot in terms of xlim, ylim, axes labels and their font size style, color of the lines, etc.

fground
MATLAB-style arguments - for the trajectories in foreground.
tag
Plot handle. String which identifies the handle of the plot which is about to be created. The default is tag=’pl_mmd’. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.

datatooltip
If datatooltip is not empty the user can use the mouse in order to have information about the unit selected, the step in which the unit enters the search and the associated label. If datatooltip is a list, it is possible to control the aspect of the data cursor (see MATLAB function datacursormode() for more details or see the examples below). The default options are DisplayStyle="Window" and SnapToDataVertex="on".

trace
Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

Value
none

Author(s)
FSDA team, <valentin.todorov@chello.at>

Examples

## Not run:
X <- iris[,1:4]
out <- fsmult(X, monitoring=TRUE)

## Produce monitoring covariances plot with all the default options
covplot(out)

## End(Not run)
diabetes

Diabetes data

Description
The diabetes dataset, introduced by Reaven and Miller (1979), consists of 145 observations (patients). For each patient three measurements are reported: plasma glucose response to oral glucose, plasma insulin response to oral glucose, degree of insulin resistance.

Usage
```
data("diabetes")
```

Format
A data frame with the following variables:

- **glucose** Area under plasma glucose curve after a three hour oral glucose tolerance test (OGTT).
- **insulin** Area under plasma insulin curve after a three hour oral glucose tolerance test (OGTT).
- **sspg** Steady state plasma glucose.
- **class** The type of diabetes: Normal, Overt, and Chemical.

Source

Examples
```
library(rrcov)
data(diabetes)
head(diabetes)
plot(CovMcd(diabetes[, 1:3]), which="pairs", col=diabetes$class)
```

emilia2001
Demographic data from the 341 municipalities in Emilia Romagna (an Italian region).

Description
A data set containing 28 demographic variables for 341 municipalities in Emilia Romagna (an Italian region).

Usage
```
data(emilia2001)
```
Format

A data frame with 341 rows and 28 variables. The variables are as follows:

- less10: population aged less than 10
- more75: population aged more than 75
- single: single-member families
- divorced: divorced
- widows: widows and widowers
- graduates: population aged more than 25 who are graduates
- no_education: of those aged over 6 having no education
- employed: activity rate
- unemployed: unemployment rate
- increase_popul: standardised natural increase in population
- migration: standardised change in population due to migration
- birth_92_94: average birth rate over 1992-94
- fecundity: three-year average birth rate amongst women of child-bearing age
- houses: occupied houses built since 1982
- houses_2WCs: occupied houses with 2 or more WCs
- houses_heating: occupied houses with fixed heating system
- TV: TV licence holders
- cars: number of cars for 100 inhabitants
- luxury_cars: luxury cars
- hotels: working in hotels and restaurants
- banking: working in banking and finance
- income: average declared income amongst those filing income tax returns
- income_tax_returns: inhabitants filing income tax returns
- factories: residents employed in factories and public services
- factories_more10: employees employed in factories with more than 10 employees
- factories_more50: employees employed in factories with more than 50 employees
- artisanal: artisanal enterprises
- entrepreneurs: entrepreneurial and skilled self-employed among those of working age

fishery

Fishery data.

Description

The fishery data consist of 677 transactions of a fishery product in Europe. For each transaction the Value in 1000 euro and the quantity in Tons are reported.

Usage

data(fishery)

Format

A data frame with 677 rows and 2 variables

flea

Flea

Description

Flea-beetle measurements

Usage

data(flea)

Format

A data frame with 74 rows and 7 variables: six explanatory and one response variable - species. The variables are as follows:

- tars1: width of the first joint of the first tarsus in microns (the sum of measurements for both tarsi)
- tars2: the same for the second joint
- head: the maximal width of the head between the external edges of the eyes in 0.01 mm
- ade1: the maximal width of the aedeagus in the fore-part in microns
- ade2: the front angle of the aedeagus (1 unit = 7.5 degrees)
- ade3: the aedeagus width from the side in microns
- species, which species is being examined - Concinna, Heptapotamica, Heikertingeri

References

Examples

data(flea)
head(flea)

Description

A data set on air pressure in the Alps and the boiling point of water (Weisberg, 1985). There are 17 observations on the boiling point of water at different pressures, obtained from measurements at a variety of elevations in the Alps. The purpose of the experiment was to allow prediction of pressure from boiling point, which is easily measured, and so to provide an estimate of altitude: the higher the altitude, the lower the pressure. The dataset is characterized by one clear outlier.

Usage

data(forbes)

Format

A data frame with 17 rows and 2 variables The variables are as follows:

- x: boiling point
- y: 100 x log(pressure)

References


Examples

data(forbes)
plot(y~x, data=forbes)
Description

An object of class `fsdalms.object` holds information about the result of a call to `fsreg`.

Value

The object itself is basically a list with the following components:

- `rew` If `rew=TRUE` all subsequent output refers to reweighted else no reweighting is done.
- `beta` p-by-1 vector containing the estimated regression parameters.
- `bs` p x 1 vector containing the units forming subset associated with bLMS (bLTS).
- `residuals` residuals.
- `scale` scale estimate of the residuals.
- `weights` Vector like y containing weights. The elements of this vector are 0 or 1. These weights identify the h observations which are used to compute the final LTS (LMS) estimate. \( \text{sum(weights)=h} \) if there is not a perfect fit otherwise \( \text{sum(weights)} \) can be greater than h.
- `h` The number of observations that have determined the LTS (LMS) estimator, i.e. the value of h.
- `outliers` vector containing the list of the units declared as outliers using confidence level specified in input scalar `conflev`.
- `conflev` confidence level which is used to declare outliers. Remark: `conflev` will be used to draw the horizontal lines (confidence bands) in the plots. Default value is 0.975.
- `singsub` Number of subsets without full rank. Notice that if this number is greater than 0.1*(number of subsamples) a warning is produced.
- `X` the data matrix X
- `y` the response vector y

The object has class "fsdalms".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~, data=hbk, method="LMS"))
class(out)
summary(out)
## End(Not run)
```
Description

An object of class \texttt{fsdalts.object} holds information about the result of a call to \texttt{fsreg}.

Value

The object itself is basically a \texttt{list} with the following components:

- \texttt{rew} If \texttt{rew=TRUE} all subsequent output refers to reweighted else no reweighting is done.
- \texttt{beta} \(p\)-by-1 vector containing the estimated regression parameters.
- \texttt{bs} \(p\times1\) vector containing the units forming subset associated with bLMS (bLTS).
- \texttt{residuals} residuals.
- \texttt{scale} scale estimate of the residuals.
- \texttt{weights} Vector like \(y\) containing weights. The elements of this vector are 0 or 1. These weights identify the \(h\) observations which are used to compute the final LTS (LMS) estimate. sum(weights)=\(h\) if there is not a perfect fit otherwise sum(weights) can be greater than \(h\).
- \texttt{h} The number of observations that have determined the LTS (LMS) estimator, i.e. the value of \(h\).
- \texttt{outliers} vector containing the list of the units declared as outliers using confidence level specified in input scalar \texttt{conflev}.
- \texttt{conflev} confidence level which is used to declare outliers. Remark: \texttt{conflev} will be used to draw the horizontal lines (confidence bands) in the plots. Default value is 0.975
- \texttt{singsub} Number of subsets with out full rank. Notice that if this number is greater than \(0.1\times(\text{number of subsamples})\) a warning is produced
- \texttt{X} the data matrix \(X\)
- \texttt{y} the response vector \(y\)

The object has class "\texttt{fsdalts}".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="LTS"))
class(out)
summary(out)

## End(Not run)
```
Description

An object of class `fsmeda.object` holds information about the result of a call to `fsmult` when called with parameter `monitoring=TRUE`.

Value

The object itself is basically a list with the following components:

MAL: `n x (n-init+1)` matrix containing the monitoring of Each row represents the distance Mahalanobis distance for the corresponding unit.

BB: `n x (n-init+1)` matrix containing the information about the units belonging to the subset at each step of the forward search. The first column contains the indexes of the units forming subset in the initial step and each further column - the indexes of the units forming the corresponding step. The last column contains the units forming subset in the final step (all units).

md: `n-by-1` vector containing the estimates of the robust Mahalanobis distances (in squared units). This vector contains the distances of each observation from the location of the data, relative to the scatter matrix cov.

mmd: `(n-init) x 3` matrix, which contains the monitoring of minimum MD or (m+1)th ordered MD at each step of the forward search.

- 1st column = fwd search index (from init to n-1)
- 2nd column = minimum MD
- 3rd column = (m+1)th-ordered MD

msr: `(n-init+1) x 3` matrix which contains the monitoring of maximum MD or m-th ordered MD at each step of the forward search.

- 1st column = fwd search index (from init to n)
- 2nd column = maximum MD
- 3rd column = mth-ordered MD

gap: `(n-init+1) x 3` matrix which contains the monitoring of the gap (difference between minMD outside subset and max inside).

- 1st column = fwd search index (from init to n)
- 2nd column = min MD - max MD
- 3rd column = (m+1)th-ordered MD - mth ordered distance

Loc: `(n-init+1) x (p+1)` matrix which contains the monitoring of the estimated means at each step of the fwd search.

S2cov: `(n-init+1) x (p*(p+1)/2+1)` matrix which contains the monitoring of the of the elements of the covariance matrix in each step of the forward search.
- 1st column = fwd search index (from init to n)
- 2nd column = monitoring of $S[1,1]$
- 3rd column = monitoring of $S[1,2]$
- ...
- last column = monitoring of $S[p,p]$

detS: $(n\text{-init+1}) \times 2$ matrix which contains the monitoring of the determinant of the covariance matrix in each step of the forward search.

Un: $(n\text{-init})\times 11$ matrix which contains the unit(s) included in the subset at each step of the fwd search. REMARK: in every step the new subset is compared with the old subset. Un contains the unit(s) present in the new subset but not in the old one. Un[1,2] for example contains the unit included in step init+1. Un[end,2] contains the units included in the final step of the search.

X: the data matrix X.
The object has class "fsmeda".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsmult(hbk[,1:3], monitoring=TRUE))
class(out)
summary(out)
## End(Not run)
```

---

**fsmmmmdrs**

Depends: 

**Performs random start monitoring of minimum Mahalanobis distance**

### Description

The trajectories originate from many different random initial subsets and provide information on the presence of groups in the data. Groups are investigated by monitoring the minimum Mahalanobis distance outside the forward search subset.

### Usage

```r
fsmmmmdrs(
  x,
  plot = FALSE,
  init,
  bsbsteps,
  nsimul = 200,
  nocheck = FALSE,
  numpool,
  cleanpool = FALSE,
  msg = FALSE,
  ```


```r
trace = FALSE,
...
```

**Arguments**

**x**
An n x p data matrix (n observations and p variables). Rows of x represent observations, and columns represent variables. Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

**plot**
Plots the random starts minimum Mahalanobis distance with 1 If plot=FALSE (default) or plot=0 no plot is produced. The scale (ylim) for the y axis is defined as follows:
- ylim[2] is the maximum between the values of mmd in steps \([n*0.2 n]\) and the final value of the 99 per cent envelope multiplied by 1.1.
- ylim[1] is the minimum between the values of mmd in steps \([n*0.2 n]\) and the 1 per cent envelope multiplied by 0.9.

Remark: the plot which is produced is very simple. In order to control a series of options in this plot (including the y scale) and in order to connect it dynamically to the other forward plots it is necessary to use function `mmdrsplot`.

**init**
Point where to start monitoring required diagnostics. If init is not specified it will be set equal to \((p+1)\).

**bsbsteps**
A vector which specifies for which steps of the forward search it is necessary to save the units forming subset for each random start. If bsbsteps = 0 for each random start we store the units forming subset in all steps. The default is store the units forming subset in all steps if n <= 500 else to store the units forming subset at step init and steps which are multiple of 100. For example, if n = 753 and init = 6, units forming subset are stored for m=init, 100, 200, 300, 400, 500 and 600.

Remark: The vector bsbsteps must contain numbers from init to n. if min(bsbsteps) < init a warning message will be issued.

**nsimul**
Number of random starts. Default value is nsimul=200.

**nocheck**
It controls whether to perform checks on matrix Y. If nocheck=TRUE, no check is performed.

**numpool**
If numpool > 1, the routine automatically checks if the Parallel Computing Toolbox is installed and distributes the random starts over numpool parallel processes. If numpool <= 1, the random starts are run sequentially. By default, numpool is set equal to the number of physical cores available in the CPU (this choice may be inconvenient if other applications are running concurrently). The same happens if the numpool value chosen by the user exceeds the available number of cores.

Remark: up to R2013b, there was a limitation on the maximum number of cores that could be addressed by the parallel processing toolbox (8 and, more recently, 12). From R2014a, it is possible to run a local cluster of more than 12 workers.
REMARK: Unless you adjust the cluster profile, the default maximum number of workers is the same as the number of computational (physical) cores on the machine.

REMARK: In modern computers the number of logical cores is larger than the number of physical cores. By default, MATLAB is not using all logical cores because, normally, hyper-threading is enabled and some cores are reserved to this feature.

REMARK: It is because of Remarks 3 that we have chosen as default value for numpool the number of physical cores rather than the number of logical ones. The user can increase the number of parallel pool workers allocated to the multiple start monitoring by:

- setting the NumWorkers option in the local cluster profile settings to the number of logical cores (Remark 2). To do so go on the menu Home|Parallel|Manage Cluster Profile and set the desired "Number of workers to start on your local machine".
- setting numpool to the desired number of workers

Therefore, "if a parallel pool is not already open*, UserOption numpool (if set) overwrites the number of workers set in the local/current profile. Similarly, the number of workers in the local/current profile overwrites default value of numpool obtained as feature('numCores') (i.e. the number of physical cores).

cleanpool

Set cleanpool cleanpool=TRUE if the parallel pool has to be cleaned after the execution of the random starts. Otherwise (default) cleanpool=FALSE. Clearly this option has an effect just if previous option numpool > 1.

msg

Level of output to sidplay. It controls whether to display or not messages about random start progress. More precisely, if previous option numpool > 1, then a progress bar is displayed, on the other hand a message will be displayed on the screen when 10

REMARK: in order to create the progress bar when nparpool > 1 the program writes on a temporary .txt file in the folder where the user is working. Therefore it is necessary to work in a folder where the user has write permission. If this is not the case and the user (say) is working without write permission in folder C:/Program Files/MATLAB the following message will appear on the screen:

*Error using ProgressBar (line 57) Do you have write permissions for C:/Program Files/MATLAB?*

trace

Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

Value

Returns an object of class fsmmmdrs.object.

Author(s)

FSDA team, <valentin.todorov@chello.at>
**References**


**Examples**

```r
## Not run:
data(hbk, package="robustbase")
out <- fsmmmdrs(hbk[,1:3])
class(out)
summary(out)
## End(Not run)
```

---

**fsmmmdrs.object**

**Description of fsmmmdrs.object Objects**

**Description**

An object of class `fsmmmdrs.object` holds information about the result of a call to `fsmmmdrs`.

**Value**

The object itself is basically a `list` with the following components:

- `mmdrs`: Minimum Mahalanobis distance, (n-init) by (nsimul+1) matrix which contains the monitoring of minimum Mahalanobis distance at each step of the forward search.
  - 1st column = fwd search index (from init to n-1)
  - 2nd column = minimum Mahalanobis distance for random start 1
  - 3rd column ...
  - nsimul+1 column minimum Mahalanobis distance for random start nsimul

- `BBrs`: Units belonging to the subset at the steps specified by input option bsbsteps. If bsbsteps=0 `BBrs` has size n-by-(n-init+1)-by-nsimul. In this case `BBrs[,j]` with j=1, 2, ..., nsimul has the following structure:
  - 1st row = has number 1 in correspondence of the steps in which unit 1 is included inside subset and a missing value for the other steps
  - ...
```
fsmult

Gives an automatic outlier detection procedure in multivariate analysis

Description

Gives an automatic outlier detection procedure in multivariate analysis and performs forward search in multivariate analysis with exploratory data

Usage

fsmult(
  x,
  bsb,
  monitoring = FALSE,
  crit = c("md", "biv", "uni"),
  rf = 0.95,
  init,
  plot = FALSE,
  bonflev,
  msg = TRUE,
  nocheck = FALSE,
  scaled = FALSE,
  trace = FALSE,
  ...
)

Examples

## Not run:
data(hbk, package="robustbase")
out <- fsmmmdrs(hbk[,1:3])
class(out)
summary(out)

## End(Not run)
Arguments

x
An n x p data matrix (n observations and p variables). Rows of x represent observations, and columns represent variables.
Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

bsb
List of units forming the initial subset or size of the initial subset. If monitoring=FALSE the default is to start the search with p+1 units, containing those observations which are not outlying on any scatterplot, found as the intersection of all points lying within a robust contour containing a specified portion of the data (Riani and Zani 1997) and inside the univariate boxplot.
Remark: if bsb is a vector, the option crit is ignored.

monitoring
Wheather to perform monitoring of Mahalanobis distances and other specific quantities

crit
If specified, the criterion to be used to initialize the search.

- If crit="md" the units which form initial subset are those which have the smallest m0 pseudo Mahalanobis distances computed using procedure unibiv() (bivariate robust ellipses).
- If crit="biv" sorting is done first in terms of times units fell outside robust bivariate ellipses and then in terms of pseudoMD. In other words, the units forming initial subset are chosen first among the set of those which never fell outside robust bivariate ellipses then among those which fell only once outside bivariate ellipses ... up to reach m0.
- If crit="uni" sorting is done first in terms of times units fell outside univariate boxplots and then in terms of pseudoMD. In other words, the units forming initial subset are chosen first among the set of those which never fell outside univariate boxplots then among those which fell only once outside univariate boxplots... up to reach m0.

Remark: as the user can see the starting point of the search is not going to affect at all the results of the analysis. The user can explore this point with his own datasets.
Remark: if crit="biv" the user can also supply in scalar rf (see below) the confidence level of the bivariate ellipses.

rf
Confidence level for bivariate ellipses. The default is 0.95. This option is useful only if crit='biv'.

init
Point where to start monitoring required diagnostics. Note that if a vector m0 is supplied, init >= length(m0). If init is not specified it will be set equal to floor(n*0.6).

plot
Plots the minimum Mahalanobis distance. If plot=FALSE (default) or plot=0 no plot is produced. If plot=TRUE the plot of minimum MD with envelopes based on n observations and the scatterplot matrix with the outliers highlighted is produced. If plot=2 the additional plots of envelope resuperimposition are produced. If plot is a list it may contain the following fields:

- ylim vector with two elements controlling minimum and maximum on the y axis. Default value is " (automatic scale)
• xlim vector with two elements controlling minimum and maximum on the x axis. Default value is "(automatic scale)
• resuper vector which specifies for which steps it is necessary to show the plots of resuperimposed envelopes if resuper is not supplied a plot of each step in which the envelope is resuperimposed is shown. Example: if resuper = c(85 87) plots of resuperimposed envelopes are shown at steps m=85 and m=87
• ncoord If ncoord=1 plots are shown in normal coordinates else (default) plots are shown in traditional mmd coordinates
• labeladd If labeladd=1, the outliers in the spm are labelled with the unit row index. The default value is labeladd="", i.e. no label is added
• nameY character vector containing the labels of the variables. As default value, the labels which are added are Y1,...,Yp.
• lwd controls line width of the curve which contains the monitoring of minimum Mahalanobis distance. Default is lwd=2.
• lwdenv Controls linewidth of the envelopes. Default is lwdenv=2.

bonflev Option that might be used to identify extreme outliers when the distribution of the data is strongly non normal. In these circumstances, the general signal detection rule based on consecutive exceedances cannot be used. In this case bonflev can be:
1. a scalar smaller than 1, which specifies the confidence level for a signal and a stopping rule based on the comparison of the minimum deletion residual with a Bonferroni bound. For example if bonflev=0.99 the procedure stops when the trajectory exceeds for the first time the 99 per cent bonferroni bound.
2. a scalar value greater than 1. In this case the procedure stops when the residual trajectory exceeds for the first time this value.

Default value is empty, which means to rely on general rules based on consecutive exceedances.

msg It controls whether to display or not messages on the screen. If msg=TRUE (default) messages about the progression of the search are displayed on the screen otherwise only error messages will be displayed.

nocheck It controls whether to perform checks on matrix Y. If nocheck=TRUE, no check is performed.

scaled Controls whether to monitor scaled Mahalanobis distances (only if monitoring=TRUE). If scaled=TRUE Mahalanobis distances monitored during the search are scaled using ratio of determinant. If scaled=2 Mahalanobis distances monitored during the search are scaled using asymptotic consistency factor. The default is scaled=FALSE, that is Mahalanobis distances are not scaled.

trace Whether to print intermediate results. Default is trace=FALSE.

Value

Depending on the input parameter monitoring, one of the following objects will be returned:
fsmult

1. fsmult.object
2. fsmeda.object

Author(s)
FSDA team, <valentin.todorov@chello.at>

References

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsmult(hbk[,1:3]))
class(out)
summary(out)

## Generate contaminated data (200,3)
n <- 200
p <- 3
set.seed(123456)
X <- matrix(rnorm(n*p), nrow=n)
Xcont <- X
out1 <- fsmult(Xcont, trace=TRUE) # no plots (plot defaults to FALSE)
names(out1)

(out1 <- fsmult(Xcont, trace=TRUE, plot=TRUE)) # identical to plot=1

## plot=1 - minimum MD with envelopes based on n observations
## and the scatterplot matrix with the outliers highlighted
(out1 <- fsmult(Xcont, trace=TRUE, plot=1))

## plot=2 - additional plots of envelope resuperimposition
(out1 <- fsmult(Xcont, trace=TRUE, plot=2))

## plots is a list: plots showing envelope superimposition in normal coordinates.
(out1 <- fsmult(Xcont, trace=TRUE, plot=list(ncoord=1)))

## Choosing an initial subset formed by the three observations with
## the smallest Mahalanobis Distance.
```
(out1 <- fsmult(Xcont, m0=5, crit="md", trace=TRUE))

## fsmult() with monitoring
(out2 <- fsmult(Xcont, monitoring=TRUE, trace=TRUE))
names(out2)

## Monitor the exceedances from m=200 without showing plots.
n <- 1000
p <- 10
Y <- matrix(rnorm(10000), ncol=10)
(out <- fsmult(Y, init=200))

## Forgery Swiss banknotes examples.
data(swissbanknotes)

## Monitor the exceedances of Minimum Mahalanobis Distance
(out1 <- fsmult(swissbanknotes[101:200,], plot=1))

## Control minimum and maximum on the x axis
(out1 <- fsmult(swissbanknotes[101:200,], plot=list(xlim=c(60,90))))

## Monitor the exceedances of Minimum Mahalanobis Distance using
## normal coordinates for mmd.
(out1 <- fsmult(swissbanknotes[101:200,], plot=list(ncoord=1)))

## End(Not run)

---

**Description**

An object of class `fsmult.object` holds information about the result of a call to `fsmult`.

**Value**

The object itself is basically a **list** with the following components:

- **outliers**: kx1 vector containing the list of the k units declared as outliers or NULL if the sample is homogeneous.
- **loc**: p-by-1 vector containing location of the data.
- **cov**: p-by-p robust estimate of covariance matrix.
- **md**: n-by-1 vector containing the estimates of the robust Mahalanobis distances (in squared units). This vector contains the distances of each observation from the location of the data, relative to the scatter matrix `cov`.
- **mmd**: (n-init)-by-2 matrix. 1st col is the forward search index; 2nd col is the value of minimum Mahalanobis Distance in each step of the fwd search.
fsr.object

Description of fsr Objects

Description

An object of class `fsr.object` holds information about the result of a call to `fsreg`.

Value

The object itself is basically a `list` with the following components:

- `beta` : p-by-1 vector containing the estimated regression parameters (in step n-k).
- `scale` : scalar containing the estimate of the scale (sigma).
- `residuals` : residuals.
- `fittedvalues` : fitted values.
- `outliers` : kx1 vector containing the list of the k units declared as outliers or NULL if the sample is homogeneous.
- `mdr` : (n-init) x 2 matrix 1st col = fwd search index, 2nd col = value of minimum deletion residual in each step of the fwd search.
Un is an \((n\text{-init}) \times 11\) matrix which contains the unit(s) included in the subset at each step of the fwd search. REMARK: in every step the new subset is compared with the old subset. Un contains the unit(s) present in the new subset but not in the old one. Un(1,2) for example contains the unit included in step init+1. Un(end,2) contains the units included in the final step of the search.

\text{nout} is a 2 \times 5 matrix containing the number of times mdr went out of particular quantiles. First row contains quantiles 1 99 99.9 99.99 99.999. Second row contains the frequency distribution.

\text{constr} This output is produced only if the search found at a certain step is a non singular matrix \(X\). In this case the search run in a constrained mode, that is including the units which produced a singular matrix in the last \(n\text{-constr}\) steps. out.constr is a vector which contains the list of units which produced a singular \(X\) matrix.

\(X\) is the data matrix \(X\)

\(y\) is the response vector \(y\)

The object has class "fsr".

**Examples**

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="FS"))
class(out)
summary(out)
## End(Not run)
```

---

**fsrbase**

**Description**

An automatic outlier detection procedure in linear regression

**Usage**

```r
fsrbase(x, ...)
```

**S3 method for class 'formula'**

```r
fsrbase(formula, data, subset, weights, na.action, 
        model = TRUE, x.ret = FALSE, y.ret = FALSE, 
        contrasts = NULL, offset, ...)
```

**Default S3 method:**

```r
fsrbase(x, y, bsb, intercept = TRUE, 
        monitoring = FALSE, control, trace = FALSE, 
        ...)
```
Arguments

formula  a formula of the form \( y \sim x_1 + x_2 + \ldots \).
data  data frame from which variables specified in formula are to be taken.
subset  an optional vector specifying a subset of observations to be used in the fitting process.
weights  an optional vector of weights to be used in the fitting process. **NOT USED YET.**
na.action  a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The “factory-fresh” default is na.omit. Another possible value is NULL, no action. Value na.exclude can be useful.
model, x.ret, y.ret  logicals indicating if the model frame, the model matrix and the response are to be returned, respectively.
contrasts  an optional list. See the contrasts.arg of model.matrix.default.
offset  this can be used to specify an \emph{a priori} known component to be included in the linear predictor during fitting. An offset term can be included in the formula instead or as well, and if both are specified their sum is used.
x  Predictor variables. Matrix. Matrix of explanatory variables (also called 'regressors') of dimension n x (p-1) where p denotes the number of explanatory variables including the intercept. Rows of X represent observations, and columns represent variables. By default, there is a constant term in the model, unless you explicitly remove it using input option intercept=FALSE, so do not include a column of 1s in X. Missing values (NA’s) and infinite values (Inf’s) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.
y  Response variable. Vector. Response variable, specified as a vector of length n, where n is the number of observations. Each entry in y is the response for the corresponding row of X. Missing values (NA’s) and infinite values (Inf’s) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.
bsb  Initial subset - vector of indices. If bsb=0 (default) then the procedure starts with p units randomly chosen. If bsb is not 0 the search will start with m0=length(bsb).
intercept  Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.
monitoring  wheather to perform monitoring for several quantities in each step of the forward search. Default is monitoring=FALSE.
control  A control object (S3) containing estimation options, as returned by FSR_control. Use the function FSR_control and see its help page. If the control object is supplied, the parameters from it will be used. If parameters are passed also in the invocation statement, they will override the corresponding elements of the control object.
trace  Whether to print intermediate results. Default is trace=FALSE.
...  Potential further optional arguments, see the help of the function FSR_control.
Value

Depending on the input parameter `monitoring`, one of the following objects will be returned:

1. `fsr.object`
2. `fsreda.object`

Author(s)

FSDA team

References


Examples

```r
## Not run:
n <- 200
p <- 3
X <- matrix(data=rnorm(n*p), nrow=n, ncol=p)
y <- matrix(data=rnorm(n*1), nrow=n, ncol=1)
(out = fsrbase(X, y))

## Now we use the formula interface:
(out1 = fsrbase(y~X, control=FSR_control(plot=FALSE)))

## Or use the variables in a data frame
(out2 = fsrbase(Y~, data=hbk, control=FSR_control(plot=FALSE)))

## let us compare to the LTS solution
(out3 = ltsReg(Y~, data=hbk))

## Now compute the model without intercept
(out4 = fsrbase(Y~.-1, data=hbk, control=FSR_control(plot=FALSE)))

## And compare again with the LTS solution
(out5 = ltsReg(Y~.-1, data=hbk))

## using default (optional arguments)
(out6 = fsrbase(Y~.-1, data=hbk, control=FSR_control(plot=FALSE, nsamp=1500, h=50)))

## End(Not run)
```
Description

An object of class `fsreda.object` holds information about the result of a call to `fsreg`.

Value

The object itself is basically a `list` with the following components:

- **RES**: n x (n-init+1) matrix containing the monitoring of scaled residuals: the first row is the residual for the first unit, ..., n-th row is the residual for the n-th unit.
- **LEV**: (n+1) x (n-init+1) matrix containing the monitoring of leverage: the first row is the leverage for the first unit, ..., n-th row is the leverage for the n-th unit.
- **BB**: n x (n-init+1) matrix containing the information about the units belonging to the subset at each step of the forward search: first col contains indexes of the units forming subset in the initial step; ...; last column contains units forming subset in the final step (all units).
- **mdr**: n-init x 3 matrix which contains the monitoring of minimum deletion residual or (m+1)-ordered residual at each step of the forward search: first col is the fwd search index (from init to n-1); 2nd col = minimum deletion residual; 3rd col = (m+1)-ordered residual.
  
  Remark: these quantities are stored with sign, that is the min deletion residual is stored with negative sign if it corresponds to a negative residual.
- **msr**: n-init+1 x 3 matrix which contains the monitoring of maximum studentized residual or m-th ordered residual: first col is the fwd search index (from init to n); 2nd col = maximum studentized residual; 3rd col = (m)-ordered studentized residual.
- **nor**: (n-init+1) x 4 matrix containing the monitoring of normality test in each step of the forward search: first col = fwd search index (from init to n); 2nd col = Asymmetry test; 3rd col = Kurtosis test; 4th col = Normality test.
- **Bols**: (n-init+1) x (p+1) matrix containing the monitoring of estimated beta coefficients in each step of the forward search.
- **S2**: (n-init+1) x 5 matrix containing the monitoring of S2 or R2 and F-test in each step of the forward search:
  1. 1st col = fwd search index (from init to n);
  2. 2nd col = monitoring of S2;
  3. 3rd col = monitoring of R2;
  4. 4th col = monitoring of rescaled S2. In this case the estimated of sigma^2 at step m is divided by the consistency factor (to make the estimate asymptotically unbiased)
  5. 5th col = monitoring of F test. Note that an asymptotic unbiased estimate of sigma^2 is used.
In this case the estimated of $s^2$ at step $m$ is divided by the consistency factor (to make the estimate asymptotically unbiased).

**coo**  
(n-init+1) x 3 matrix containing the monitoring of Cook or modified Cook distance in each step of the forward search:  
1. 1st col = fwd search index (from init to n);  
2. 2nd col = monitoring of Cook distance;  
3. 3rd col = monitoring of modified Cook distance.

**Tols**  
(n-init+1) x (p+1) matrix containing the monitoring of estimated t-statistics (as specified in option input ‘tstat’) in each step of the forward search

**Un**  
(n-init) x 11 matrix which contains the unit(s) included in the subset at each step of the fwd search.  
REMARK: in every step the new subset is compared with the old subset. Un contains the unit(s) present in the new subset but not in the old one. For example, Un contains the unit included in step init+1 Un(end,2) contains the units included in the final step of the search.

**betaINT**  
Confidence intervals for the elements of $\beta$. betaINT is a (n-init+1)-by-2*length(confint)-by-p 3D array. Each third dimension refers to an element of beta:  
1. betaINT[,1] is associated with first element of beta;  
2. ...;  
3. betaINT[,p] is associated with last element of beta.  
The first two columns contain the lower and upper confidence limits associated with conflev(1). Columns three and four contain the lower and upper confidence limits associated with conflev(2); ...; The last two columns contain the lower and upper confidence limits associated with conflev(end). For example betaINT[,3:4,5] contain the lower and upper confidence limits for the fifth element of beta using confidence level specified in the second element of input option conflev.

**sigma2INT**  
Confidence interval for $s^2$.  
1. 1st col = fwd search index;  
2. 2nd col = lower confidence limit based on conflev(1);  
3. 3rd col = upper confidence limit based on conflev(1);  
4. 4th col = lower confidence limit based on conflev(2);  
5. 5th col = upper confidence limit based on conflev(2);  
6. ...  
7. penultimate col = lower confidence limit based on conflev(end);  
8. last col = upper confidence limit based on conflev(end).

**X**  
the data matrix X

**y**  
the response vector y

The object has class "fsreda".
Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="FS", monitoring=TRUE))
class(out)
summary(out)
## End(Not run)
```

---

**FSReda_control**

*Creates an FSReda_control object*

**Description**

Creates an object of class FSReda_control to be used with the fsreg() function, containing various control parameters.

**Usage**

```r
FSReda_control(intercept = TRUE, init, nocheck = FALSE,
tstat = c("trad", "scal"), conflev = c(0.95, 0.99))
```

**Arguments**

- **intercept**
  Indicator for constant term. Scalar. If `intercept=TRUE`, a model with constant term will be fitted (default), else, no constant term will be included.

- **init**
  Search initialization, scalar. It specifies the initial subset size to start monitoring exceedances of minimum deletion residual, if `init` is not specified it set equal to: \( p+1 \), if the sample size is smaller than 40 or \( \min(3*p+1, \text{floor}(0.5*(n+p+1))) \), otherwise. For example, if `init=100`, the procedure starts monitoring from step \( m=100 \).

- **nocheck**
  Check input arguments, scalar. If `nocheck=TRUE` no check is performed on matrix \( y \) and matrix \( X \). Notice that \( y \) and \( X \) are left unchanged. In other words the additional column of ones for the intercept is not added. As default `nocheck=FALSE`.

- **tstat**
  The kind of t-statistics which have to be monitored. `tstat="trad"` implies monitoring of traditional t statistics (out$Tols). In this case the estimate of \( s^2 \) at step \( m \) is based on \( s^2_m \) (notice that \( s^2_m=s^2 \) when \( m/n \) is small) `tstat="scal"` (default) implies monitoring of rescaled t statistics. In this case the estimate of \( s^2 \) at step \( m \) is based on \( s^2_m/\text{vartruncnorm}(m/n) \) where \( \text{vartruncnorm}(m/n) \) is the variance of the truncated normal distribution.

- **conflev**
  Confidence level which is used to declare units as outliers. Usually `conflev=0.95, 0.975, 0.99` (individual alpha) or `conflev=1-0.05/n, 1-0.025/n, 1-0.01/n` (simultaneous alpha). Default value is 0.975.
Details

Creates an object of class `FSReda_control` to be used with the `fsreg()` function, containing various control parameters.

Value

An object of class "FSReda_control" which is basically a list with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also as `FSR_control`, `MMreg_control` and `LXS_control`

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="FS", monitoring=TRUE,
             control=FSReda_control(tstat="scal")))
## End(Not run)
```

## fsreg

### fsreg: an automatic outlier detection procedure in linear regression

Description

An automatic outlier detection procedure in linear regression

Usage

```r
fsreg(x, ...)
```

### S3 method for class 'formula'

```r
fsreg(formula, data, subset, weights, na.action,
      model = TRUE, x.ret = FALSE, y.ret = FALSE,
      contrasts = NULL, offset, ...)
```

### Default S3 method:

```r
fsreg(x, y, bsb, intercept = TRUE,
      family = c("homo", "hetero", "bayes"),
      method = c("FS", "S", "MM", "LTS", "LMS"),
      monitoring = FALSE, control, trace = FALSE, ...
)```
Arguments

**formula**
a formula of the form y \sim x1 + x2 +....

**data**
data frame from which variables specified in formula are to be taken.

**subset**
an optional vector specifying a subset of observations to be used in the fitting process.

**weights**
an optional vector of weights to be used in the fitting process. **NOT USED YET.**

**na.action**
a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The “factory-fresh” default is na.omit. Another possible value is NULL, no action. Value na.exclude can be useful.

**model, x.ret, y.ret**
logicals indicating if the model frame, the model matrix and the response are to be returned, respectively.

**contrasts**
an optional list. See the contrasts.arg of model.matrix.default.

**offset**
this can be used to specify an *a priori* known component to be included in the linear predictor during fitting. An offset term can be included in the formula instead or as well, and if both are specified their sum is used.

**family**
family of robust regression models, can be 'homo' for homoscedastic (same variance) regression model, 'hetero' for heteroskedastic regression model or 'bayes' Bayesian linear regression. The default is family='homo' for homoscedastic regression model.

**method**
robust regression estimation model, can be 'FS' for Forward search, 'S' for S regression, 'MM' for MM regression, 'LMS' or 'LTS'. The default is method='FS' for forward search estimation.

**monitoring**
whether to perform monitoring for several quantities in each step of the forward search or for series of values of the breakdown point in case of S estimates or for series of values of the efficiency in case of MM estimates. Default is monitoring=FALSE.

**y**
Response variable. Vector. Response variable, specified as a vector of length n, where n is the number of observations. Each entry in y is the response for the corresponding row of X. Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

**x**
Predictor variables. Matrix. Matrix of explanatory variables (also called 'regressors') of dimension n x (p-1) where p denotes the number of explanatory variables including the intercept. Rows of X represent observations, and columns represent variables. By default, there is a constant term in the model, unless you explicitly remove it using input option intercept=FALSE, so do not include a column of 1s in X. Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

**bsb**
Initial subset - vector of indices. If bsb=0 (default) then the procedure starts with p units randomly chosen. If bsb is not 0 the search will start with m0=length(bsb).
intercept  Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.

ccontrol  A control object (S3) containing estimation options. If the control object is supplied, the parameters from it will be used. If parameters are passed also in the invocation statement, they will override the corresponding elements of the control object.

trace  Whether to print intermediate results. Default is trace=FALSE.

Value

Depending on the input parameters family and method, one of the following objects will be returned:

1. `fsr.object`
2. `sreg.object`
3. `mmreg.object`
4. `fsdalms.object`
5. `fsdalts.object`
6. `fsreda.object`
7. `sregeda.object`
8. `mmregeda.object`

Author(s)

FSDA team

References


Examples

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

n <- 200
p <- 3

X <- matrix(data=rnorm(n*p), nrow=n, ncol=p)
y <- matrix(data=rnorm(n*1), nrow=n, ncol=1)
(out = fsreg(X, y))

## Now we use the formula interface:
(out1 = fsreg(y~X, control=FSR_control(plot=FALSE)))
```
## Or use the variables in a data frame

```r
(out2 = fsreg(Y~., data=hbk, control=FSR_control(plot=FALSE)))
```

## Let us compare to the LTS solution

```r
library(robustbase)
(out3 = ltsReg(Y~., data=hbk))
```

## Now compute the model without intercept

```r
(out4 = fsreg(Y~.-1, data=hbk, control=FSR_control(plot=FALSE)))
```

## And compare again with the LTS solution

```r
(out5 = ltsReg(Y~.-1, data=hbk))
```

## Using default (optional arguments)

```r
(out6 = fsreg(Y~.-1, data=hbk, control=FSR_control(plot=FALSE, nsamp=1500, h=50)))
```

## End(Not run)

---

### fsrfan

**Robust transformations for regression**

### Description

The transformations for negative and positive responses were determined by Yeo and Johnson (2000) by imposing the smoothness condition that the second derivative of $zYJ (\lambda)$ with respect to $y$ be smooth at $y = 0$. However some authors, for example Weisberg (2005), query the physical interpretability of this constraint which is oftentimes violated in data analysis. Accordingly, Atkinson et al. (2019) and (2020) extend the Yeo-Johnson transformation to allow two values of the transformations parameter: $\lambda_N$ for negative observations and $\lambda_P$ for non-negative ones.

FSRfan monitors:

1. the t test associated with the constructed variable computed assuming the same transformation parameter for positive and negative observations fixed. In short we call this test, "global score test for positive observations".
2. the t test associated with the constructed variable computed assuming a different transformation for positive observations keeping the value of the transformation parameter for negative observations fixed. In short we call this test, "test for positive observations".
3. the t test associated with the constructed variable computed assuming a different transformation for negative observations keeping the value of the transformation parameter for positive observations fixed. In short we call this test, "test for negative observations".
4. the F test for the joint presence of the two constructed variables described in points 2) and 3).
5. the F likelihood ratio test based on the MLE of $\lambda_P$ and $\lambda_N$. In this case the residual sum of squares of the null model based on a single transformation parameter $\lambda$ is compared with the residual sum of squares of the model based on data transformed data using MLE of $\lambda_P$ and $\lambda_N$. 
Usage

fsrfan(x, ...)

## S3 method for class 'formula'
fsrfan(
  formula,
  data,
  subset,
  weights,
  na.action,
  model = TRUE,
  x.ret = FALSE,
  y.ret = FALSE,
  contrasts = NULL,
  offset,
  ...
)

## Default S3 method:
fsrfan(
  x,
  y,
  intercept = TRUE,
  plot = FALSE,
  family = c("BoxCox", "YJ", "YJpn", "YJall"),
  la = c(-1, -0.5, 0, 0.5, 1),
  lms,
  alpha = 0.75,
  h,
  init,
  msg = FALSE,
  nocheck = FALSE,
  nsamp = 1000,
  conflev = 0.99,
  xlab,
  ylab,
  main,
  xlim,
  ylim,
  lwd = 2,
  lwd.env = 1,
  trace = FALSE,
  ...
)

## S3 method for class 'fsrfan'
plot(
  x,
conflev = 0.99,
xlim, ylim,
xlab = "Subset of size m",
ylab = "Score test statistic",
main = "Fan plot",
col, lty, lwd = 2.5,
lwd.env = 1,
...)

Arguments

x An $n \times p$ data matrix ($n$ observations and $p$ variables). Rows of $x$ represent observations, and columns represent variables. Missing values (NA’s) and infinite values (Inf’s) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

... potential further arguments passed to lower level functions.

formula a formula of the form $y \sim x1 + x2 + ...$
data data frame from which variables specified in formula are to be taken.

subset an optional vector specifying a subset of observations to be used in the fitting process.

weights an optional vector of weights to be used

NOT USED YET.

na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The “factory-fresh” default is na.omit. Another possible value is NULL, no action. Value na.exclude can be useful.

model logical indicating if the model frame, is to be returned.

x.ret logical indicating if the the model matrix is to be returned.

y.ret logical indicating if the response is to be returned.

contrasts an optional list. See the contrasts.arg of model.matrix.default.

offset this can be used to specify an a priori known component to be included in the linear predictor during fitting. An offset term can be included in the

y Response variable. A vector with $n$ elements that contains the response variable.

intercept wheather to use constant term (default is intercept=TRUE

plot If plot=FALSE (default) no plot is produced. If plot=TRUE a fan plot is shown.

family string which identifies the family of transformations which must be used. Possible values are c(‘BoxCox’, ‘YJ’, ‘YJpn’, ‘YJall’). Default is ‘BoxCox’. The Box-Cox family of power transformations equals $(y^\lambda - 1)/\lambda$ for $\lambda$ not equal to zero, and $\log(y)$ if $\lambda = 0$. The Yeo-Johnson (YJ) transformation is
the Box-Cox transformation of \( y + 1 \) for nonnegative values, and of \(|y| + 1\) with parameter \( 2 - \lambda \) for \( y \) negative. Remember that BoxCox can be used only if input \( y \) is positive. Yeo-Johnson family of transformations does not have this limitation. If \( \text{family} = 'YJpn' \) Yeo-Johnson family is applied but in this case it is also possible to monitor (in the output arguments \( \text{Scorep} \) and \( \text{Scoren} \)) the score test for positive and negative observations respectively. If \( \text{family} = 'YJall' \), it is also possible to monitor the joint F test for the presence of the two constructed variables for positive and negative observations.

\( \lambda \) values of the transformation parameter for which it is necessary to compute the score test. Default value of lambda is \( \lambda = c(-1, -0.5, 0, 0.5, 1) \), i.e., the five most common values of lambda.

\( \text{lms} \) how to find the initial subset to initialize the search. If \( \text{lms} = 1 \) (default) Least Median of Squares (LMS) is computed, else Least Trimmed Squares (LTS) is computed. If \( \text{lms} \) is matrix of size \( p - 1 + \text{intercept} \times \text{length}(\lambda) \) it contains in column \( j = 1, \ldots, \text{length}(\lambda) \) the list of units forming the initial subset for the search associated with \( \lambda(j) \). In this case the input option \( \text{nsamp} \) is ignored.

\( \alpha \) the percentage (roughly) of squared residuals whose sum will be minimized, by default \( \alpha = 0.5 \). In general, \( \alpha \) must between 0.5 and 1.

\( h \) The number of observations that have determined the least trimmed squares estimator, scalar. \( h \) is an integer greater or equal than \( p \) but smaller than \( n \). Generally \( h = \lfloor 0.5 \times (n + p + 1) \rfloor \) (default value).

\( \text{init} \) Search initialization. It specifies the initial subset size to start monitoring the value of the score test. If \( \text{init} \) is not specified it will be set equal to: \( p + 1 \), if the sample size is smaller than 40 or \( \min(3 \times p + 1, \text{floor}(0.5 \times (n + p + 1))) \), otherwise.

\( \text{msg} \) Controls whether to display or not messages on the screen. If \( \text{msg} = \text{TRUE} \) messages are displayed on the screen. If \( \text{msg} = 2 \), detailed messages are displayed, for example the information at iteration level.

\( \text{nocheck} \) Whether to check input arguments. If \( \text{nocheck} = \text{TRUE} \) no check is performed on matrix \( y \) and matrix \( X \). Notice that \( y \) and \( X \) are left unchanged. In other words the additional column of ones for the intercept is not added. The default is \( \text{nocheck} = \text{FALSE} \).

\( \text{nsamp} \) number of subsamples which will be extracted to find the robust estimator. If \( \text{nsamp} = 0 \) all subsets will be extracted. They will be \( n \) choose \( p \).

Remark: if the number of all possible subset is \(<1000\) the default is to extract all subsets otherwise just 1000. If \( \text{nsamp} \) is a matrix of size \( r \times 1 \), it contains in the rows the subsets which sill have to be extracted. For example, if \( p = 3 \) and \( \text{nsamp} = c(2, 4, 9; 23, 45, 49; 90, 34, 1) \) the first subset is made up of units \( c(2, 4, 9) \), the second subset of units \( c(23, 45, 49) \) and the third subset of units \( c(90 34 1) \).

\( \text{conflev} \) Confidence level for the bands (default is 0.99, that is, we plot two horizontal lines corresponding to values -2.58 and 2.58).

\( \text{xlab} \) A label for the X-axis, default is 'Subset size m'

\( \text{ylab} \) A label for the Y-axis, default is 'Score test statistic'

\( \text{main} \) A label for the title, default is 'Fan plot'
xlim  Minimum and maximum for the X-axis
ylim  Minimum and maximum for the Y-axis
lwd  The line width of the curves which contain the score test, a positive number, default is lwd=2
lwd.env  The line width of the lines associated with the envelopes, a positive number, default is lwd.env=1
trace  Whether to print intermediate results. Default is trace=FALSE.
col  a vector specifying the colors for the lines, each one corresponding to a la value. if length(col) < length(la), the colors will be recycled.
lty  a vector specifying the line types for the lines, each one corresponding to a la value. if length(col) < length(la), the colors will be recycled.

Value

An S3 object of class fsrfan.object will be returned which is basically a list containing the following elements:

1. la: vector containing the values of lambda for which fan plot is constructed
2. bs: matrix of size p X length(la) containing the units forming the initial subset for each value of lambda
3. Score: a matrix containing the values of the score test for each value of the transformation parameter:
   - 1st col = fwd search index;
   - 2nd col = value of the score test in each step of the fwd search for la[1]
   - ...
4. Scorep: matrix containing the values of the score test for positive observations for each value of the transformation parameter.
   Note: this output is present only if input option family='YJpn' or family='YJall'.
5. Scoren: matrix containing the values of the score test for negative observations for each value of the transformation parameter.
   Note: this output is present only if input option 'family' is 'YJpn' or 'YJall'.
6. Scoreb: matrix containing the values of the score test for the joint presence of both constructed variables (associated with positive and negative observations) for each value of the transformation parameter. In this case the reference distribution is the \( F \) with 2 and subset.size - p degrees of freedom.
   Note: this output is present only if input option family='YJall'.
7. Un: a three-dimensional array containing length(la) matrices of size \( \text{retnUn}=(n\text{-init}) \times \text{retpUn}=11 \). Each matrix contains the unit(s) included in the subset at each step in the search associated with the corresponding element of la.
   REMARK: at each step the new subset is compared with the old subset. Un contains the unit(s) present in the new subset but not in the old one.

Author(s)

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References


Examples

```r
## Not run:
data(wool)
XX <- wool
y <- XX[, ncol(XX)]
X <- XX[, 1:(ncol(XX)-1), drop=FALSE]

out <- fsrfan(X, y) # call 'fsrfan' with all default parameters
out <- fsrfan(cycles~, data=wool) # use the formula interface

set.seed(10)
out <- fsrfan(cycles~, data=wool, plot=TRUE) # call 'fsrfan' and produce the plot

plot(out) # use the plot method on the fsrfan object
plot(out, conflev=c(0.9, 0.95, 0.99)) # change the confidence level in the plot method

##======================
##
## fsrfan() with all default options.
##
## Store values of the score test statistic for the five most common
## values of \$\lambda\$. Produce also a fan plot and display it on the screen.
## Common part to all examples: load 'wool' data set.
data(wool)
head(wool)
dim(wool)

## The function fsrfan() stores the score test statistic.
## In this case we use the five most common values of lambda are considered

out <- fsrfan(cycles~, data=wool)
plot(out)
  # fanplot(out) # Not yet implemented in fsdaR

## The fan plot shows the log transformation is diffused throughout the data
## and does not depend on the presence of particular observations.

##
## Example specifying 'lambda'.
```
## Produce a fan plot for each value of 'lambda' in the vector 'la'.
## Extract in matrix 'Un' the units which entered the search in each step

data(wool)
out <- fsrfan(cycles~., data=wool, la=c(-1, -0.5, 0, 0.5), plot=TRUE)
plot(out)

out$Un[,2,]

## Example specifying the confidence level and the initial starting point for monitoring.
## Construct the fan plot specifying the confidence level and the initial starting point
## for monitoring.
data(wool)
out <- fsrfan(cycles~., data=wool, init=ncol(wool)+1, nsamp=0, conflev=0.95, plots=TRUE)
plot(out, conflev=0.95)

## Example with starting point based on LTS.
## Extract all subsamples, construct a fan plot specifying the confidence level
## and the initial starting point for monitoring based on p+2 observations,
## strong line width for lines associated with the confidence bands.
data(wool)
out <- fsrfan(cycles~., data=wool, init=ncol(wool)+1, nsamp=0, lms=0, lwd.env=3, plot=TRUE)
plot(out, lwd.env=3)

## Fan plot using the loyalty cards data.
## In this example, 'la' is the vector containing the most common values
## of the transformation parameter.
## Store the score test statistics for the specified values of lambda
## and automatically produce the fan plot
data(loyalty)
head(loyalty)
dim(loyalty)

la is a vector containing the most common values of the transformation parameter
out <- fsrfan(amount_spent~., data=loyalty, la=c(0, 1/3, 0.4, 0.5),
ininit=ncol(loyalty)+1, plot=TRUE, lwd=3)
plot(out, lwd=3)

## The fan plot shows that even if the third root is the best value of the transformation
## parameter at the end of the search, in earlier steps it lies very close to the upper
## rejection region. The best value of the transformation parameter seems to be the one
## associated with la=0.4, which is always the confidence bands but at the end of search,
## due to the presence of particular observations it goes below the lower rejection line.

## Compare BoxCox with Yeo and Johnson transformation.
## Store values of the score test statistic for the five most common
## values of lambda. Produce also a fan plot and display it on the screen.
## Common part to all examples: load wool dataset.
data(wool)

## Store the score test statistic using Box Cox transformation.
outBC <- fsrfan(cycles~, data=wool, nsamp=0)

## Store the score test statistic using Yeo and Johnson transformation.
outYJ <- fsrfan(cycles~, data=wool, family="YJ", nsamp=0)

## Not yet fully implemented
## fanplot(outBC, main="Box Cox")
## fanplot(outYJ, main="Yeo and Johnson")

plot(outBC, main="Box Cox")
plot(outYJ, main="Yeo and Johnson")

cat("Maximum difference in absolute value: ",
    max(max(abs(outYJ$Score - outBC$Score), na.rm=TRUE)), "\n")

======================
## Call 'fsrfan' with Yeo-Johnson (YJ) transformation
out <- fsrfan(cycles~, data=wool, family="YJ")
plot(out)

## End(Not run)

---

fsrfan.object  
*Objects returned by the function fsrfan*

**Description**

An object of class *fsrfan.object* holds information about the result of a call to *fsrfan*.

**Value**

The functions `print()` and `summary()` are used to obtain and print a summary of the results. An object of class *fsrfan* is a list containing at least the following components:

1. 1a vector containing the values of lambda for which fan plot is constructed
2. bs matrix of size $p \times \text{length}(1a)$ containing the units forming the initial subset for each value of lambda
3. Score a matrix containing the values of the score test for each value of the transformation parameter:
   - 1st col = fwd search index;
• 2nd col = value of the score test in each step of the fwd search for la[1]
• ...

4. Scorep matrix containing the values of the score test for positive observations for each value of the transformation parameter.
   Note: this output is present only if input option family='YJpn' or family='YJall'.

5. Scoren matrix containing the values of the score test for negative observations for each value of the transformation parameter.
   Note: this output is present only if input option 'family' is 'YJpn' or 'YJall'.

6. Scoreb matrix containing the values of the score test for the joint presence of both constructed variables (associated with positive and negative observations) for each value of the transformation parameter. In this case the reference distribution is the $F$ with $2$ and $\text{subset.size} - p$ degrees of freedom.
   Note: this output is present only if input option family='YJall'.

7. Un a three-dimensional array containing length(la) matrices of size retnUn=(n-init) X retpUn=11. Each matrix contains the unit(s) included in the subset at each step in the search associated with the corresponding element of la.
   REMARK: at each step the new subset is compared with the old subset. Un contains the unit(s) present in the new subset but not in the old one.

Examples

```r
## Not run:
data(wool)
XX <- wool
y <- XX[, ncol(XX)]
X <- XX[, 1:(ncol(XX)-1), drop=FALSE]

out <- fsrfan(X, y)

class(out)
summary(out)
```

### FSR_control

* **FSR_control**

* **Description**

  Creates an object of class FSR_control to be used with the fsreg() function, containing various control parameters.
Usage

FSR_control(intercept = TRUE, h, nsamp = 1000, lms = 1, init, nocheck = FALSE, bonflev = "", msg = TRUE, bsmfullrank = TRUE, plot = FALSE, bivarfit = FALSE, multivarfit = FALSE, labeladd = FALSE, nameX, namey, ylim, xlim)

Arguments

intercept  
Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.

h  
The number of observations that have determined the least trimmed squares estimator, scalar. h is an integer greater or equal than p but smaller than n. Generally if the purpose is outlier detection h=[0.5*(n+p+1)] (default value). h can be smaller than this threshold if the purpose is to find subgroups of homogeneous observations. In this function the LTS/LMS estimator is used just to initialize the search.

nsamp  
Number of subsamples which will be extracted to find the robust estimator, scalar. If nsamp=0 all subsets will be extracted. They will be \((n choose p)\). If the number of all possible subset is <1000 the default is to extract all subsets otherwise just 1000.

lms  
Criterion to use to find the initial subset to initialize the search (LMS, LTS with concentration steps, LTS without concentration steps or subset supplied directly by the user). The default value is 1 (Least Median of Squares is computed to initialize the search). On the other hand, if the user wants to initialize the search with LTS with all the default options for concentration steps then lms=2. If the user wants to use LTS without concentration steps, lms can be a scalar different from 1 or 2. If lms is a list it is possible to control a series of options for concentration steps (for more details see option lms inside LXS_control). If, on the other hand, the user wants to initialize the search with a prespecified set of units there are two possibilities:

1. lms can be a vector with length greater than 1 which contains the list of units forming the initial subset. For example, if the user wants to initialize the search with units 4, 6 and 10 then lms=c(4, 6, 10);
2. lms is a struct which contains a field named bsb which contains the list of units to initialize the search. For example, in the case of simple regression through the origin with just one explanatory variable, if the user wants to initialize the search with unit 3 then lms=list(bsb=3).

init  
Search initialization, scalar. It specifies the initial subset size to start monitoring exceedances of minimum deletion residual, if init is not specified it set equal to: \(p+1\), if the sample size is smaller than 40 or \(\min(3*p+1, \lceil 0.5*(n+p+1) \rceil)\), otherwise. For example, if init=100, the procedure starts monitoring from step \(m=100\).

nocheck  
Check input arguments, scalar. If nocheck=TRUE no check is performed on matrix y and matrix X. Notice that y and X are left unchanged. In other words the additional column of ones for the intercept is not added. As default nocheck=FALSE.
bonflev  Option to be used if the distribution of the data is strongly non normal and, thus, the general signal detection rule based on consecutive exceedances cannot be used. In this case bonflev can be:

1. a scalar smaller than 1 which specifies the confidence level for a signal and a stopping rule based on the comparison of the minimum MD with a Bonferroni bound. For example if bonflev=0.99 the procedure stops when the trajectory exceeds for the first time the 99% bonferroni bound.
2. A scalar value greater than 1. In this case the procedure stops when the residual trajectory exceeds for the first time this value.

Default value is ", which means to rely on general rules based on consecutive exceedances.

msg Controls whether to display or not messages on the screen If msg==1 (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.

bsbmfullrank How to behave in case subset at step m (say bsbm) produces a singular X. In other words, this options controls what to do when rank(X[bsbm,]) is smaller then number of explanatory variables. If bsbmfullrank=1 (default) these units (whose number is say mnofullrank) are constrained to enter the search in the final n-mnofullrank steps else the search continues using as estimate of beta at step m the estimate of beta found in the previous step.

plot Plot on the screen. Scalar. If plot=TRUE the plot of minimum deletion residual with envelopes based on n observations and the scatterplot matrix with the outliers highlighted is produced. If plot=2 the user can also monitor the intermediate plots based on envelope superimposition. If plot=FALSE (default) no plot is produced.

bivarfit Wheather to superimpose bivariate least square lines on the plot (if plot=TRUE. This option adds one or more least squares lines, based on SIMPLE REGRESSION of y on Xi, to the plots of y|Xi. The default is bivarfit=FALSE: no line is fitted. If bivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. If bivarfit=2, two OLS lines are fitted: one to all points and another to the group of the genuine observations. The group of the potential outliers is not fitted. If bivarfit=0 one OLS line is fitted to each group. This is useful for the purpose of fitting mixtures of regression lines. If bivarfit='i1' or bivarfit='i2', etc. an OLS line is fitted to a specific group, the one with index 'i' equal to 1, 2, 3 etc. Again, useful in case of mixtures.

multivarfit Wheather to superimpose multivariate least square lines. This option adds one or more least square lines, based on MULTIVARIATE REGRESSION of y on X, to the plots of y|Xi. The default is multivarfit=FALSE: no line is fitted. If multivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. The line added to the scatter plot y|Xi is avconst + Ci*Xi, where Ci is the coefficient of Xi in the multivariate regression and avconst is the effect of all the other explanatory variables different from Xi evaluated at their centroid (that is overline(y)'C)). If multivarfit=2, same action as with multivarfit=1 but this time we also add the line based on the group of unselected observations (i.e. the normal units).
labeladd Add outlier labels in plot. If labeladd=TRUE, we label the outliers with the unit row index in matrices X and y. The default value is labeladd=FALSE, i.e. no label is added.

nameX Add variable labels in plot. A vector of strings of length p containing the labels of the variables of the regression dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically.

namey Add response label. A string containing the label of the response.

ylim Control y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.

xlim Control x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.

Details

Creates an object of class FSR_control to be used with the fsreg() function, containing various control parameters.

Value

An object of class "FSR_control" which is basically a list with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also Sreg_control, MMreg_control, LXS_control, FSReda_control, Sregeda_control and MMregeda_control.

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="FS", control=FSR_control(h=56, nsamp=500, lms=2)))
summary(out)
## End(Not run)
```
**geyser2**

*Old Faithful Geyser Data.*

**Description**

A bivariate data set obtained from the Old Faithful Geyser, containing the eruption length and the length of the previous eruption for 271 eruptions of this geyser in minutes.

**Usage**

data(geyser2)

**Format**

A data frame with 271 rows and 2 variables The variables are as follows:

- Eruption length: The eruption length in minutes.
- Previous eruption length: The length of the previous eruption in minutes.

**References**


---

**hawkins**

*Hawkins data.*

**Description**

These data, simulated by Hawkins, consist of 128 observations and eight explanatory variables ($X_1$, $\ldots$, $X_8$) and one dependent variable, $y$.

**Usage**

data(hawkins)

**Format**

A data frame with 128 rows and 9 variables
Description

Data on the logged survival time of 108 patients undergoing liver surgery, together with four potential explanatory variables. Data are composed of 54 observations plus other 54 observations, introduced to check the model fitted to the first 54. Their comparison suggests there is no systematic difference between the two sets. However by looking at some FS plots (Riani and Atkinson, 2007), we conclude that these two groups are significantly different.

Usage

data(hospital)

Format

A data frame with 108 rows and 5 variables. The variables are as follows:

• X1
• X2
• X3
• X4
• y


@references A. C. ATKINSON, M. RIANI, Robust Diagnostic Regression Analysis. Springer-Verlag, New York, 2000.

Description

Income data taken from the United States Census Bureau. The data are a random sample of 200 observations referred to four variables. The goal is to predict HTOTVAL.

Usage

data(Income1)
Income2

Format
A data frame with 200 rows and 4 variables. The variables are as follows:

- H_NUMPER: Number of persons in household
- HOTHVAL: All other types of income except HEARNVAL Recode - Total other household income
- HSSVAL: household income - social security
- HTOTVAL: total household income (dollar amount)

Source

Examples
```r
data(Income1)
head(Income1)
```

Description
A sample of 200 observations of full time employees from a municipality in Northern Italy who have declared extra income from investment sources. The variables are as follows. The goal is the possibility in predicting income level based on the individual’s personal information.

Usage
data(Income2)

Format
A data frame with 200 rows and 6 variables. The variables are as follows:

- Age: Age of the person (the minimum is 19 and the maximum is 73).
- Education: Number of years of education (the minimum value of 5 is primary school, and the maximum value is 16 bachelor degree)
- Gender: A factor - Male or Female
- ExtraGain: Income from investment sources (profit-losses) apart from wages/salary
- Hours: total number of declared hours worked during the week. The minimum value is 35 and the maximum is 99
- Income: total yearly income (Euro amount)
Examples

```r
data(Income2)
head(Income2)
```

data(Income2)

```r
default(Income2)
```

levfwdplot

Plots the trajectories of the monitored scaled (squared) residuals

Description

Plots the trajectories of the monitored scaled (squared) residuals

Usage

```r
levfwdplot(out, 
  xlim, ylim, xlab, ylab, main, lwd, lty, col, cex.lab, cex.axis, 
  xvalues, 
  fg.thresh, fg.unit, fg.labstep, fg.lwd, fg.lty, fg.col, fg.mark, fg.cex, 
  bg.thresh, bg.style, 
  xground=c("lev", "res"), tag, datatooltip, label, nameX, namey, msg, databrush, 
  standard, fground, bground, ...) 
```

Arguments

- `out` An object containing monitoring of leverage, `fsreda.object`. The needed elements of out are
  1. `LEV`: matrix containing the leverage monitored in each step of the forward search. Every row is associated with a unit. This matrix can be created using function `fsreg()` with method=“FS”, monitoring=TRUE.
  2. `Un`: (for FSR only) - matrix containing the order of entry in the subset of each unit (required only when datatooltip is true or databrush is not empty).
  3. `y`: a vector containing the response (required only when option databrush is requested).
  4. `X`: a matrix containing the explanatory variables (required only when option databrush is requested).
  5. `Bols`: (n-init+l) x (p+1) matrix containing the estimated beta coefficients monitored in each step of the robust procedure (required only when option databrush is requested and suboption multivarfit is requested).
- `ylim` Control y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.
- `xlim` Control x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.
- `xlab` a title for the x axis
- `ylab` a title for the y axis
**levfwdplot**

- **main**: an overall title for the plot
- **lwd**: The line width, a positive number, defaulting to 1
- **lty**: The line type. Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.
- **col**: colors to be used for the highlighted units
- **cex.lab**: The magnification to be used for x and y labels relative to the current setting of cex
- **cex.axis**: The magnification to be used for axis annotation relative to the current setting of cex
- **xvalues**: values for the x axis. Numeric vector of ncol(RES) controlling the x axis coordinates. The default value of xvalues is \((nrow(RES) - ncol(RES) + 1):nrow(RES)\)
- **fg.thresh**: (alternative to fg.unit) numeric vector of length 1 or 2 which specifies the highlighted trajectories. If \(\text{length}(\text{fthresh}) == 1\) the highlighted trajectories are those of units that throughout the search had at least once a residual greater (in absolute value) than thresh. The default value is \(\text{fg.thresh}=2.5\). If \(\text{length}(\text{fthresh}) == 2\) the highlighted trajectories are those of units that throughout the search had a residual at least once bigger than \(\text{fg.thresh}[2]\) or smaller than \(\text{fg.thresh}[1]\).
- **fg.unit**: (alternative to fg.thresh), vector containing the list of the units to be highlighted. If fg.unit is supplied, fg.thresh is ignored.
- **fg.labstep**: numeric vector which specifies the steps of the search where to put labels for the highlighted trajectories (units). The default is to put the labels at the initial and final steps of the search. iflabstep=' ' means no label.
- **fg.lwd**: The line width for the highlighted trajectories (units). Default is 1.
- **fg.lty**: The line type for the highlighted trajectories (units). Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.
- **fg.col**: colors to be used for the highlighted units.
- **fg.mark**: Controls whether to plot highlighted trajectories as symbols. if fg.mark==TRUE each line is plotted using a different symbol else no marker is used (default).
- **fg.cex**: controls the font size of the labels of the trajectories in foreground.
- **bg.thresh**: numeric vector of length 1 or 2 which specifies how to define the unimportant trajectories. Unimportant trajectories will be plotted using a colormap, in grey or will be hidden. If \(\text{length}(\text{thresh}) == 1\) the irrelevant units are those which always had a residual smaller (in absolute value) than thresh. If \(\text{length}(\text{bthresh}) == 2\) the irrelevant units are those which always had a residual greater than \(\text{bthresh}(1)\) and smaller than \(\text{bthresh}(2)\). The default is: \(\text{bg.thresh}=2.5\) if \(n > 100\) and \(\text{bg.thresh}=-\text{Inf}\) if \(n <= 100\) i.e. to treat all trajectories as important if \(n <= 100\) and, if \(n > 100\), to reduce emphasis only to trajectories having in all steps of the search a value of scaled residual smaller than 2.5.
- **bg.style**: specifies how to plot the unimportant trajectories as defined in option bthresh.
1. **bg.style="faint"**: unimportant trajectories are plotted using a colormap.
2. **bg.style="hide"**: unimportant trajectories are hidden.
3. **bg.style="greyish"**: unimportant trajectories are displayed in a faint grey.

When \( n > 100 \) the default option is **bg.style='faint'**. When \( n <= 100 \) and **bg.thresh == -Inf** option bstyle is ignored. Remark: bgground=" is equivalent to -Inf that is all trajectories are considered relevant.

**tag**

Plot handle. String which identifies the handle of the plot which is about to be created. The default is to use tag 'pl_resfwd'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.

**xground**

trajectories to highlight in connection with resfwdplot. If **xground="lev"** (default), the levfwdplot trajectories are put in foreground or in background depending on the leverage values. If **xground="res"**, the levfwdplot trajectories are put in foreground or in background depending on the residual values. See options bg.thresh and fg.thresh.

**datatooltip**

Interactive clicking. It is inactive if this parameter is missing or empty. The default is datatooltip=TRUE, i.e. the user can select with the mouse an individual residual trajectory in order to have information about the corresponding unit. The information displayed depends on the estimator in use.

For example for class **fsreda.object** the information concerns the label and the step of the search in which the unit enters the subset. If datatooltip is a list it may contain the following fields:

1. **DisplayStyle** determines how the data cursor displays. Possible values are 'datatip' and 'window' (default). 'datatip' displays data cursor information in a small yellow text box attached to a black square marker at a data point you interactively select. 'window' displays data cursor information for the data point you interactively select in a floating window within the figure.
2. **SnapToDataVertex**: specifies whether the data cursor snaps to the nearest data value or is located at the actual pointer position. Possible values are SnapToDataVertex='on' (default) and SnapToDataVertex='off'.
3. **LineColor**: controls the color of the trajectory selected with the mouse. It can be an RGB triplet of values between 0 and 1, or character vector indicating a color name. Note that a RGB vector can be conveniently chosen with our MATLAB class FSColor, see documentation.
4. **SubsetLinesColor**: enables to control the color of the trajectories of the units that are in the subset at a given step of the search (if levfwdplot() is applied to an object of class **fsreda.object**) or have a weight greater than 0.9 (if levfwdplot() is applied to an object of class **sregeda.object** or **mmregeda.object**). This can be done (repeatedly) with a left mouse click in proximity of the step of interest. A right mouse click will terminate the selection by marking with a up-arrow the step corresponding to the highlighted lines. The highlighted lines by default are in red, but a different color can be specified as RGB triplet or character of color name. Note that a
RGB vector can be conveniently chosen with our MATLAB class FSCColor, see documentation. By default SubsetLinesColor="", i.e. the modality is not active. Any initialization for SubsetLinesColor which cannot be interpreted as RGB vector will be converted to blue, i.e. SubsetLinesColor will be forced to be [0 0 1]. If SubsetLinesColor is not empty the previous optionLineColor is ignored.

**label**
Character vector containing the labels of the units (optional argument used when datatooltip=TRUE. If this field is not present labels row1, ..., rown will be automatically created and included in the pop up datatooltip window).

**nameX**
Add variable labels in plot. A vector of strings of length p containing the labels of the variables of the regression dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically

**namey**
Add response label. A string containing the label of the response

**msg**
Controls whether to display or not messages on the screen If msg==1 (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.

**databrush**
interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is a scalar or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the y|X plot, i.e. a matrix of scatter plots of y against each column of X, grouped according to the selection(s) done by brushing. If the plot y|X does not exist it is automatically created. In addition, brushed units are automatically highlighted in the minimum deletion residual plot if it is already open. The extension to the following plots will be available in future versions of the toolbox:

1. monitoring leverage plot;
2. maximum studentized residual;
3. s^2 and R^2;
4. Cook distance and modified Cook distance;
5. deletion t statistics.

Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist="").

If databrush=list(...), it is possible to use all optional arguments of function selectdataFS() and the following optional argument:

1. persist. Persist is an empty value or a character containing 'on' or 'off'. The default value is persist="", that is brushing is allowed only once. If persist="on" or persist="off" brushing can be done as many time as the user requires. If persist='on' then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If persist='off' every time a new brush is performed units previously brushed are removed.
2. bivarfit. Wheather to superimpose bivariate least square lines on the plot (if plot=TRUE). This option adds one or more least squares lines, based on SIMPLE REGRESSION of y on Xi, to the plots of y|Xi. The default is bivarfit=FALSE: no line is fitted. If bivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. If bivarfit=2, two OLS lines are fitted: one to all points and another to the group of the genuine observations. The group of the potential outliers is not fitted. If bivarfit=0 one OLS line is fitted to each group. This is useful for the purpose of fitting mixtures of regression lines. If bivarfit='i1' or bivarfit='i2', etc. an OLS line is fitted to a specific group, the one with index 'i' equal to 1, 2, 3 etc. Again, useful in case of mixtures.

3. multivarfit. Wheather to superimpose multivariate least square lines. This option adds one or more least square lines, based on MULTIVARIATE REGRESSION of y on X, to the plots of y|Xi. The default is multivarfit=FALSE: no line is fitted. If bivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. The line added to the scatter plot y|Xi is avconst + Ci*Xi, where Ci is the coefficient of Xi in the multivariate regression and avconst is the effect of all the other explanatory variables different from Xi evaluated at their centroid (that is overline(y)'C)). If multivarfit=2, same action as with multivarfit=1 but this time we also add the line based on the group of unselected observations (i.e. the normal units).

4. labeladd. Add outlier labels in plot. If labeladd=TRUE, we label the outliers with the unit row index in matrices X and y. The default value is labeladd=FALSE, i.e. no label is added.

standard (MATLAB-style arguments) appearance of the plot in terms of xlim, ylim, axes labels and their font size style, color of the lines, etc.
fground MATLAB-style arguments for the fground trajectories in foreground.
bground MATLAB-style arguments for the fground trajectories in background.
... potential further arguments passed to lower level functions.

Details
No details

Value
No value returned

Author(s)
FSDA team

Examples
## Not run:

n <- 100
loyalty

```r
y <- rnorm(n)
X <- matrix(rnorm(n*4), nrow=n)
out <- fsreg(y~X, method="LTS")
out <- fsreg(y~X, method="FS", bsb=out$bs, monitoring=TRUE)
levfwdplot(out)

## End(Not run)
```

---

**Description**

The loyalty data consist of 509 observations on the behaviour of customers with loyalty cards from a supermarket chain in Northern Italy. The response \( y \) is the amount in euros spent at the shop over six months and the explanatory variables are: \( X_1 \), the number of visits to the supermarket in the six month period; \( X_2 \), the age of the customer; \( X_3 \), the number of members of the customers’ family. To find out more about this data set please see Atkinson and Riani (2006), JCGS.

**Usage**

`data("loyalty")`

**Format**

A data frame with 509 observations on the following 4 variables.

- **visits** the number of visits to the supermarket in the six month period
- **age** the age of the customer
- **family** the number of members of the customers’ family
- **amount_spent** the amount in euros spent at the shop over six months

**Details**

To find out more about this data set please see Atkinson and Riani (2006), JCGS.

**Source**

The data are themselves a random sample from a larger database. The sample of 509 observations is available at [http://www.riani.it/trimmed/](http://www.riani.it/trimmed/).

**References**

Examples

data(loyalty)

LXS_control

Description

Creates an object of class LXS_control to be used with the fsreg() function, containing various control parameters.

Usage

LXS_control(intercept = TRUE, lms, h, bdp, nsamp, rew = FALSE, conflev = 0, msg = TRUE, nocheck = FALSE, nomes = FALSE, plot = FALSE)

Arguments

intercept Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.

lms Criterion to use to find the initial subset to initialize the search (LMS, LTS with concentration steps, LTS without concentration steps or subset supplied directly by the user). The default value is 1 (Least Median of Squares is computed to initialize the search). On the other hand, if the user wants to initialize the search with LTS with all the default options for concentration steps then lms=2. If the user wants to use LTS without concentration steps, lms can be a scalar different from 1 or 2. If lms is a list it is possible to control a series of options for concentration steps (for more details see option lms inside LXS_control). If, on the other hand, the user wants to initialize the search with a prespecified set of units there are two possibilities:

1. lms can be a vector with length greater than 1 which contains the list of units forming the initial subset. For example, if the user wants to initialize the search with units 4, 6 and 10 then lms=c(4, 6, 10);
2. lms is a struct which contains a field named bsb which contains the list of units to initialize the search. For example, in the case of simple regression through the origin with just one explanatory variable, if the user wants to initialize the search with unit 3 then lms=list(bsb=3).

h The number of observations that have determined the least trimmed squares estimator, scalar. h is an integer greater or equal than p but smaller then n. Generally if the purpose is outlier detection h=[0.5*(n+p+1)] (default value). h can be smaller than this threshold if the purpose is to find subgroups of homogeneous observations. In this function the LTS/LMS estimator is used just to initialize the search.
Breakdown point. It measures the fraction of outliers the algorithm should resist. In this case any value greater than 0 but smaller or equal than 0.5 will do fine. If on the other hand the purpose is subgroups detection then bdp can be greater than 0.5. In any case however \( n^*(1-bdp) \) must be greater than \( p \). If this condition is not fulfilled an error will be given. Please specify \( h \) or \( bdp \) not both.

Number of subsamples which will be extracted to find the robust estimator, scalar. If \( nsamp=0 \) all subsets will be extracted. They will be \( (n \choose p) \). If the number of all possible subset is <1000 the default is to extract all subsets otherwise just 1000.

LXS reweighted - if \( rew=1 \) the reweighted version of LTS (LMS) is used and the output quantities refer to the reweighted version else no reweighting is performed (default).

Confidence level which is used to declare units as outliers, usually \( conflev=0.95, 0.975, 0.99 \) (individual alpha) or \( 1-0.05/n, 1-0.025/n, 1-0.01/n \) (simultaneous alpha). Default value is 0.975.

Controls whether to display or not messages on the screen If \( msg=1 \) (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.

Check input arguments, scalar. If \( nocheck=TRUE \) no check is performed on matrix \( y \) and matrix \( X \). Notice that \( y \) and \( X \) are left unchanged. In other words the additional column of ones for the intercept is not added. As default \( nocheck=FALSE \).

It controls whether to display or not on the screen messages about estimated time to compute LMS (LTS). If \( nomes=1 \) no message about estimated time to compute LMS (LTS) is displayed, else if \( nomes=0 \) (default), a message about estimated time is displayed.

Plot on the screen. Scalar. If \( plot=TRUE \) the plot of minimum deletion residual with envelopes based on \( n \) observations and the scatterplot matrix with the outliers highlighted is produced. If \( plot=2 \) the user can also monitor the intermediate plots based on envelope superimposition. If \( plot=FALSE \) (default) no plot is produced.

Details

Creates an object of class \( \text{FSR\_control} \) to be used with the \( \text{fsreg()} \) function, containing various control parameters.

Value

An object of class "LXS\_control" which is basically a \text{list} with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also as \( \text{Sreg\_control}, \text{MMreg\_control} \) and \( \text{FSR\_control} \)
Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~, data=hbk, method="LMS", control=LXS_control(h=56, nsamp=500, lms=2)))
## End(Not run)
```

---

### M5data

**Mixture M5 Data.**

**Description**

A bivariate data set obtained from three normal bivariate distributions with different scales and proportions 1:2:2. One of the components is strongly overlapping with another one. A 10% noise is added uniformly distributed in a rectangle containing the three normal components and not strongly overlapping with the three mixture components. A precise description of the M5 data set can be found in Garcia-escudero et al. (2008).

**Usage**

```r
data(M5data)
```

**Format**

A data frame with 2000 rows and 3 variables: The first two columns are the two variables. The last column is the true classification vector where symbol "0" stands for the contaminating data points.

**Source**


---

### malfwdplot

**Plots the trajectories of scaled Mahalanobis distances along the search**

**Description**

Plots the trajectories of scaled Mahalanobis distances along the forward search.
**Usage**

```r
malfwdplot(
  out,
  xlim, ylim, xlab, ylab, main, lwd, lty, col, cex.lab, cex.axis, subsize,
  fg.thresh, fg.unit, fg.labstep, fg.lwd, fg.lty, fg.col, fg.mark,
  fg.cex, bg.thresh, bg.style, standard, fground, bgground, tag,
  datatooltip, label, nameX, databrush, conflev, trace = FALSE,
  ...
)
```

**Arguments**

- `out`: An object of S3 class `fsmeda.object` returned by `fsmult` with `monitoring=TRUE` - a list containing the monitoring of minimum Mahalanobis distance.

The needed elements of `out` are

1. **MAL**: matrix containing the squared Mahalanobis distances monitored in each step of the forward search. Every row is associated with a unit (row of data matrix X).
2. **Un**: matrix containing the order of entry of each unit (necessary if datatooltip or databrush is selected).
3. **X**: The data matrix.
xlim Controls the x scale in the plot. xlim is a vector with two elements controlling minimum and maximum on the x-axis. Default is to use automatic scale.

ylim Controls the y scale in the plot. ylim is a vector with two elements controlling minimum and maximum on the y-axis. Default is to use automatic scale.

xlab A title for the x axis

ylab A title for the y axis, default is "Squared Mahalanobis distances".

main An overall title for the plot

lwd The line width, a positive number, defaulting to 1

lty The line type. Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.

col Colors to be used for the highlighted units

cex.lab The magnification to be used for x and y labels relative to the current setting of cex

cex.axis The magnification to be used for axis annotation relative to the current setting of cex

subsize Numeric vector containing the subset size with length equal to the number of columns of matrix of mahalanobis distances. The default value of subsize is (nrow(MAL) - ncol(MAL) + 1):nrow(MAL)

fg.thresh (alternative to fg.unit) numeric vector of length 1 or 2 which specifies the highlighted trajectories. If length(fg.thresh) == 1 the highlighted trajectories are those of units that throughout the search had at least once a mahalanobis distance greater than fg.thresh. The default value is fg.thresh=2.5. If length(fg.thresh) == 2 the highlighted trajectories are those of units that throughout the search had a mahalanobis distance at least once bigger than fg.thresh[2] or smaller than fg.thresh[1].

fg.unit (alternative to fg.thresh), vector containing the list of the units to be highlighted. If fg.unit is supplied, fg.thresh is ignored.

fg.labstep numeric vector which specifies the steps of the search where to put labels for the highlighted trajectories (units). The default is to put the labels at the initial and final steps of the search. fg.labstep="" means no label.

fg.lwd The line width for the highlighted trajectories (units). Default is 1.

fg.lty The line type for the highlighted trajectories (units). Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.

fg.col colors to be used for the highlighted units.

fg.mark Controls whether to plot highlighted trajectories as symbols. if fg.mark=TRUE each line is plotted using a different symbol else no marker is used (default).

fg.cex Controls the font size of the labels of the trajectories in foreground. If fg.cex=0 no labels will be shown - equivalent to fg.labstop="".
bg.thresh Numeric vector of length 1 or 2 which specifies how to define the unimportant trajectories. Unimportant trajectories will be plotted using a colormap, in greyish or will be hidden. If length(bg.thresh) == 1 the irrelevant units are those which always had a mahalanobis distance smaller than bg.thresh. If length(bg.thresh) == 2 the irrelevant units are those which always had a mahalanobis distance greater than bg.thresh[1] and smaller than bg.thresh[2]. The default is bg.thresh=2.5 if n > 100 and bg.thresh=-Inf if n <= 100 i.e. to treat all trajectories as important if n <= 100 and, if n > 100, to reduce emphasis only to trajectories having in all steps of the search a value of mahalanobis distance smaller than 2.5.

bg.style Specifies how to plot the unimportant trajectories as defined in option bg.thresh.
1. bg.style="faint": unimportant trajectories are plotted using a colormap.
2. bg.style="hide": unimportant trajectories are hidden.
3. bg.style="greyish": unimportant trajectories are displayed in a faint grey.

When n > 100 the default option is bg.style='faint'. When n <= 100 and bg.thresh == -Inf option bg.style is ignored. Remark: bground=" is equivalent to -Inf that is all trajectories are considered relevant.

standard MATLAB-style arguments - appearance of the plot in terms of xlim, ylim, axes labels and their font size style, color of the lines, etc.

fground MATLAB-style arguments - for the trajectories in foregound.
bground MATLAB-style arguments - for the trajectories in backgroud.
tag Plot handle. String which identifies the handle of the plot which is about to be created. The default is to use tag 'pl_resfwd'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.
datatooltip Interactive clicking. It is inactive if this parameter is set to FALSE. The default is datatooltip=TRUE, the user can select with the mouse an individual mahalanobis distance trajectory in order to have information about the corresponding unit, the associated label and the step of the search in which the unit enters the subset. If datatooltip is a list it may contain the following fields:
1. DisplayStyle determines how the data cursor displays. Possible values are 'datatip' and 'window' (default). 'datatip' displays data cursor information in a small yellow text box attached to a black square marker at a data point you interactively select. 'window' displays data cursor information for the data point you interactively select in a floating window within the figure.
2. SnapToDataVertex: specifies whether the data cursor snaps to the nearest data value or is located at the actual pointer position. Possible values are SnapToDataVertex='on' (default) and SnapToDataVertex='off'.
3. LineColor: controls the color of the trajectory selected with the mouse. It can be an RGB triplet of values between 0 and 1, or character vector indicating a color name. Note that a RGB vector can be conveniently chosen with our MATLAB class FSColor, see documentation.
4. **SubsetLinesColor**: enables to control the color of the trajectories of the units that are in the subset at a given step of the search (if `resfwdplot()` is applied to an object of class `fsreda.object`) or have a weight greater than 0.9 (if `resfwdplot()` is applied to an object of class `sregeda.object` or `mmregeda.object`). This can be done (repeatedly) with a left mouse click in proximity of the step of interest. A right mouse click will terminate the selection by marking with a up-arrow the step corresponding to the highlighted lines. The highlighted lines by default are in red, but a different color can be specified as RGB triplet or character of color name. Note that a RGB vector can be conveniently chosen with our MATLAB class FSColor, see documentation. By default **SubsetLinesColor**="", i.e. the modality is not active. Any initialization for **SubsetLinesColor** which cannot be interpreted as RGB vector will be converted to blue, i.e. **SubsetLinesColor** will be forced to be [0 0 1]. If **SubsetLinesColor** is not empty the previous option **LineColor** is ignored.

**label**  
Character vector containing the labels of the units (optional argument used when `datatooltip=TRUE`). If this field is not present labels row1, ..., rown will be automatically created and included in the pop up datatooltip window).

**nameX**  
Add variable labels in plot. A vector of strings of length p containing the labels of the variables in the dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically

**databrush**  
Interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is TRUE or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the scatterplot matrix. If the scatterplot matrix does not exist it is automatically created. In addition, brushed units can be highlighted in the monitoring Mahalanobis distance plot. Note that the window style of the other figures is set equal to that which contains the monitoring Mahalanobis distance plot. In other words, if the monitoring Mahalanobis distance plot is docked all the other figures will be docked too.

If `databrush=TRUE` the default selection tool is a rectangular brush and it is possible to brush only once (that is `persist="")`. If `databrush=list(...)`, it is possible to use all optional arguments of the MATLAB function `selectdataFS()` and the following optional arguments:

- **persist**: Persist is an empty value or a character containing 'on' or 'off'. The default value is `persist="", that is brushing is allowed only once. If `persist="on"` or `persis="off"` brushing can be done as many time as the user requires. If `persist='on'` then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If `persist='off'` every time a new brush is performed units previously brushed are removed.

- **label**: add labels of brushed units in the monitoring plot.

- **labeladd**: add labels of brushed units in the scatterplot matrix. If this option is '1', we label the units of the last selected group with the unit row index in the matrix X. The default value is `labeladd="", i.e. no label is added.

**conflev**  
Confidence interval for the horizontal bands. It can be a vector of different confidence level values, e.g. c(0.95, 0.99, 0.999). The confidence interval is based
On the $\chi^2$ distribution.

trace Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

Value

none

Author(s)

FSDA team, <valentin.todorov@chello.at>

References


Examples

## Not run:
## Produce monitoring MD plot with all the default options.
## Generate input structure for malfwdplot
n <- 100
p <- 4
Y <- matrix(rnorm(n*p), ncol=p)
Y[1:10,] <- Y[1:10,] + 4

out <- fsmult(Y, monitoring=TRUE, init=30)

## Produce monitoring MD plot with all the default options
malfwdplot(out)

## End(Not run)

malindexplot Plots the trajectory of minimum Mahalanobis distance (mmd)

Description

Plots the trajectory of minimum Mahalanobis distance (mmd)
Usage

malindexplot(
    out,
    p,
    xlab,
    ylab,
    main,
    nameX,
    conflev,
    numlab,
    tag,
    trace = FALSE,
    ...
)

Arguments

out  a numeric vector or an object of S3 class (one of `fsmult.object`, `smult.object` or `mmmult.object`) returned by one of the functions `fsmult` or `smult` or `mmmult` - a list containing the monitoring of minimum Mahalanobis distance
p   If `out` is a vector, `p` is the number of variables of the original data matrix which have been used to compute md.
xlab A title for the x axis
ylab A title for the y axis
main An overall title for the plot
nameX Add variable labels in the plot. A vector of strings of length `p` containing the labels of the variables of the original data matrix X. If it is empty (default) the sequence X1, ..., Xp will be created automatically
conflev confidence interval for the horizontal bands. It can be a vector of different confidence level values, e.g. c(0.95, 0.99, 0.999). The confidence interval is based on the chi^2 distribution.
numlab Number of points to be labeled in the plot. If `numlab` is a single number, e.g. `numlab=5`, the units with the k largest md are labelled in the plots. If `numlab` is a vector, the units indexed by the vector are labelled in the plot. Default is `numlab=5`, i.e. the 5 units units with the largest md are labelled. Use `numlab=NA` for no labelling.
tag Tag of the figure which will host the malindexplot. The default tag is `tag="pl_malindex"`. 
trace Whether to print intermediate results. Default is `trace=FALSE`.
...
 potential further arguments passed to lower level functions.

Value

none

Author(s)

FSDA team, <valentin.todorov@chello.at>
mdrplot

References


Examples

```r
## Not run:
## Mahalanobis distance plot of 100 random numbers.
## Numbers are from from the chisq with 5 degrees of freedom
malindexplot(rchisq(100, 5), 5)
## End(Not run)
```

**mdrplot**

*Plots the trajectory of minimum deletion residual (mdr)*

Description

Plots the trajectory of minimum deletion residual (mdr).

Usage

```r
mdrplot(out, quant = c(0.01, 0.5, 0.99), sign = TRUE,
   mplus1 = FALSE, envm,
   xlim, ylim, xlab, ylab, main,
   lwdenv, lwd, cex.lab, cex.axis,
   tag, datatooltip, label, nameX, namey, databrush,
   ...)
```

Arguments

- `out`: An object returned by FSReda() (see `FSReda_control`).
- `quant`:
  1. `mdr`: Minimum deletion residual. A matrix containing the monitoring of minimum deletion residual in each step of the forward search. The first column of mdr must contain the fwd search index.
  2. `Un`: (for FSR only) - matrix containing the order of entry in the subset of each unit (required only when datatooltip is true or databrush is not empty).
  3. `y`: a vector containing the response (required only when option databrush is requested).
  4. `X`: a matrix containing the explanatory variables (required only when option databrush is requested).
  5. `Bols`: (n-init+1) x (p+1) matrix containing the estimated beta coefficients monitored in each step of the robust procedure (required only when option databrush is requested and suboption multivarfit is requested).
Quantiles for which envelopes have to be computed. The default is to produce 1%, 50% and 99% envelopes. In other words the default is quant=c(0.01, 0.5, 0.99).

Whether to use MDR with sign: if sign=TRUE (default) we distinguish steps for which minimum deletion residual was associated with positive or negative value of the residual. Steps associated with positive values of mdr are plotted in black, while other steps are plotted in red.

Whether to plot the (m+1)-th order statistic. Specifies if it is necessary to plot the curve associated with (m+1)-th order statistic.

Sample size for drawing enevlopes. Specifies the size of the sample which is used to superimpose the envelope. The default is to add an envelope based on all the observations (size n envelope).

Control y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.

Control x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.

a title for the x axis

a title for the y axis

an overall title for the plot

Controls the width of the lines associated with the envelopes, default is lwdenv=1.

Controls the linewidth of the curve which contains the monitoring of minimum deletion residual.

The magnification to be used for x and y labels relative to the current setting of cex

The magnification to be used for axis annotation relative to the current setting of cex

Plot handle. String which identifies the handle of the plot which is about to be created. The default is to use tag 'pl_mdr'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.

If datatooltip is not empty the user can use the mouse in order to have information about the unit selected, the step in which the unit enters the search and the associated label. If datatooltip is a list, it is possible to control the aspect of the data cursor (see MATLAB function datacursormode() for more details or see the examples below). The default options areDisplayStyle="Window" and SnapToDataVertex="on".

Character vector containing the labels of the units (optional argument used when datatooltip=TRUE. If this field is not present labels row1, ..., rown will be automatically created and included in the pop up datatooltip window).

Add variable labels in plot. A vector of strings of length p containing the labels of the variables of the regression dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically
**namey**

Add response label. A string containing the label of the response.

**databrush**

Interactive mouse brushing. If `databrush` is missing or empty (default), no brushing is done. The activation of this option (databrush is a scalar or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the y|X plot, i.e. a matrix of scatter plots of y against each column of X, grouped according to the selection(s) done by brushing. If the plot y|X does not exist it is automatically created. In addition, brushed units are automatically highlighted in the minimum deletion residual plot if it is already open. The extension to the following plots will be available in future versions of the toolbox:

1. monitoring leverage plot;
2. maximum studentized residual;
3. $s^2$ and $R^2$;
4. Cook distance and modified Cook distance;
5. deletion t statistics.

Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If `databrush=TRUE` the default selection tool is a rectangular brush and it is possible to brush only once (that is `persist=""`).

If `databrush=list(...)`, it is possible to use all optional arguments of function `selectdataFS()` and the following optional argument:

1. **persist**. Persist is an empty value or a character containing 'on' or 'off'. The default value is `persist=""`, that is brushing is allowed only once. If `persist="on"` or `persist="off"` brushing can be done as many time as the user requires. If `persist='on'` then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If `persist='off'` every time a new brush is performed units previously brushed are removed.

2. **bivarfit**. Whether to superimpose bivariate least square lines on the plot (if `plot=TRUE`). This option adds one or more least squares lines, based on SIMPLE REGRESSION of y on Xi, to the plots of y|Xi. The default is `bivarfit=FALSE`: no line is fitted. If `bivarfit=1`, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. If `bivarfit=2`, two OLS lines are fitted: one to all points and another to the group of the genuine observations. The group of the potential outliers is not fitted. If `bivarfit=0` one OLS line is fitted to each group. This is useful for the purpose of fitting mixtures of regression lines. If `bivarfit='i1'` or `bivarfit='i2'`, etc. an OLS line is fitted to a specific group, the one with index 'i' equal to 1, 2, 3 etc. Again, useful in case of mixtures.

3. **multivarfit**. Whether to superimpose multivariate least square lines. This option adds one or more least square lines, based on MULTIVARIATE REGRESSION of y on X, to the plots of y|Xi. The default is `multivarfit=FALSE`: no line is fitted. If `multivarfit=1`, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. The line added to the scatter
plot \( y | X_i \) is \( \text{avconst} + C_i * X_i \), where \( C_i \) is the coefficient of \( X_i \) in the multivariate regression and \( \text{avconst} \) is the effect of all the other explanatory variables different from \( X_i \) evaluated at their centroid (that is overline(y)'C)). If \( \text{multivarfit}=2 \), same action as with \( \text{multivarfit}=1 \) but this time we also add the line based on the group of unselected observations (i.e. the normal units).

4. labeladd. Add outlier labels in plot. If \( \text{labeladd}=\text{TRUE} \), we label the outliers with the unit row index in matrices \( X \) and \( y \). The default value is \( \text{labeladd}=\text{FALSE} \), i.e. no label is added.

... potential further arguments passed to lower level functions.

Details
No details

Value
No value returned

Author(s)
FSDA team

Examples

```r
## Not run:
n <- 100
y <- rnorm(n)
X <- matrix(rnorm(n*4), nrow=n)
out <- fsreg(y~X, method="LTS")
out <- fsreg(y~X, method="FS", bsb=out$bs, monitoring=TRUE)
mdrplot(out)
## End(Not run)
```

---

**mmdplot**

*Plots the trajectory of minimum Mahalanobis distance (mmd)*

Description

Plots the trajectory of minimum Mahalanobis distance (mmd)
mmdplot

Usage

mmdplot(
  out,
  quant = c(0.01, 0.5, 0.99),
  mplus1 = FALSE,
  envm,
  lwd,
  lwdenv,
  xlim,
  ylim,
  tag,
  datatooltip,
  label,
  xlab,
  ylab,
  main,
  nameX,
  cex.lab,
  cex.axis,
  databrush,
  trace = FALSE,
  ...
)

Arguments

out An object of S3 class \textit{fsmeda.object} returned by \textit{fsmult} with monitoring=TRUE - a list containing the monitoring of minimum Mahalanobis distance
quant Quantiles for which envelopes have to be computed. The default is to produce 1%, 50% and 99% envelopes. In other words the default is quant=c(0.01, 0.5, 0.99).
mplus1 Weather to plot the (m+1)-th order statistic.
envm Sample size for drawing envelopes. Specifies the size of the sample which is used to superimpose the envelope. The default is to add an envelope based on all the observations (size n envelope).
lwd Controls the line width of the curve which contains the monitoring of minimum deletion residual.
lwdenv Controls the width of the lines associated with the envelopes. Default is lwdenv=1
xlim Control the x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.
ylim Control the y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.
tag Plot handle. String which identifies the handle of the plot which is about to be created. The default is tag='pl_mmd'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.
datatooltip
If datatooltip is not empty the user can use the mouse in order to have information about the unit selected, the step in which the unit enters the search and the associated label. If datatooltip is a list, it is possible to control the aspect of the data cursor (see MATLAB function `datacursormode()` for more details or see the examples below). The default options are `DisplayStyle="Window"` and `SnapToDataVertex="on"`.

label
Row labels. Character vector containing the labels of the units (optional argument used when `datatooltip=TRUE`). If this field is not present labels row1, ..., rown will be automatically created and included in the pop up datatooltip window.

xlab
A title for the x axis

ylab
A title for the y axis

main
An overall title for the plot

nameX
Add variable labels in the plot. A vector of strings of length p containing the labels of the variables of the original data matrix X. If it is empty (default) the sequence X1, ..., Xp will be created automatically

cex.lab
The magnification to be used for x and y labels relative to the current setting of cex

cex.axis
The magnification to be used for axis annotation relative to the current setting of cex

databrush
Interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is TRUE or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the scatterplot matrix. If the scatterplot matrix does not exist it is automatically created. In addition, brushed units can be highlighted in the monitoring MD plot. Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If `databrush=TRUE` the default selection tool is a rectangular brush and it is possible to brush only once (that is `persist=""`).

Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too

If `databrush=TRUE` the default selection tool is a rectangular brush and it is possible to brush only once (that is `persist=""`).

If `databrush=list(...)`, it is possible to use all optional arguments of the MATLAB function `selectdataF5()` and the following optional arguments:

- **persist**: This option can be an empty value or a character containing 'on' or 'off'. The default value is `persist=""`, that is brushing is allowed only once. If `persist="on"` or `persist="off"` brushing can be done as many time as the user requires. If `persist='on'` then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If `persist='off'` every time a new brush is performed units previously brushed are removed.
mmdrsplot

• labeladd: add labels of brushed units in the scatterplot matrix. If this option is '1', we label the units of the last selected group with the unit row index in the matrix X. The default value is labeladd="", i.e. no label is added.

trace

Whether to print intermediate results. Default is trace=FALSE.

...

potential further arguments passed to lower level functions.

Value

none

Author(s)

FSDA team, <valentin.todorov@chello.at>

References


Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsmult(hbk[,1:3], monitoring=TRUE))
mmdplot(out)
## End(Not run)
```

mmdrsplot

Plots the trajectories of minimum Mahalanobis distances from different starting points

Description

Plots the trajectories of minimum Mahalanobis distances from different starting points

Usage

```r
mmdrsplot(
  out,
  quant = c(0.01, 0.5, 0.99),
  envm,
  lwd,
  lwdenv,
  xlim,
  ylim,
  tag,
)```
Arguments

out An object of S3 class `fsmmmdrs.object` returned by `fsmmmdrs` - a list containing the following elements:

- `mmdrs` = a matrix of size \((n-ninit)-(nsimul+1)\) containing the monitoring of minimum Mahalanobis distance in each step of the forward search for each of the nsimul random starts. The first column of mmdrs must contain the forward search index. This matrix can be created using function `fsmmmdrs`.
- `BBrs` = 3D array of size \(n\)-by-\(n\)-(init)-by-\(nsimul\) containing units forming subset for each random start. This field is necessary if datatooltip is true or databrush is not empty.
- `X` = \(n\)-by-\(v\) matrix containing the original data matrix. This field is necessary if datatooltip is true or databrush is not empty.

quant Quantiles for which envelopes have to be computed. The default is to produce 1%, 50% and 99% envelopes. In other words the default is `quant=c(0.01, 0.5, 0.99)`.

envm Sample size for drawing envelopes. Specifies the size of the sample which is used to superimpose the envelope. The default is to add an envelope based on all the observations (size \(n_{envelope}\)).

lwd Controls the linewidth of the curve which contains the monitoring of minimum deletion residual.

lwdenv line width: a scalar which controls the width of the lines associated with the envelopes. Default is `lwdenv=1`

xlim Control the x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.

ylim Control the y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.

tag Plot handle. String which identifies the handle of the plot which is about to be created. The default is `tag='pl_mmdrs'`. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the
new plot overwrites the existing one in the same window else a new window is created.

datatooltip If datatooltip is not empty the user can use the mouse in order to have information about the unit selected, the step in which the unit enters the search and the associated label. If datatooltip is a list, it is possible to control the aspect of the data cursor (see MATLAB function datacursormode() for more details or see the examples below). The default options are DisplayStyle="Window" and SnapToDataVertex="on".

label Row labels. Character vector containing the labels of the units (optional argument used when datatooltip=TRUE. If this field is not present labels row1, ..., rown will be automatically created and included in the pop up datatooltip window).

xlab A title for the x axis

ylab A title for the y axis

envlab wheather to label the envelopes. If envlab is true (default) labels of the confidence envelopes which are used are added on the y axis.

main An overall title for the plot

nameX Add variable labels in the plot. A vector of strings of length p containing the labels of the variables of the original data matrix X. If it is empty (default) the sequence X1, ..., Xp will be created automatically

cex.lab The magnification to be used for x and y labels relative to the current setting of cex

cex.axis The magnification to be used for axis annotation relative to the current setting of cex

databrush Interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is TRUE or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the scatterplot matrix. If the scatterplot matrix does not exist it is automatically created. In addition, brushed units can be highlighted in the monitoring MD plot. Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist="").

Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist="").

If databrush=list(...), it is possible to use all optional arguments of the MATLAB function selectdataFS() and the following optional arguments:

1. persist. Persist is an empty value or a character containing 'on' or 'off'. The default value is persist="", that is brushing is allowed only once. If persist="on" or persist="off" brushing can be done as many times as the user requires. If persist='on' then the unit(s) currently brushed are
added to those previously brushed. It is possible, every time a new brushing
is done, to use a different color for the brushed units. If persist='off'
every time a new brush is performed units previously brushed are removed.

`scaled` Wheather to use scaled or unscaled envelopes. If `scaled=TRUE` the envelopes
are produced for scaled Mahalanobis distances (no consistency factor is applied)
else the traditional consistency factor is applied. Default is `scaled=FALSE`

`trace` Whether to print intermediate results. Default is `trace=FALSE`.

... potential further arguments passed to lower level functions.

Value

`none`

Author(s)

FSDA team, <valentin.todorov@chello.at>

References

Atkinson, A.C., Riani, M. and Cerioli, A. (2004), 'Exploring multivariate data with the forward
search', Springer Verlag, New York.

Examples

```r
## Not run:
data(hbk, package="robustbase")
out <- fsmmdrs(hbk[,1:3])
mmdrsplot(out)
## End(Not run)
```

### mmmult

Computes MM estimators in multivariate analysis with auxiliary S-scale

Description

Computes MM estimators in multivariate analysis with auxiliary S-scale

Usage

```r
mmmult(
  x,
  monitoring = FALSE,
  plot = FALSE,
  eff,
  conflev = 0.975,
  nocheck = FALSE,
```
Arguments

\textbf{x} \quad \text{An n x p data matrix (n observations and p variables). Rows of x represent observations, and columns represent variables. Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.}

\textbf{monitoring} \quad \text{Whether to perform monitoring of Mahalanobis distances and other specific quantities}

\textbf{plot} \quad \text{Plots the Mahalanobis distances against index number. If plot=FALSE (default) or plot=0 no plot is produced. The confidence level used to draw the confidence bands for the MD is given by the input option conflev. If conflev is not specified a nominal 0.975 confidence interval will be used. If plot=2 a scatter plot matrix with the outliers highlighted is produced. If plot is a list it may contain the following fields:}

- \textbf{labeladd} If labeladd=1, the outliers in the spm are labelled with the unit row index. The default value is labeladd="", i.e. no label is added
- \textbf{nameY} character vector containing the labels of the variables. As default value, the labels which are added are Y1, ...Yp.

\textbf{eff} \quad \text{Defining the nominal efficiency (i.e. a number between 0.5 and 0.99). The default value is eff=0.95.}

\textbf{conflev} \quad \text{Confidence level which is used to declare units as outliers (scalar). Usually conflev=0.95, conflev=0.975 or conflev=0.99 (individual alpha) conflev=1-0.05/n, conflev=1-0.025/n or conflev=1-0.01/n (simultaneous alpha). Default value is conflev=0.975.}

\textbf{nocheck} \quad \text{It controls whether to perform checks on matrix Y. If nocheck=TRUE, no check is performed.}

\textbf{trace} \quad \text{Whether to print intermediate results. Default is trace=FALSE.}

... potential further arguments passed to lower level functions.

Details

This function follows the lines of MATLAB/R code developed during the years by many authors. For more details see http://www.econ.kuleuven.be/public/NDBAE06/programs/ and the R package \texttt{CovMMest} The core of these routines, e.g. the resampling approach, however, has been completely redesigned, with considerable increase of the computational performance.

Value

Depending on the input parameter monitoring, one of the following objects will be returned:

1. \texttt{mmmult.object}
2. \texttt{mmmulteda.object}
Author(s)
FSDA team, <valentin.todorov@chello.at>

References

Examples

## Not run:
data(hbk, package="robustbase")
(out <- mmmult(hbk[,1:3]))
class(out)
summary(out)

## Generate contaminated data (200,3)
n <- 200
p <- 3
set.seed(123456)
X <- matrix(rnorm(n*p), nrow=n)
Xcont <- X

out1 <- mmmult(Xcont, trace=TRUE) # no plots (plot defaults to FALSE)
names(out1)

## plots is a list: the spm shows the labels of the outliers.
(out1 <- mmmult(Xcont, trace=TRUE, plot=list(labeladd="1")))

## plots is a list: the spm uses the variable names provided by 'nameY'.
(out1 <- mmmult(Xcont, trace=TRUE, plot=list(nameY=c("A", "B", "C"))))

## mmmult() with monitoring
(out2 <- mmmult(Xcont, monitoring=TRUE, trace=TRUE))
names(out2)

## Forgery Swiss banknotes examples.
data(swissbanknotes)
(out1 <- mmmult(swissbanknotes[101:200,], plot=TRUE))
(out1 <- mmmult(swissbanknotes[101:200,], plot=list(labeladd="1")))

## End(Not run)

### Description of mmmult.object Objects

**Description**

An object of class `mmmult.object` holds information about the result of a call to `mmmult`.

**Value**

The object itself is basically a list with the following components:

- `loc`:
  - p-by-1 vector containing MM estimate of location.
- `shape`:
  - p-by-p matrix with MM estimate of the shape matrix.
- `cov`:
  - matrix with MM estimate of the covariance matrix. Remark: `covariance = auxscale^2 * shape`.
- `weights`:
  - A vector containing the estimates of the weights.
- `outliers`:
  - A vector containing the list of the units declared as outliers using confidence level specified in input scalar `conflev`.
- `Sloc`:
  - A vector with S estimate of location.
- `Sshape`:
  - A matrix with S estimate of the shape matrix.
- `Scov`:
  - A matrix with S estimate of the covariance matrix.
- `auxscale`:
  - S estimate of the scale.
- `md`:
  - n-by-1 vector containing the estimates of the robust Mahalanobis distances (in squared units).
- `conflev`:
  - Confidence level that was used to declare outliers.
- `X`:
  - the data matrix X

The object has class "mmmult".

**Examples**

```r
## Not run:
data(hbk, package="robustbase")
(out <- mmmult(hbk[,1:3]))
class(out)
summary(out)

## End(Not run)
```
mmulteda.object

Description

An object of class `mmulteda.object` holds information about the result of a call to `mmult` with `monitoring=TRUE`.

Value

The object itself is basically a list with the following components:

- **Loc**: length(\(\mathbf{eff}\))-by-\(p\) matrix containing MM estimate of location for each value of \(\mathbf{eff}\).
- **Shape**: \(p\)-by-\(p\)-by-length(\(\mathbf{eff}\)) 3D array containing robust estimate of the shape for each value of \(\mathbf{eff}\). Remark: \(\det(\mathbf{shape})=1\).
- **Scale**: length(\(\mathbf{eff}\)) vector containing robust estimate of the scale for each value of \(\mathbf{eff}\).
- **Cov**: \(p\)-by-\(p\)-by-length(\(\mathbf{eff}\)) 3D array containing robust estimate of covariance matrix for each value of \(\mathbf{eff}\). Note that \(\text{scale}(i)^2 \times \text{shape}[,,i] = \) robust estimate of covariance matrix.
- **Bs**: \((p+1)\)-by-length(\(\mathbf{eff}\)) matrix containing the units forming best subset for each value of \(\mathbf{eff}\).
- **MAL**: \(n\)-by-length(\(\mathbf{eff}\)) matrix containing the estimates of the robust Mahalanobis distances (in squared units) for each value of \(\mathbf{eff}\).
- **Outliers**: \(n\)-by-length(\(\mathbf{eff}\)) matrix containing flags for the outliers. Boolean matrix containing the list of the units declared as outliers for each value of \(\mathbf{eff}\) using confidence level specified in input scalar `conflev`.
- **Weights**: \(n \times \text{length}(\mathbf{eff})\) matrix containing the weights for each value of \(\mathbf{eff}\).
- **conflev**: Confidence level that was used to declare outliers.
- **singsub**: Number of subsets without full rank. Notice that \(\text{singsub} > 0.1 \times \text{(number of subsamples)}\) produces a warning.
- **eff**: vector which contains the values of \(\mathbf{eff}\) which have been used.
- **X**: the data matrix \(X\).

The object has class "mmulteda".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- mmmult(hbk[,1:3], monitoring=TRUE))
class(out)
summary(out)

## End(Not run)
```
Description

An object of class \texttt{mmreg.object} holds information about the result of a call to \texttt{fsreg} with \texttt{method=}"MM".

Value

The object itself is basically a \texttt{list} with the following components:

- \texttt{beta}: p-by-1 vector containing the MM estimate of regression coefficients.
- \texttt{auxscale}: scalar, S estimate of the scale (or supplied external estimate of scale, if option \texttt{InitialEst} is not empty).
- \texttt{residuals}: residuals.
- \texttt{fittedvalues}: fitted values.
- \texttt{weights}: n x 1 vector. Weights assigned to each observation.
- \texttt{Sbeta}: p x 1 vector containing S estimate of regression coefficients (or supplied initial external estimate of regression coefficients, if option \texttt{InitialEst} is not empty).
- \texttt{Ssingsub}: Number of subsets without full rank in the S preliminary part. Notice that \texttt{out.singsub} > 0.1*(number of subsamples) produces a warning.
- \texttt{outliers}: kx1 vector containing the list of the k units declared as outliers or NULL if the sample is homogeneous.
- \texttt{conflev}: Confidence level which is used to declare units as outliers. Usually \texttt{conflev=0.95, 0.975, 0.99} (individual alpha) or \texttt{conflev=1-0.05/n, 1-0.025/n, 1-0.01/n} (simultaneous alpha). Default value is 0.975.
- \texttt{rhofunc}: Specifies the rho function which has been used to weight the residuals. If a different rho function is specified for S and MM loop then instead of \texttt{rhofunc} we will have \texttt{rhofuncS} and \texttt{rhofuncMM}.
- \texttt{rhofuncparam}: Vector which contains the additional parameters for the specified rho function which has been used. For hyperbolic rho function the value of k =sup CVC. For Hampel rho function the parameters a, b and c. If a different rho function is specified for S and MM loop then instead of \texttt{rhofuncparam} we will have \texttt{rhofuncparamS} and \texttt{rhofuncparamMM}.
- \texttt{X}: the data matrix X
- \texttt{y}: the response vector y

The object has class "mmreg".
Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="MM"))
class(out)
summary(out)
## End(Not run)
```

---

**mmregeda.object**  
*Description of mmregeda Objects*

**Description**

An object of class `mmregeda.object` holds information about the result of a call to `fsreg` when method="MM" and monitoring=TRUE.

**Value**

The object itself is basically a `list` with the following components:

- `auxscale` scalar, S estimate of the scale (or supplied external estimate of scale, if option InitialEst is not empty).
- `Beta` p x length(eff) matrix containing MM estimate of regression coefficients for each value of eff.
- `RES` n x length(eff) matrix containing the monitoring of scaled residuals for each value of eff.
- `Weights` n x length(eff) matrix containing the estimates of the weights for each value of eff.
- `Outliers` Boolean matrix containing the list of the units declared as outliers for each value of eff using confidence level specified in input scalar conflev.
- `conflev` Confidence level which is used to declare units as outliers. Remark: conflev will be used to draw the horizontal line (confidence band) in the plot.
- `Singsub` Number of subsets without full rank. Notice that Notice that singsub > 0.1*(number of subsamples) produces a warning
- `rhofunc` string identifying the rho function which has been used.
- `rhofuncparam` vector which contains the additional parameters for the specified rho function which have been used. For hyperbolic rho function the value of k =sup CVC. For Hampel rho function the parameters a, b and c.
- `eff` vector containing the value of eff which have been used.
- `X` the data matrix X
- `y` the response vector y

The object has class "mmregeda".
Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="MM", monitoring=TRUE))
class(out)
summary(out)
## End(Not run)
```

**MMregeda_control**

*Creates an MMregeda_control object*

**Description**

Creates an object of class `MMregeda_control` to be used with the `fsreg()` function, containing various control parameters.

**Usage**

```r
MMregeda_control(intercept = TRUE, InitialEst, Soptions, eff, effshape,
rhofunc = c("bisquare", "optimal", "hyperbolic", "hampel", "mdpd", "AS"),
rhofuncparam, refsteps = 3, tol = 1e-07, conflev, nocheck = FALSE, plot = FALSE)
```

**Arguments**

- `intercept`: Indicator for constant term. Scalar. If `intercept=TRUE`, a model with constant term will be fitted (default), else, no constant term will be included.

- `InitialEst`: Starting values of the MM-estimator, a list with the following elements: `loc`, a $p \times 1$ vector, location vector estimate and `scale`, a scaler, estimate of the scale. If empty (default) the program will use S estimators. In this last case it is possible to specify the options given in function `Sreg`.

- `Soptions`: Options to pass to `Sreg`, an `Sreg_control` object. The options are: `Srhofunc`, `Snsamp`, `Srefsteps`, `Sreftol`, `Srnodstepsbestr`, `Sreftolbestr`, `Sminsctol`, `Sbestr`. See function `Sreg_control` for more details on these options.

- `eff`: Vector defining nominal efficiency (i.e. a series of numbers between 0.5 and 0.99). The default value is the sequence `seq(0.5, 0.99, 0.01)`

- `effshape`: Location or scale efficiency. If `effshape=1` efficiency refers to shape efficiency else (default) efficiency refers to location efficiency.

- `rhofunc`: Specifies the rho function which must be used to weight the residuals. Possible values are 'bisquare' 'optimal' 'hyperbolic' 'hampel'.
1. 'bisquare' uses Tukey's rho and psi functions. See TBrho and TBpsi.
2. 'optimal' uses optimal rho and psi functions. See OPTrho and OPTpsi.
3. 'hyperbolic' uses hyperbolic rho and psi functions. See HYPrho and HYPpsi.
4. 'hampel' uses Hampel rho and psi functions. See HArho and HApssi.

The default is 'bisquare'.

```r
rhofuncparam
```

Additional parameters for the specified rho function. For hyperbolic rho function it is possible to set the value of \( k = \sup CVC \) (the default value of \( k \) is 4.5).

For Hampel rho function it is possible to define parameters \( a, b \) and \( c \) (the default values are \( a=2, b=4, c=8 \)).

```r
refsteps
```

Number of refining iterations in each subsample (default is \( refsteps=3 \)). \( refsteps = 0 \) means "raw-subsampling" without iterations.

```r
tol
```

Scalar controlling tolerance in the MM loop. The default value is \( tol=1e-6 \).

```r
conflev
```

Confidence level which is used to declare units as outliers. Usually \( conflev=0.95, 0.975, 0.99 \) (individual alpha) or \( conflev=1-0.05/n, 1-0.025/n, 1-0.01/n \) (simultaneous alpha). Default value is 0.975.

```r
nocheck
```

Check input arguments, scalar. If \( nocheck=TRUE \) no check is performed on matrix \( y \) and matrix \( X \). Notice that \( y \) and \( X \) are left unchanged. In other words the additional column of ones for the intercept is not added. As default \( nocheck=FALSE \).

```r
plot
```

Plot on the screen. Scalar. If \( plots=TRUE \) the plot of minimum deletion residual with envelopes based on \( n \) observations and the scatterplot matrix with the outliers highlighted is produced. If \( plots=2 \) the user can also monitor the intermediate plots based on envelope superimposition. If \( plots=FALSE \) (default) no plot is produced.

Details

Creates an object of class \texttt{MMregeda\_control} to be used with the \texttt{fsreg()} function, containing various control parameters.

Value

An object of class "\texttt{MMregeda\_control}" which is basically a \texttt{list} with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also as \texttt{FSR\_control}, \texttt{Sreg\_control}, \texttt{MMreg\_control} and \texttt{LXS\_control}
MMreg_control

Description

Creates an object of class MMreg_control to be used with the fsreg() function, containing various control parameters for calling the MATLAB function MMreg().

Usage

MMreg_control(intercept = TRUE, InitialEst, eff, effshape, rhofunc = c("bisquare", "optimal", "hyperbolic", "hampel", "mdpd", "AS"), rhofuncparam, refsteps = 3, tol = 1e-07, conflev, nocheck = FALSE, Smsg = TRUE, plot = FALSE)

Arguments

intercept Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.

InitialEst Starting values of the MM-estimator, a list with the following elements: 1oc, a $p x 1$ vector, location vector estimate and scale, a scaler, estimate of the scale. If empty (default) the program will use S estimators. In this last case it is possible to specify the options given in function Sreg.

eff Scalar defining nominal efficiency (i.e. a number between 0.5 and 0.99). The default value is 0.95.

effshape Location or scale efficiency. If effshape=1 efficiency refers to shape efficiency else (default) efficiency refers to location efficiency.

rhofunc Specifies the rho function which must be used to weight the residuals. Possible values are 'bisquare' 'optimal' 'hyperbolic' 'hampel'.
1. 'bisquare' uses Tukey's rho and psi functions. See TBrho and TBpsi.
2. 'optimal' uses optimal rho and psi functions. See OPTrho and OPTpsi.
3. 'hyperbolic' uses hyperbolic rho and psi functions. See HYPrho and HYPpsi.
4. 'hampel' uses Hampel rho and psi functions. See HARho and HAPsi.

The default is 'bisquare'.

Examples

## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="MM", monitoring=TRUE, control=MMregeda_control(eff=seq(0.75, 0.99, 0.01))))
## End(Not run)
rhofuncparam  Additional parameters for the specified rho function. For hyperbolic rho function it is possible to set up the value of \( k = \sup CVC \) (the default value of \( k \) is 4.5).
For Hampel rho function it is possible to define parameters \( a, b \) and \( c \) (the default values are \( a=2, b=4, c=8 \)).

refsteps  Number of refining iterations in each subsample (default is \( \text{refsteps}=3 \)). \( \text{refsteps}=0 \) means "raw-subsampling" without iterations.

tol  Scalar controlling tolerance in the MM loop. The default value is \( \text{tol}=1e-6 \)

conflev  Confidence level which is used to declare units as outliers. Usually \( \text{conflev}=0.95, 0.975, 0.99 \) (individual alpha) or \( \text{conflev}=1-0.05/n, 1-0.025/n, 1-0.01/n \) (simultaneous alpha). Default value is 0.975

nocheck  Check input arguments, scalar. If \( \text{nocheck}=TRUE \) no check is performed on matrix \( y \) and matrix \( X \). Notice that \( y \) and \( X \) are left unchanged. In other words the additional column of ones for the intercept is not added. As default \( \text{nocheck}=FALSE \).

Smsg  Controls whether to display or not messages on the screen If \( \text{Smsg}=TRUE \) (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.

plot  Plot on the screen. Scalar. If \( \text{plots}=TRUE \) the plot of minimum deletion residual with envelopes based on \( n \) observations and the scatterplot matrix with the outliers highlighted is produced. If \( \text{plots}=2 \) the user can also monitor the intermediate plots based on envelope superimposition. If \( \text{plots}=FALSE \) (default) no plot is produced.

Details

Creates an object of class \texttt{MMreg\_control} to be used with the \texttt{fsreg()} function, containing various control parameters.

Value

An object of class "\texttt{MMreg\_control}" which is basically a \texttt{list} with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also as \texttt{FSR\_control}, \texttt{MMreg\_control} and \texttt{LXS\_control}

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="MM", control=MMreg_control(eff=0.99, rhofunc="optimal")))
## End(Not run)
```
**multiple_regression**

*Multiple regression data showing the effect of masking (Atkinson and Riani, 2000).*

**Description**

There are 60 observations on a response $y$ with the values of three explanatory variables. The scatter plot matrix of the data shows $y$ increasing with each of $x_1$, $x_2$ and $x_3$. The plot of residuals against fitted values shows no obvious pattern. However the FS finds that there are 6 masked outliers.

**Usage**

```r
data(multiple_regression)
```

**Format**

A data frame with 60 rows and 4 variables The variables are as follows:

- X1
- X2
- X3
- y


**mussels**

*Mussels data.*

**Description**

These data, introduced by Cook and Weisberg (1994), consist of 82 observations on horse mussels from New Zealand. The variables are shell length, width, height, mass and muscle mass.

**Usage**

```r
data(mussels)
```

**Format**

A data frame with 82 rows and 5 variables
myrng

Set seed for the MATLAB random number generator

Description

Initializes the MATLAB random generator

Usage

myrng(seed)

Arguments

seed a single value, interpreted as an integer

Value

Integer, the seed value with which the MATLAB random number generator was initialized.

Author(s)

FSDA team, <valentin.todorov@chello.at>

Examples

## Not run:
data(wool)
XX <- wool
y <- XX[, ncol(XX)]
X <- XX[, 1:(ncol(XX)-1), drop=FALSE]
seed <- myrng() # initialized the RNG and keep the seed
myrng(seed) # repeat the computations with the same seed
(out2 <- fsreg(X, y, method="LTS")

all.equal(out1, out2)

## End(Not run)
The poison data (by Box and Cox, 1964) are about the time to death of animals in a 3 x 4 factorial experiment with four observations at each factor combination. There are no outliers or influential observations that cannot be reconciled with the greater part of the data by a suitable transformation.

Usage

data(poison)

Format

A data frame with 48 rows and 7 variables: six explanatory and one response variable.

Source


Examples

data(poison)
head(poison)

psifun

Finds the tuning constant(s) associated to the supplied breakdown point or asymptotic efficiency for different psi functions

Description

Finds the tuning constant(s) associated to the supplied breakdown point or asymptotic efficiency or computes breakdown point and efficiency associated with the supplied constant(s) for the following psi functions: TB=Tukey biweight, HA=Hampel, HU=Huber, HYP=Hyperbolic, OPT=Optimal, PD=mdpd.
Usage

psifun(
  u = vector(mode = "double", length = 0),
  p = 1,
  fun = c("TB", "bisquare", "biweight", "HA", "hampel", "HU", "huber", "HYP",
           "hyperbolic", "OPT", "optimal", "PD", "mdpd"),
  bdp,
  eff,
  const,
  param,
  trace = FALSE
)

Arguments

u optional vector containing scaled residuals or Mahalanobis distances for the n units of the sample. If not provided, rho, psi, psider, psix and weights are not computed
p number of variables (p=1 for regression)
fun psi function class. One of TB, HA, HU, HP, OPT or PD.
bdp requested breakdown point
eff requested asymptotic efficiency
const tuning constant c
param additional parameters
trace whether to print intermediate results. Default is trace=FALSE.

Value

A list will be returned containing the following elements:

1. class: link function which has been used. Possible values are 'bisquare', 'optimal', 'hyperbolic', 'hampel', 'huber' or 'mdpd'
2. bdp: breakdown point
3. eff: asymptotic efficiency
4. c1: consistency factor (and other parameters) associated to required breakdown point or nominal efficiency.
5. kc1: Expectation of rho associated with c1 to get a consistent estimator at the model distribution kc1 = E(rho) = sup(rho)*bdp
6. rho: vector of length n which contains the rho function associated to the residuals or Mahalanobis distances for the n units of the sample. Empty if u is not provided.
7. psi: vector of length n which contains the psi function associated with the residuals or Mahalanobis distances for the n units of the sample. Empty if u is not provided.
8. psider: vector of length n which contains the derivative of the psi function associated with the residuals or Mahalanobis distances for the n units of the sample. Empty if u is not provided.
9. \texttt{psix}: vector of length \( n \) which contains the psi function multiplied by \( u \) associated with the residuals or Mahalanobis distances for the \( n \) units of the sample. Empty if \( u \) is not provided.

10. \texttt{wei}: vector of length \( n \) which contains the weights associated with the residuals or Mahalanobis distances for the \( n \) units of the sample. Empty if \( u \) is not provided.

\textbf{Author(s)}

FSDA team, <valentin.todorov@chello.at>

\textbf{References}


\textbf{Examples}

```r
## Not run:
## Find c for given bdp for the Tukey biweight function
## The constant c associated to a breakdown point of
## 50 percent in regression is
##  c=1.54764980928226
##               psifun(bdp=0.5)
##               psifun(c=1.54764980928226)

## Find c for given bdp for the Hampel function
##               psifun(bdp=0.5, fun="hampel")

## Plot Huber rho function.
##               x <- seq(-3, 3, 0.001)
##               c <- 1.345;
##               HUc1 <- psifun(u=x, p=1, fun="HU", const=c)
##               rhoHU <- HUc1$rho
##               plot(x, rhoHU, type="l", lty="solid", lwd=2, col="blue",
##               xlab="u", ylab="rho \(\{u,1.345\}\)"
##               ylim=c(0.16, 4.5))
##               lines(x, x^2/2, type="l", lty="dotted", lwd=1.5, col="red")
##               legend(-1, 4.6, legend=c("Huber rho function", "u^2/2"),
##               lty=c("solid", "dotted"), lwd=c(2,1.5), col=c("blue", "red"))
```
regspmplot

Interactive scatterplot matrix for regression

Description

Produces an interactive scatterplot of the response y against each variable of the predictor matrix X.

Usage

regspmplot(
  y,
  X,
  group,
  plot,
  namey,
  nameX,
  col,
  cex,
  pch,
  labeladd,
  legend,
  xlim,
  ylim,
  tag,
  datatooltip,
  databrush,
  subsize,
  selstep,
  selunit,
  trace = FALSE,
  ...
)
Arguments

**y**
response variable or an object containing the response, the predictors and possibly other variable resulting from monitoring of regression.

If `y` is a vector, a data matrix `X` must be present as an argument. If `y` is a list containing just `y` and `X`, the call is equivalent to `regspmplot(y, X)`. Otherwise `y` must be an object of S3 class `fsreda.object` returned by `fsreg` with `monitoring=TRUE` - a list containing the monitoring along a search.

**X**
Predictor variables. Data matrix of explanatory variables (also called 'regressors') of dimension `n` by `p` if the argument `y` is a vector. The rows of `X` represent observations, and the columns represent variables.

**group**
Grouping variable. Vector with `n` elements. Specifies a grouping variable defined as a categorical variable (factor), numeric, or array of strings, or string matrix, and it must have the same number of rows as `X`. This grouping variable determines the marker and color assigned to each point. Remark: if `group` is used to distinguish a set of outliers from a set of good units, the id number for the outliers should be the larger (see optional field `labeladd` of parameter `plot` for details).

**plot**
This option controls the names which are displayed in the margins of the scatterplot matrix as well as the labels of the legend. If `plot=FALSE`, then `namey`, `nameX` and `labeladd` are both set to the empty string (default), and no label and no name is added to the plot. If `plot=TRUE` the names `y`, and `X1,..., Xp` are added to the margins of the the scatter plot matrix else nothing is shown. If `plot` is a list, it is possible to control not only the names but also, point labels, colors and symbols. More precisely list `plot` may contain the following elements:

1. `labeladd` - see parameter `labeladd`
2. `namey` - a character string containing the response variable name. See parameter `namey`.
3. `nameX` - a vector of character strings containing the labels of the explanatory variables. As default value, the labels which are added are `Y1, ..., Yp`. See parameter `nameX`.
4. `clr` - see parameter `col`
5. `sym` - see parameter `pch`
6. `siz` - see parameter `cex`
7. `doleg` - see parameter `legend`
8. `xlimx` - see parameter `xlim`
9. `ylimy` - see parameter `ylim`

**namey**
a character string with the name of the response variable

**nameX**
a vector of character strings with the names of the explanatory variables

**col**
color specification for the data point. Can be different for each group. By default, the order of the colors is `blue`, `red`, `black`, `magenta`, `green`, `cyan` and `yellow`.

**cex**
the size of the symbols used for plotting. By default `cex=1` the symbol size depends on the number of plots and the size of the figure window. Values larger than 1 will increase the size and values smaller than 1 will decrease the size.
specification of the symbols to use. For example, if there are three groups, and
pch=c(1, 3, 4), the first group will be plotted with a circle, the second with a
plus, and the third with a 'x' (see ?pch or ?points for a list of symbols. NOTE:
not all symbols available in R can be mapped to the symbols in MATLAB.

labeladd
logical, controls whether the elements belonging to the last group in the scat-
terplot matrix are labelled with their unit row index or their rowname. The row-
name is taken from the parameter label or if it is missing, from the sequence
1:n. The default value is labeladd=FALSE, i.e. no label is added.

legend
logical, controls where a legend is shown or not.

xlim
x limits. A vector with two elements controlling minimum and maximum on the
x axis. By default automatic scale is used.

ylim
y limits. A vector with two elements controlling minimum and maximum on the
y axis. By default automatic scale is used.

tag
Plot handle. String which identifies the handle of the plot which is about to be
created. The default is tag='pl_mmd'. Notice that if the program finds a plot
which has a tag equal to the one specified by the user, then the output of the
new plot overwrites the existing one in the same window else a new window is
created.

datatooltip
If datatooltip is not empty the user can use the mouse in order to have infor-
mation about the unit selected, the step in which the unit enters the search and
the associated label. If datatooltip is a list, it is possible to control the aspect of
the data cursor (see MATLAB function datacursormode() for more details or
see the examples below). The default options are DisplayStyle="Window" and
SnapToDataVertex="on".

databrush
Interactive mouse brushing. If databrush is missing or empty (default), no brush-
ing is done. The activation of this option (databrush is TRUE or a list) enables
the user to select a set of trajectories in the current plot and to see them high-
lighted in the scatterplot matrix. If the scatterplot matrix does not exist it is
automatically created. In addition, brushed units can be highlighted in the mon-
itoring MD plot. Note that the window style of the other figures is set equal to
that which contains the monitoring residual plot. In other words, if the moni-
toring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is
possible to brush only once (that is persist="").

Note that the window style of the other figures is set equal to that which contains
the monitoring residual plot. In other words, if the monitoring residual plot is
docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is
possible to brush only once (that is persist="").

If databrush=list(...), it is possible to use all optional arguments of the
MATLAB function selectdataFS() and the following optional arguments:

• persist: This option can be an empty value or a character containing
  'on' or 'off'. The default value is persist="", that is brushing is al-
  lowed only once. If persist="on" or persist="off" brushing can be done
  as many time as the user requires. If persist='on' then the unit(s) cur-
  rently brushed are added to those previously brushed. It is possible, every
time a new brushing is done, to use a different color for the brushed units. If persist='off' every time a new brush is performed units previously brushed are removed.

- labeladd: add labels of brushed units in the scatterplot matrix. If this option is '1', we label the units of the last selected group with the unit row index in the matrix X. The default value is labeladd='", i.e. no label is added.

subsize x axis control, a numeric vector containing the subset size with length equal to the number of columns of matrix residuals. If it is not specified it will be set equal to (nrow(residuals) - ncol(residuals) + 1) : nrow(residuals).

selstep Text shown in selected steps, a numeric vector which specifies for which steps of the forward search textlabels are added in the monitoring residual plot after a brushing action in the yXplot. The default is to write the labels at the initial and final step. The default is selstep=c(m0, n) where m0 and n are respectively the first and final step of the search.

selunit Unit labelling. A vector of strings, a string, or a numeric vector for labelling units. If out is an object the threshold is associated with the trajectories of the residuals monitored along the search else it refers to the values of the response variable. If it is a vector of strings, only the lines associated with the units that in at least one step of the search had a residual smaller than selunit[1] or greater than sellunit[2] will have a textbox. If it is a string it specifies the threshold above which labels have to be put. For example selunit='2.6' means that the text labels are written only for the units which have in at least one step of the search a value of the scaled residual greater than 2.6 in absolute value. If it is a numeric vector it contains the list of the units for which it is necessary to put the text labels. The default value of selunit is string selunit='2.5' if y is an object else it is an empty value.

trace Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

Value

none

Author(s)

FSDA team, <valentin.todorov@chello.at>

See Also

spmplot, mdrplot, resfwdplot

Examples

## Not run:
## Example of the use of function regspmplot with all the default options
## regspmplot() with first argument vector y and no option.
## In the first example as input there are two matrices: y and X respectively
## A simple plot is created

```r
n <- 100
p <- 3
X <- matrix(data=rnorm(n*p), nrow=n, ncol=p)
y <- matrix(data=rnorm(n*1), nrow=n, ncol=1)
regspmplot(y, X)
```

## Example of the use of function regspmplot with first argument
## vector y and third argument group.
## Different groups are shown in the yXplot

```r
group <- rep(0, n)
group[1:(n/2)] <- rep(1, n/2)
regspmplot(y, X, group)
```

## Example of the use of function regspmplot with first argument
## vector y, third argument group and fourth argument plot
## (Ex1) plot=TRUE

```r
regspmplot(y, X, group, plot=TRUE)
```

## (Ex1) Set the scale for the x axes, the y axis and control symbol type
## regspmplot(y, X, group, xlim=c(-1,2), ylim=c(0,2), pch=c(10,11), trace=TRUE)

## When the first input argument is an object.
## In the following example the input is an object which also contains
## information about the forward search.
```r
(out <- fsreg(y~X, method="LMS", control=LXS_control(nsamp=1000)))
(out <- fsreg(y~X, bsb=out$bs, monitoring=TRUE))
```

```r
regspmplot(out, plot=0)
```

## End(Not run)

---

### resfwdplot

Plots the trajectories of the monitored scaled (squared) residuals

#### Description

Plots the trajectories of the monitored scaled (squared) residuals

#### Usage

```r
resfwdplot(out, xlim, ylim, xlab, ylab, main, lwd, lty, col, cex.lab, cex.axis, values, fg.thresh, fg.unit, fg.labstep, fg.lwd, fg.lty, fg.col, fg.mark, fg.cex,```
Arguments

out An object returned by one of the monitoring functions (see `FSReda_control`, `Sregeda_control` and `MMregeda_control`). The object is one of `fsreda.object`, `sregeda.object` or `mmregeda.object`.

The needed elements of `out` are

1. RES: matrix containing the residuals monitored in each step of the forward search or any other robust procedure. Every row is associated with a residual (unit). This matrix can be created using function `FSReda`, `Sregeda`, `MMregeda`.
2. Un: (for FSR only) - matrix containing the order of entry in the subset of each unit (required only when datatooltip is true or databrush is not empty).
3. bdp: (for Sreg only) - vector containing a sequence of breakdown point values to monitor on.
4. eff: (for MMreg only) - vector containing a sequence of efficiency values to monitor on.
5. y: a vector containing the response (required only when option databrush is requested).
6. X: a matrix containing the explanatory variables (required only when option databrush is requested).
7. Bols: (n-init+1) x (p+1) matrix containing the estimated beta coefficients monitored in each step of the robust procedure (required only when option databrush is requested and suboption multivarfit is requested).

ylim Control y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.

xlim Control x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.

xlab a title for the x axis

ylab a title for the y axis

main an overall title for the plot

lwd The line width, a positive number, defaulting to 1

lty The line type. Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.

col colors to be used for the highlighted units

cex.lab The magnification to be used for x and y labels relative to the current setting of cex

cex.axis The magnification to be used for axis annotation relative to the current setting of cex
xvalues values for the x axis. Numeric vector of ncol(RES) controlling the x axis coordinates. The default value of xvalues is (nrow(RES) - ncol(RES) + 1):nrow(RES)

fg.thresh (alternative to fg.unit) numeric vector of length 1 or 2 which specifies the highlighted trajectories. If length(fg.thresh) == 1 the highlighted trajectories are those of units that throughout the search had at least once a residual greater (in absolute value) than thresh. The default value is fg.thresh=2.5. If length(fg.thresh) == 2 the highlighted trajectories are those of units that throughout the search had a residual at least once bigger than fg.thresh[2] or smaller than fg.thresh[1].

fg.unit (alternative to fg.thresh), vector containing the list of the units to be highlighted. If fg.unit is supplied, fg.thresh is ignored.

fg.labstep numeric vector which specifies the steps of the search where to put labels for the highlighted trajectories (units). The default is to put the labels at the initial and final steps of the search. fg.labstep='' means no label.

fg.lwd The line width for the highlighted trajectories (units). Default is 1.

fg.lty The line type for the highlighted trajectories (units). Line types can either be specified as an integer (1=solid (default), 2=dashed, 3=dotted, 4=dotdash, 5=longdash, 6=twodash) or as one of the character strings "solid", "dashed", "dotted", "dotdash", "longdash", or "twodash". The latter two are not supported by Matlab.

fg.col colors to be used for the highlighted units.

fg.mark Controls whether to plot highlighted trajectories as symbols. if fg.mark==TRUE each line is plotted using a different symbol else no marker is used (default).

fg.cex controls the font size of the labels of the trajectories in foreground.

bg.thresh numeric vector of length 1 or 2 which specifies how to define the unimportant trajectories. Unimportant trajectories will be plotted using a colormap, in grey or will be hidden. If length(bg.thresh) == 1 the irrelevant units are those which always had a residual smaller (in absolute value) than thresh. If length(bg.thresh) == 2 the irrelevant units are those which always had a residual greater than bg.thresh[1] and smaller than bg.thresh[2]. The default is: bg.thresh=2.5 if n > 100 and bg.thresh=-Inf if n <= 100 i.e. to treat all trajectories as important if n <= 100 and, if n > 100, to reduce emphasis only to trajectories having in all steps of the search a value of scaled residual smaller than 2.5.

bg.style specifies how to plot the unimportant trajectories as defined in option bg.thresh.

  1. bg.style="faint": unimportant trajectories are plotted using a colormap.
  2. bg.style="hide": unimportant trajectories are hidden.
  3. bg.style="greyish": unimportant trajectories are displayed in a faint grey.

When n>100 the default option is bg.style='faint'. When n <= 100 and bg.thresh == -Inf option bstyle is ignored. Remark: bground=0 is equivalent to -Inf that is all trajectories are considered relevant.

tag Plot handle. String which identifies the handle of the plot which is about to be created. The default is to use tag 'pl_resfwd'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.
datatooltip Interactive clicking. It is inactive if this parameter is missing or empty. The default is datatooltip=TRUE, i.e. the user can select with the mouse an individual residual trajectory in order to have information about the corresponding unit. The information displayed depends on the estimator in use. For example for class fsreda.object the information concerns the label and the step of the search in which the unit enters the subset. If datatooltip is a list it may contain the following fields:

1. DisplayStyle determines how the data cursor displays. Possible values are 'datatip' and 'window' (default). 'datatip' displays data cursor information in a small yellow text box attached to a black square marker at a data point you interactively select. 'window' displays data cursor information for the data point you interactively select in a floating window within the figure.

2. SnapToDataVertex: specifies whether the data cursor snaps to the nearest data value or is located at the actual pointer position. Possible values are SnapToDataVertex='on' (default) and SnapToDataVertex='off'.

3. LineColor: controls the color of the trajectory selected with the mouse. It can be an RGB triplet of values between 0 and 1, or character vector indicating a color name. Note that a RGB vector can be conveniently chosen with our MATLAB class FSCColor, see documentation.

4. SubsetLinesColor: enables to control the color of the trajectories of the units that are in the subset at a given step of the search (if resfwdplot() is applied to an object of class fsreda.object) or have a weight greater than 0.9 (if resfwdplot() is applied to an object of class sregeda.object or mmregeda.object). This can be done (repeatedly) with a left mouse click in proximity of the step of interest. A right mouse click will terminate the selection by marking with a up-arrow the step corresponding to the highlighted lines. The highlighted lines by default are in red, but a different color can be specified as RGB triplet or character of color name. Note that a RGB vector can be conveniently chosen with our MATLAB class FSCColor, see documentation. By default SubsetLinesColor="", i.e. the modality is not active. Any initialization for SubsetLinesColor which cannot be interpreted as RGB vector will be converted to blue, i.e. SubsetLinesColor will be forced to be [0 0 1]. If SubsetLinesColor is not empty the previous option LineColor is ignored.

label Character vector containing the labels of the units (optional argument used when datatooltip=TRUE). If this field is not present labels row1, ..., rown will be automatically created and included in the pop up datatooltip window).

nameX Add variable labels in plot. A vector of strings of length p containing the labels of the variables of the regression dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically

namey Add response label. A string containing the label of the response

msg Controls whether to display or not messages on the screen If msg==1 (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.

databrush interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is a scalar or a list) enables
The user to select a set of trajectories in the current plot and to see them highlighted in the y|X plot, i.e. a matrix of scatter plots of y against each column of X, grouped according to the selection(s) done by brushing. If the plot y|X does not exist it is automatically created. In addition, brushed units are automatically highlighted in the minimum deletion residual plot if it is already open. The extension to the following plots will be available in future versions of the toolbox:

1. monitoring leverage plot;
2. maximum studentized residual;
3. s^2 and R^2;
4. Cook distance and modified Cook distance;
5. deletion t statistics.

Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist="on").

If databrush=list(...), it is possible to use all optional arguments of function selectdataFS() and the following optional argument:

1. persist. Persist is an empty value or a character containing 'on' or 'off'. The default value is persist="", that is brushing is allowed only once. If persist="on" or persist="off" brushing can be done as many times as the user requires. If persist="on" then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If persist="off" every time a new brush is performed units previously brushed are removed.

2. bivarfit. Wheather to superimpose bivariate least square lines on the plot (if plot=TRUE. This option adds one or more least squares lines, based on SIMPLE REGRESSION of y on Xi, to the plots of y|Xi. The default is bivarfit=FALSE: no line is fitted. If bivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. If bivarfit=2, two OLS lines are fitted: one to all points and another to the group of the genuine observations. The group of the potential outliers is not fitted. If bivarfit=0 one OLS line is fitted to each group. This is useful for the purpose of fitting mixtures of regression lines. If bivarfit='i1' or bivarfit='i2', etc. an OLS line is fitted to a specific group, the one with index 'i' equal to 1, 2, 3 etc. Again, useful in case of mixtures.

3. multivarfit. Wheather to superimpose multivariate least square lines. This option adds one or more least square lines, based on MULTIVARIATE REGRESSION of y on X, to the plots of y|Xi. The default is multivarfit=FALSE: no line is fitted. If multivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. The line added to the scatter plot y|Xi is avconst + Ci*Xi, where Ci is the coefficient of Xi in the multivariate regression and avconst is the effect of all the other explanatory variables different from Xi evaluated at their centroid (that is overline(y))(C). If multivarfit=2, same action as with multivarfit=1 but this time we also
add the line based on the group of unselected observations (i.e. the normal units).

4. labeladd. Add outlier labels in plot. If labeladd=TRUE, we label the outliers with the unit row index in matrices X and y. The default value is labeladd=FALSE, i.e. no label is added.

standard (MATLAB-style arguments) appearance of the plot in terms of xlim, ylim, axes labels and their font size style, color of the lines, etc.

fground MATLAB-style arguments for the fground trajectories in foreground.

bground MATLAB-style arguments for the fground trajectories in background.

... potential further arguments passed to lower level functions.

Details

No details

Value

No value returned

Author(s)

FSDA team

Examples

## Not run:

n <- 100
y <- rnorm(n)
X <- matrix(rnorm(n*4), nrow=n)

out <- fsreg(y~X, method="LTS")
out <- fsreg(y~X, method="FS", bsb=out$bs, monitoring=TRUE)
resfwdplot(out)

## End(Not run)

**resindexplot**

Plots the residuals from a regression analysis versus index number or any other variable

Description

The function `resindexplot()` plots the residuals from a regression analysis versus index number or any other variable. The residuals come from an output object of any of the regression functions or a simply a vector of values. In order to use the databrush option, the residuals must come from one of the fsdaR regression functions.
resindexplot

Usage

resindexplot(out, x, xlim, ylim, xlab, ylab, main, numlab, indlab, conflev, cex.axis, cex.lab, lwd, nameX, namey, tag, col, cex, databrush, ...)

Arguments

out  A vector containing the residuals from a regression analysis or an object returned by one of the regression functions (see FSR_control, LXS_control, Sreg_control and MMreg_control). The object is one of fsr.object, fsdalts.object, fsdalms.object, sreg.object or mmreg.object. The needed elements of out are at least residuals, but if the option databrush is used, also X and y will be needed.

x  The vector to be plotted on the x-axis. As default the sequence 1:length(residuals) will be used

xlim  Control x scale in plot. Vector with two elements controlling minimum and maximum on the x axis. Default is to use automatic scale.

ylim  Control y scale in plot. Vector with two elements controlling minimum and maximum on the y axis. Default is to use automatic scale.

xlab  a title for the x axis

ylab  a title for the y axis

main  an overall title for the plot

numlab  Number of points to be identified in plots (see also indlab). By default the five points with largest values will be identified. If numlab is a single number containing scalar k, the units with the k largest residuals are labelled in the plots. If numlab is a vector, the units inside vector numlab are labelled in the plots. The default value of numLab=5 and the units with the 5 largest residuals will be labelled. If numlib=0 or numlib=NULL no labelling will be done.

indlab  Which points to be identified in plots (see also numlab) - the units with indexes in the vector indlab are labelled in the plots.

conflev  Confidence interval for the horizontal bands (a numeric vector). It can be a vector of different confidence level values.

Remark: confidence interval is based on the chi^2 distribution

cex.axis  The magnification to be used for axis annotation relative to the current setting of cex

cex.lab  The magnification to be used for x and y labels relative to the current setting of cex

lwd  The line width, a positive number, defaulting to 1

tag  Figure tag (character). Tag of the figure which will host the resindexplot. The default tag is pl_resindex.

col  Fill color for markers that are closed shapes (circle, square, diamond, pentagram, hexagram, and the four triangles). Can be ‘none’ or ‘auto’ or color name(string) or RGB triplet.

cex  Size of the point symbols. The magnification to be used relative to the current setting of cex.
nameX  Add variable labels in plot. A vector of strings of length p containing the labels of the variables of the regression dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically.

namey  Add response label. A string containing the label of the response.

databrush  Interactive mouse brushing. If databrush is missing or empty (default) or databrush=FALSE, no brushing is done. The activation of this option (databrush is a scalar or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the y|X plot, i.e. a matrix of scatter plots of y against each column of X, grouped according to the selection(s) done by brushing. If the plot y|X does not exist it is automatically created. In addition, brushed units are automatically highlighted in the minimum deletion residual plot if it is already open. The extension to the following plots will be available in future versions of the package:

1. monitoring leverage plot;
2. maximum studentized residual;
3. s^2 and R^2;
4. Cook distance and modified Cook distance;
5. deletion t statistics.

Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist="").

If databrush=list(...), it is possible to use all optional arguments of function selectdataFS() and the following optional argument:

1. persist  Persist is an empty value or a character containing 'on' or 'off'. The default value is persist="", that is brushing is allowed only once. If persist="on" or persist="off" brushing can be done as many times as the user requires. If persist='on' then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If persist='off' every time a new brush is performed units previously brushed are removed.

2. bivarfit  This option adds one or more least square lines based on SIMPLE REGRESSION to the plots of y|X, depending on the selected groups. The default is bivarfit=FALSE: no line is fitted. If bivarfit=1, a single OLS line is fitted to all points of each bivariate plot in the scatter matrix y|X. If bivarfit=2, two OLS lines are fitted: one to all points and another to the group of the genuine observations. The group of the potential outliers is not fitted. If bivarfit=0 one OLS line is fitted to each group. This is useful for the purpose of fitting mixtures of regression lines. If bivarfit='i1' or bivarfit='i2', etc. an OLS line is fitted to a specific group, the one with index 'i' equal to 1, 2, 3 etc. Again, useful in case of mixtures.

3. multivarfit  Whether to superimpose multivariate least square lines. This option adds one or more least square lines, based on MULTIVARIATE REGRESSION of y on X, to the plots of y|Xi. The default is multivarfit=FALSE: no line is fitted. If bivarfit=1, a single OLS line is fitted to all points of...
score

Computes the score test for transformation in regression

Description

Computes the score test for transformation in regression

Usage

score(x, ...)

## S3 method for class 'formula'
score(
  formula, 
  data, 
  subset, 
  ...
weights, na.action, model = TRUE, contrasts = NULL, offset, ...
)

## Default S3 method:
score(
x, y, intercept = TRUE, la = c(-1, -0.5, 0, 0.5, 1), lik = FALSE, nocheck = FALSE, trace = FALSE, ...
)

Arguments

x An \( n \times p \) data matrix (\( n \) observations and \( p \) variables). Rows of \( x \) represent observations, and columns represent variables. Missing values (NA's) and infinite values (Inf’s) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

... potential further arguments passed to lower level functions.

formula a formula of the form \( y \sim x_1 + x_2 + \ldots \).

data a data frame from which variables specified in formula are to be taken.

subset an optional vector specifying a subset of observations to be used in the fitting process.

weights an optional vector of weights to be used

NOT USED YET.

na.action a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The “factory-fresh” default is na.omit. Another possible value is NULL, no action. Value na.exclude can be useful.

model logical indicating if the model frame, is to be returned.

contrasts an optional list. See the contrasts.arg of model.matrix.default.

offset this can be used to specify an \( a \ priori \) known component to be included in the linear predictor during fitting. An offset term can be included in the

y Response variable. A vector with \( n \) elements that contains the response variable.

intercept wheather to use constant term (default is intercept=TRUE

...
la values of the transformation parameter for which it is necessary to compute the score test. Default value of lambda is la=c(-1, -0.5, 0, 0.5, 1), i.e., the five most common values of lambda.

lik likelihood for the augmented model. If true the value of the likelihood for the augmented model will be calculated and returned otherwise (default) only the value of the score test will be given.

nocheck Whether to check input arguments. If nocheck=TRUE no check is performed on matrix y and matrix X. Notice that y and X are left unchanged. In other words the additional column of ones for the intercept is not added. The default is nocheck=FALSE.

trace Whether to print intermediate results. Default is trace=FALSE.

Value
An S3 object of class score.object will be returned which is basically a list containing the following elements:

1. la: vector containing the values of lambda for which fan plot is constructed
2. Score: a vector containing the values of the score test for each value of the transformation parameter.
3. Lik: value of the likelihood. This output is produced only if lik=TRUE.

Author(s)
FSDA team, <valentin.todorov@chello.at>

References

Examples

```r
## Not run:
data(wool)
XX <- wool
y <- XX[, ncol(XX)]
X <- XX[, 1:(ncol(XX)-1), drop=FALSE]
(out <- score(X, y))  # call 'score' with all default parameters
(out <- score(cycles-. , data=wool))  # use the formula interface
(out <- score(cycles-. , data=wool, lik=TRUE))  # return the likelihood

data(loyalty)
head(loyalty)
## la is a vector containing the values of \lambda which have to be tested
(out <- score(amount_spent-. , data=loyalty, la=c(0.25, 1/3, 0.4, 0.5)))
(out <- score(amount_spent-. , data=loyalty, la=c(0.25, 1/3, 0.4, 0.5), lik=TRUE))
```
score.object

## End(Not run)

---

**score.object**

*Objects returned by the function* score

### Description

An object of class `score.object` holds information about the result of a call to `score`.

### Value

The functions `print()` and `summary()` are used to obtain and print a summary of the results. An object of class `score` is a list containing at least the following components:

1. la: vector containing the values of lambda for which fan plot is constructed
2. Score: a vector containing the values of the score test for each value of the transformation parameter.
3. Lik: value of the likelihood. This output is produced only if lik=TRUE.

### Examples

```r
## Not run:
data(wool)
(out <- score(cycles~., data=wool, lik=TRUE))
class(out)
summary(out)
## End(Not run)
```

---

**smult**

*Computes S estimators in multivariate analysis*

### Description

Computes S estimators in multivariate analysis
Usage

\[
\text{smult}(\text{x}, \text{monitoring} = \text{FALSE}, \text{plot} = \text{FALSE}, \text{bdp}, \text{nsamp}, \text{conflev} = 0.975, \text{nocheck} = \text{FALSE}, \text{trace} = \text{FALSE}, \ldots)
\]

Arguments

\(x\) An n x p data matrix (n observations and p variables). Rows of \(x\) represent observations, and columns represent variables. Missing values (NA’s) and infinite values (Inf’s) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

\(\text{monitoring}\) Whether to perform monitoring of Mahalanobis distances and other specific quantities

\(\text{plot}\) Plots the Mahalanobis distances against index number. If \(\text{plot} = \text{FALSE}\) (default) or \(\text{plot} = 0\) no plot is produced. The confidence level used to draw the confidence bands for the MD is given by the input option \(\text{conflev}\). If \(\text{conflev}\) is not specified a nominal 0.975 confidence interval will be used. If \(\text{plot} = 2\) a scatter plot matrix with the outliers highlighted is produced. If \(\text{plot}\) is a list it may contain the following fields:

- \(\text{labeladd}\) If \(\text{labeladd} = 1\), the outliers in the \(\text{spm}\) are labelled with the unit row index. The default value is \(\text{labeladd} = "\text{"}, \text{i.e. no label is added}\n
- \(\text{nameY}\) character vector containing the labels of the variables. As default value, the labels which are added are \(Y_1, \ldots Y_p\).

\(\text{bdp}\) Measures the fraction of outliers the algorithm should resist. In this case any value greater than 0 but smaller or equal than 0.5 will do fine (default is \(\text{bdp} = 0.5\)). Note that given \(\text{bdp}\) nominal efficiency is automatically determined.

\(\text{nsamp}\) Number of subsamples which will be extracted to find the robust estimator. If \(\text{nsamp} = 0\) all subsets will be extracted. They will be \((n \text{ choose } p)\). If the number of all possible subset is <\(1000\) the default is to extract all subsets otherwise just 1000.

\(\text{conflev}\) Confidence level which is used to declare units as outliers (scalar). Usually \(\text{conflev} = 0.95, \text{conflev} = 0.975\) or \(\text{conflev} = 0.99\) (individual alpha) \(\text{conflev} = 1 - 0.05/n\), \(\text{conflev} = 1 - 0.025/n\) or \(\text{conflev} = 1 - 0.01/n\) (simultaneous alpha). Default value is \(\text{conflev} = 0.975\).

\(\text{nocheck}\) It controls whether to perform checks on matrix \(Y\). If \(\text{nocheck} = \text{TRUE}\), no check is performed.

\(\text{trace}\) Whether to print intermediate results. Default is \(\text{trace} = \text{FALSE}\).
Details

This function follows the lines of MATLAB/R code developed during the years by many authors. For more details see http://www.econ.kuleuven.be/public/NDBAE06/programs/ and the R package CovSest. The core of these routines, e.g. the resampling approach, however, has been completely redesigned, with considerable increase of the computational performance.

Value

Depending on the input parameter monitoring, one of the following objects will be returned:

1. smult.object
2. smulteda.object

Author(s)

FSDA team, <valentin.todorov@chello.at>

References


Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- smult(hbk[,1:3]))
class(out)
summary(out)

## Generate contaminated data (200,3)
n <- 200
p <- 3
set.seed(123456)
X <- matrix(rnorm(n*p), nrow=n)
Xcont <- X

out1 <- smult(Xcont, trace=TRUE) # no plots (plot defaults to FALSE)
names(out1)

## plot=TRUE - generates: (1) a plot of Mahalanobis distances against
## index number. The confidence level used to draw the confidence bands for
## the MD is given by the input option conflev. If conflev is
## not specified a nominal 0.975 confidence interval will be used and
## (2) a scatter plot matrix with the outliers highlighted.

(out1 <- smult(Xcont, trace=TRUE, plot=TRUE))
```
## plots is a list: the spm shows the labels of the outliers.
(out1 <- smult(Xcont, trace=TRUE, plot=list(labeladd="1")))

## plots is a list: the spm uses the variable names provided by 'nameY'.
(out1 <- smult(Xcont, trace=TRUE, plot=list(nameY=c("A", "B", "C"))))

## smult() with monitoring
(out2 <- smult(Xcont, monitoring=TRUE, trace=TRUE))
names(out2)

## Forgery Swiss banknotes examples.
data(swissbanknotes)

(out1 <- smult(swissbanknotes[101:200,], plot=TRUE))
(out1 <- smult(swissbanknotes[101:200,], plot=list(labeladd="1")))

## End(Not run)

---

### smult.object

**Description of smult.object Objects**

**Description**

An object of class `smult.object` holds information about the result of a call to `smult`.

**Value**

The object itself is basically a **list** with the following components:

- **loc**: p-by-1 vector containing S estimate of location.
- **shape**: p-by-p matrix containing robust estimate of the shape matrix. Remark: det(shape)=1.
- **scale**: robust estimate of the scale.
- **cov**: scale^2 * shape: robust estimate of covariance matrix.
- **bs**: a (p+1) vector containing the units forming best subset associated with S estimate of location.
- **md**: n-by-1 vector containing the estimates of the robust Mahalanobis distances (in squared units). This vector contains the distances of each observation from the location of the data, relative to the scatter matrix cov.
- **outliers**: A vector containing the list of the units declared as outliers using confidence level specified in input scalar `conflev`.
- **conflev**: Confidence level that was used to declare outliers.
- **singsub**: Number of subsets without full rank. Notice that `singsub > 0.1*(number of subsamples)` produces a warning.
weights n x 1 vector containing the estimates of the weights.
X the data matrix X

The object has class "smult".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- smult(hbk[,1:3]))
class(out)
summary(out)
## End(Not run)
```

smulteda.object Description of smulteda.object Objects

Description

An object of class smulteda.object holds information about the result of a call to smult with monitoring=TRUE.

Value

The object itself is basically a list with the following components:

- `Loc` length(bdp)-by-p matrix containing S estimate of location for each value of bdp.
- `Shape` p-by-p-by-length(bdp) 3D array containing robust estimate of the shape for each value of bdp. Remark: det(shape)=1.
- `Scale` length(bdp) vector containing robust estimate of the scale for each value of bdp.
- `Cov` p-by-p-by-length(bdp) 3D array containing robust estimate of covariance matrix for each value of bdp. Note that scale(i)^2 * shape[,i] = robust estimate of covariance matrix.
- `Bs` (p+1)-by-length(bdp) matrix containing the units forming best subset for each value of bdp.
- `MAL` n-by-length(bdp) matrix containing the estimates of the robust Mahalanobis distances (in squared units) for each value of bdp.
- `Outliers` n-by-length(bdp) matrix containing flags for the outliers. Boolean matrix containing the list of the units declared as outliers for each value of bdp using confidence level specified in input scalar conflev
- `Weights` n x length(bdp) matrix containing the weights for each value of bdp.
- `conflev` Confidence level that was used to declare outliers.
- `singsub` Number of subsets without full rank. Notice that singsub > 0.1*(number of subsamples) produces a warning.
bdp vector which contains the values of bdp which have been used.

X the data matrix X.

The object has class "smulteda".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- smult(hbk[,1:3], monitoring=TRUE))
class(out)
summary(out)
## End(Not run)
```

---

**Description**

Produces an interactive scatterplot matrix with boxplots or histograms on the main diagonal and possibly robust bivariate contours

**Usage**

```r
spmplot(
  x, group, plot, variables, col, cex, pch, labeladd, label, legend, dispopt = c("hist", "box"), tag, datatooltip, databrush, trace = FALSE,
  ...
)
```
Arguments

X  data matrix (2D array) containing n observations on p variables or an object of S3 class fsmeda.object returned by fsmult with monitoring=TRUE - a list containing the monitoring of minimum Mahalanobis distance

group  grouping variable. Vector with n elements. Specifies a grouping variable defined as a categorical variable (factor), numeric, or array of strings, or string matrix, and it must have the same number of rows as X. This grouping variable determines the marker and color assigned to each point. Remark: if group is used to distinguish a set of outliers from a set of good units, the id number for the outliers should be the larger (see optional field labeladd of parameter plot for details).

plot  controls the names which are displayed in the margins of the scatter-plot matrix, the labels of the legend the colors and the symbols. If plot is empty (plot=FALSE or plot=0 or plot=c() or plot=NULL) empty strings are displayed and no label and no name is added to the plot. If plot=TRUE or plot=1, the names Y1, ..., Yp are added to the margins of the the scatter plot matrix else nothing is added. If plot is a list, it is possible to control not only the names but also, point labels, colors and symbols. More precisely list plot may contain the following elements:
1. labeladd - see parameter labeladd
2. nameY - a character string containing the labels of the variables. As default value, the labels which are added are Y1, ..., Yp. See parameter variables.
3. clr - see parameter col
4. sym - see parameter pch
5. siz - see parameter cex
6. doleg - see parameter legend
7. label - see parameter label

variables  a character string with the names of the variables

col  color specification for the data point. Can be different for each group. By default, the order of the colors is blue, red, black, magenta, green, cyan and yellow.
cex  the size of the symbols used for plotting. By default cex=1 the symbol size depends on the number of plots and the size of the figure window. Values larger than 1 will increase the size and values smaller than 1 will decrease the size.
pch  specification of the symbols to use. For example, if there are three groups, and pch=c(1, 3, 4), the first group will be plotted with a circle, the second with a plus, and the third with a 'x' (see ?pch or ?points for a list of symbols. NOTE: not all symbols available in R can be mapped to the symbols in MATLAB.

labeladd  logical, controls wheather the elements belonging to the last group in the scatterplot matrix are labelled with their unit row index or their rowname. The rowname is taken from the parameter label or if it is missing, from the sequence 1:n. The default value is labeladd=FALSE, i.e. no label is added.

label  a character vector of length n (the number of rows in the data matrix) containing the labels of the units. If this field is empty the sequence 1:n will be used to label the units.
### legend
Logical, controls where a legend is shown or not.

### dispopt
Controls how to fill the diagonals in the plot (main diagonal of the scatter plot matrix). Set dispopt='hist' (default) to plot histograms, or dispopt='box' to plot boxplots. The style which is used for univariate boxplots is traditional, if the number of groups is less or equal 5, else it is 'compact' (plot boxes using a smaller box style designed for plots with many groups).

### tag
Plot handle. String which identifies the handle of the plot which is about to be created. The default is tag='pl_mmd'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.

### datatooltip
If datatooltip is not empty the user can use the mouse in order to have information about the unit selected, the step in which the unit enters the search and the associated label. If datatooltip is a list, it is possible to control the aspect of the data cursor (see MATLAB function datacursormode() for more details or see the examples below). The default options areDisplayStyle="Window" and SnapToDataVertex="on".

### databrush
Interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is TRUE or a list) enables the user to select a set of trajectories in the current plot and to see them highlighted in the scatterplot matrix. If the scatterplot matrix does not exist it is automatically created. In addition, brushed units can be highlighted in the monitoring MD plot. Note that the window style of the other figures is set equal to that which contains the monitoring residual plot. In other words, if the monitoring residual plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist="").

If databrush=list(...), it is possible to use all optional arguments of the MATLAB function selectdataFS() and the following optional arguments:

- **persist**: This option can be an empty value or a character containing 'on' or 'off'. The default value is persist="", that is brushing is allowed only once. If persist="on" or persist="off" brushing can be done as many time as the user requires. If persist='on' then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing is done, to use a different color for the brushed units. If persist='off' every time a new brush is performed units previously brushed are removed.

- **labeladd**: add labels of brushed units in the scatterplot matrix. If this option is '1', we label the units of the last selected group with the unit row index in the matrix X. The default value is labeladd="", i.e. no label is added.

### trace
Whether to print intermediate results. Default is trace=FALSE.

... Potential further arguments passed to lower level functions.
Author(s)

FSDA team, <valentin.todorov@chello.at>

Examples

## Not run:
## Call of spmplot() without optional parameters.
## Iris data: scatter plot matrix with univariate boxplots on the main
## diagonal.

X <- iris[,1:4]
group <- iris[,5]
spmplot(X, group, variables=c('SL','SW','PL','PW'), dispopt="box")

## Example of spmplot() called by routine fsmult().
## Generate contaminated data.

n <- 200; p <- 3
X <- matrix(rnorm(n*p), ncol=3)
Xcont <- X
Xcont[1:5,] <- Xcont[1:5,] + 3

## spmplot is called automatically by all outlier detection methods, e.g. fsmult()
out <- fsmult(Xcont, plot=TRUE);

## Now test the direct use of fsmult(). Set two groups, e.g. those obtained
## from fsmult().

group = rep(0, n)
group[out$outliers] <- 1

## option 'labeladd' is used to label the outliers
## By default, the legend identifies the groups with the identifiers
## given in vector 'group'.
## Set the colors for the two groups to blue and red.

spmplot(Xcont, group, col=c("blue", "red"), labeladd=1, dispopt="box")

## End(Not run)
sregeda.object

Value

The object itself is basically a list with the following components:

- **beta**: p-by-1 vector containing the estimated regression parameters (in step n-k).
- **scale**: scalar containing the estimate of the scale (sigma).
- **bs**: p x 1 vector containing the units forming best subset associated with S estimate of regression coefficient.
- **residuals**: residuals.
- **fittedvalues**: fitted values.
- **outliers**: k x 1 vector containing the list of the k units declared as outliers or NULL if the sample is homogeneous.
- **conflev**: Confidence level which is used to declare units as outliers. Usually conflev=0.95, 0.975, 0.99 (individual alpha) or conflev=1-0.05/n, 1-0.025/n, 1-0.01/n (simultaneous alpha). Default value is 0.975.
- **singsub**: Number of subsets without full rank. Notice that singsub > 0.1*(number of subsamples) produces a warning.
- **weights**: n x 1 vector containing the estimates of the weights.
- **rhofunc**: Specifies the rho function which has been used to weight the residuals.
- **rhofuncparam**: Vector which contains the additional parameters for the specified rho function which has been used. For hyperbolic rho function the value of k = sup CVC. For Hampel rho function the parameters a, b and c.

The object has class "sreg".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="S"))
class(out)
summary(out)
## End(Not run)
```

sregeda.object | Description of sregeda Objects

Description

An object of class sregeda.object holds information about the result of a call to fsreg when method="S" and monitoring=TRUE.
Value

The object itself is basically a list with the following components:

- **Beta**: matrix containing the S estimator of regression coefficients for each value of bdp.
- **Scale**: vector containing the estimate of the scale (sigma) for each value of bdp. This is the value of the objective function.
- **BS**: p x 1 vector containing the units forming best subset associated with S estimate of regression coefficient.
- **RES**: n x length(bdp) matrix containing the monitoring of scaled residuals for each value of bdp.
- **Weights**: n x length(bdp) matrix containing the estimates of the weights for each value of bdp.
- **Outliers**: Boolean matrix containing the list of the units declared as outliers for each value of bdp using confidence level specified in input scalar conflev.
- **conflev**: Confidence level which is used to declare units as outliers. Remark: conflev will be used to draw the horizontal line (confidence band) in the plot.
- **Singsub**: Number of subsets without full rank. Notice that singsub[bdp[jj]] > 0.1*(number of subsamples) produces a warning.
- **rhofunc**: Specifies the rho function which has been used to weight the residuals.
- **rhofuncparam**: Vector which contains the additional parameters for the specified rho function which has been used. For hyperbolic rho function the value of k = sup CVC. For Hampel rho function the parameters a, b and c.
- **X**: the data matrix X
- **y**: the response vector y

The object has class "sregeda".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="S", monitoring=TRUE))
class(out)
summary(out)
```

## End(Not run)
Sregeda_control

Description

Creates an object of class Sregeda_control to be used with the fsreg() function, containing various control parameters.

Usage

Sregeda_control(intercept = TRUE, bdp = seq(0.5, 0.01, -0.01),
                 rhofunc = c("bisquare", "optimal", "hyperbolic", "hampel", "mdpd", "AS"),
                 rhofuncparam, nsamp = 1000, refsteps = 3, reftol = 1e-06,
                 refstepsbestr = 50, reftolbestr = 1e-08,
                 minsctol = 1e-07, bestr = 5,
                 conflev, msg = TRUE, nocheck = FALSE, plot = FALSE)

Arguments

intercept Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.
bdp Breakdown point. It measures the fraction of outliers the algorithm should resist. In this case any value greater than 0 but smaller or equal than 0.5 will do fine. The default value of bdp is a sequence from 0.5 to 0.01 with step 0.01
rhofunc Specifies the rho function which must be used to weight the residuals. Possible values are 'bisquare' 'optimal' 'hyperbolic' 'hampel'.
  1. 'bisquare' uses Tukey’s rho and psi functions. See TBrho and TBpsi.
  2. 'optimal' uses optimal rho and psi functions. See OPTrho and OPTpsi.
  3. 'hyperbolic' uses hyperbolic rho and psi functions. See HYPrho and HYPpsi.
  4. 'hampel' uses Hampel rho and psi functions. See HArho and HApri.

The default is 'bisquare'.
rhofuncparam Additional parameters for the specified rho function. For hyperbolic rho function it is possible to set up the value of k = sup CVC (the default value of k is 4.5).

For Hampel rho function it is possible to define parameters a, b and c (the default values are a=2, b=4, c=8)
nsamp Number of subsamples which will be extracted to find the robust estimator, scalar. If nsamp=0 all subsets will be extracted. They will be \((n \choose p)\). If the number of all possible subset is <1000 the default is to extract all subsets otherwise just 1000.
refsteps Number of refining iterations in each subsample (default is refsteps=3). refsteps = 0 means "raw-subsampling" without iterations.
reftol Tolerance for the refining steps. The default value is 1e-6
Sregeda_control

refstepsbestr  Scalar defining number of refining iterations for each best subset (default = 50).
reftolbestr  Tolerance for the refining steps for each of the best subsets. The default value is reftolbestr=1e-8.
minsctol  Value of tolerance for the iterative procedure for finding the minimum value of the scale for each subset and each of the best subsets (It is used by subroutine minscale.m). The default value is minsctol=1e-7.
bestr  Defins the number of "best betas" to remember from the subsamples. These will be later iterated until convergence (default is bestr=5).
conflev  Confidence level which is used to declare units as outliers. Usually conflev=0.95, 0.975, 0.99 (individual alpha) or conflev=1-0.05/n, 1-0.025/n, 1-0.01/n (simultaneous alpha). Default value is 0.975
msg  Controls whether to display or not messages on the screen If msg==1 (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.
nocheck  Check input arguments, scalar. If nocheck=TRUE no check is performed on matrix y and matrix X. Notice that y and X are left unchanged. In other words the additional column of ones for the intercept is not added. As default nocheck=FALSE.
plot  Plot on the screen. Scalar. If plots=TRUE the plot of minimum deletion residual with envelopes based on n observations and the scatterplot matrix with the outliers highlighted is produced. If plots=2 the user can also monitor the intermediate plots based on envelope superimposition. If plots=FALSE (default) no plot is produced.

Details

Creates an object of class Sregeda_control to be used with the fsreg() function, containing various control parameters.

Value

An object of class "Sregeda_control" which is basically a list with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also as FSR_control, MMreg_control and LXS_control

Examples

## Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~, data=hbk, method="S", monitoring=TRUE,
control=Sregeda_control(nsamp=500, rhofunc='hyperbolic')))
Sreg_control

Creates an Sreg_control object

Description

Creates an object of class Sreg_control to be used with the fsreg() function, containing various control parameters for calling the MATLAB function Sreg().

Usage

Sreg_control(intercept = TRUE, bdp = 0.5,
            rhofunc = c("bisquare", "optimal", "hyperbolic", "hampel", "mdpd", "AS"),
            rhofuncparam, nsamp = 1000, refsteps = 3, reftol = 1e-06,
            refstepsbestr = 50, reftolbestr = 1e-08,
            minsctol = 1e-07, bestr = 5,
            conflev, msg = TRUE, nocheck = FALSE, plot = FALSE)

Arguments

intercept Indicator for constant term. Scalar. If intercept=TRUE, a model with constant term will be fitted (default), else, no constant term will be included.

bdp Breakdown point. It measures the fraction of outliers the algorithm should resist. In this case any value greater than 0 but smaller or equal than 0.5 will do fine. Note that given bdp nominal efficiency is automatically determined.

rhofunc Specifies the rho function which must be used to weight the residuals. Possible values are 'bisquare' 'optimal' 'hyperbolic' 'hampel'.
   1. 'bisquare' uses Tukey's rho and psi functions. See TBrho and TBpsi.
   2. 'optimal' uses optimal rho and psi functions. See OPTrho and OPTpsi.
   3. 'hyperbolic' uses hyperbolic rho and psi functions. See HYPrho and HYPpsi.
   4. 'hampel' uses Hampel rho and psi functions. See HArho and HAppsi.

The default is 'bisquare'.

rhofuncparam Additional parameters for the specified rho function. For hyperbolic rho function it is possible to set up the value of k = sup CVC (the default value of k is 4.5).

For Hampel rho function it is possible to define parameters a, b and c (the default values are a=2, b=4, c=8)

nsamp Number of subsamples which will be extracted to find the robust estimator, scalar. If nsamp=0 all subsets will be extracted. They will be (n choose p). If the number of all possible subset is <1000 the default is to extract all subsets otherwise just 1000.

refsteps Number of refining iterations in each subsample (default is refsteps=3). refsteps = 0 means "raw-subsampling" without iterations.
reftol  Tolerance for the refining steps. The default value is 1e-6
borfstepsbestr  Scalar defining number of refining iterations for each best subset (default = 50).
reftolbestr  Tolerance for the refining steps for each of the best subsets. The default value is reftolbestr=1e-8.
minsctol  Value of tolerance for the iterative procedure for finding the minimum value of the scale for each subset and each of the best subsets (It is used by subroutine minscale.m). The default value is minsctol=1e-7.
bestr  Defines the number of "best betas" to remember from the subsamples. These will be later iterated until convergence (default is bestr=5).
conlev  Confidence level which is used to declare units as outliers. Usually conlev=0.95, 0.975, 0.99 (individual alpha) or conlev=1-0.05/n, 1-0.025/n, 1-0.01/n (simultaneous alpha). Default value is 0.975
msg  Controls whether to display or not messages on the screen If msg=1 (default) messages are displayed on the screen about step in which signal took place else no message is displayed on the screen.
nocheck  Check input arguments, scalar. If nocheck=TRUE no check is performed on matrix $y$ and matrix $X$. Notice that $y$ and $X$ are left unchanged. In other words the additional column of ones for the intercept is not added. As default nocheck=FALSE.
plot  Plot on the screen. Scalar. If plots=TRUE the plot of minimum deletion residual with envelopes based on n observations and the scatterplot matrix with the outliers highlighted is produced. If plots=2 the user can also monitor the intermediate plots based on envelope superimposition. If plots=FALSE (default) no plot is produced.

Details

Creates an object of class Sreg_control to be used with the fsreg() function, containing various control parameters.

Value

An object of class "Sreg_control" which is basically a list with components the input arguments of the function mapped accordingly to the corresponding Matlab function.

Author(s)

FSDA team

See Also

See Also as FSR_control, MMreg_control and LXS_control

Examples

```r
# Not run:
data(hbk, package="robustbase")
(out <- fsreg(Y~., data=hbk, method="S", control=Sreg_control(bdp=0.25, nsamp=500)))

# End(Not run)
```
summary.fsdalms  

Summary Method for fsdalms objects

Description

`summary` method for class "fsdalms".

Usage

```r
## S3 method for class 'fsdalms'
summary(object, correlation = FALSE, ...)
## S3 method for class 'summary.fsdalms'
print(x, digits = max(3, getOption("digits") - 3),
      signif.stars = getOption("show.signif.stars"), ...)
```

Arguments

- `object, x` an object of class "fsdalms" (or "summary.fsdalms"); usually, a result of a call to `fsreg`.
- `correlation` logical; if TRUE, the correlation matrix of the estimated parameters is returned and printed.
- `digits` the number of significant digits to use when printing.
- `signif.stars` logical indicating if “significance stars” should be printer, see `printCoefmat`.
- `...` further arguments passed to or from other methods.

Details

`summary.fsdalms()`, the S3 method, simply returns an (S3) object of class "summary.fsdalms" for which there’s a `print` method:

`print.summary.fsdalms` prints summary statistics for the forward search (FS) regression estimates. While the function `print.fsdalms` prints only the robust estimates of the coefficients, `print.summary.fsdalms` will print also the regression table.

Value

`summary.fsdalms` returns an `summary.fsdalms` object, whereas the print methods returns its first argument via `invisible`, as all print methods do.

See Also

`fsreg, summary`
Examples

## Not run:

```r
data(Animals, package = "MASS")
brain <- Animals[c(1:24, 26:25, 27:28),]
lbrain <- log(brain)
(fs <- fsreg(brain~body, data=lbrain, method="LTS"))
summary(fs)

## compare to the result of ltsReg() from 'robustbase'
library(robustbase)
(lts <- ltsReg(brain~body, data=lbrain))
summary(lts)

## End(Not run)
```

summary.fsdalts  Summary Method for fsdalts objects

Description

`summary` method for class "fsdalts".

Usage

```r
## S3 method for class 'fsdalts'
summary(object, correlation = FALSE, ...)
## S3 method for class 'summary.fsdalts'
print(x, digits = max(3, getOption("digits") - 3),
    signif.stars = getOption("show.signif.stars"), ...)
```

Arguments

- `object, x`:
  - an object of class "fsdalts" (or "summary.fsdalts"); usually, a result of a call to `fsreg`.
- `correlation`:
  - logical; if TRUE, the correlation matrix of the estimated parameters is returned and printed.
- `digits`:
  - the number of significant digits to use when printing.
- `signif.stars`:
  - logical indicating if “significance stars” should be printed, see `printCoefmat`.
- `...`:
  - further arguments passed to or from other methods.

Details

`summary.fsdalts()` is a generic function which, for class "fsdalts", returns an (S3) object of class "summary.fsdalts" which has a `print` method:

`print.summary.fsdalts` prints summary statistics for the forward search (FS) regression estimates. While the function `print.fsdalts` prints only the robust estimates of the coefficients, `print.summary.fsdalts` will print also the regression table.
Value

`summary.fsr` returns an object, whereas the print methods returns its first argument via `invisible`, as all print methods do.

See Also

`fsreg`, `summary`

Examples

```r
## Not run:

data(Animals, package = "MASS")
brain <- Animals[1:24, 26:25, 27:28],]
lbrain <- log(brain)
(fs <- fsreg(brain~body, data=lbrain, method="LTS"))
summary(fs)

## compare to the result of ltsReg() from 'robustbase'
library(robustbase)
(lts <- ltsReg(brain~body, data=lbrain))
summary(lts)

## End(Not run)
```

### Summary Method for FSR objects

**Summary Method for FSR objects**

**Description**

`summary` method for class "fsr".

**Usage**

```r
## S3 method for class 'fsr'
summary(object, correlation = FALSE, ...)
## S3 method for class 'summary.fsr'
print(x, digits = max(3, getOption("digits") - 3),
    signif.stars = getOption("show.signif.stars"), ...)
```

**Arguments**

- `object, x` an object of class "fsr" (or "summary.fsr"); usually, a result of a call to `fsreg`
- `correlation` logical; if `TRUE`, the correlation matrix of the estimated parameters is returned and printed.
- `digits` the number of significant digits to use when printing.
- `signif.stars` logical indicating if "significance stars" should be printed, see `printCoefmat`
- `...` further arguments passed to or from other methods.
Details

`summary.fsr()`, the S3 method, simply returns an (S3) object of class "summary.fsr" for which there's a `print` method:

`print.summary.fsr` prints summary statistics for the forward search (FS) regression estimates. While the function `print.fsr` prints only the robust estimates of the coefficients, `print.summary.fsr` will print also the regression table.

Value

`summary.fsr` returns an `summary.fsr` object, whereas the `print` methods returns its first argument via `invisible`, as all `print` methods do.

See Also

`fsreg`, `summary`

Examples

```r
## Not run:
data(Animals, package = "MASS")
brain <- Animals[c(1:24, 26:25, 27:28),]
lbrain <- log(brain)
(fs <- fsreg(brain~body, data=lbrain, method="FS"))
summary(fs)
## End(Not run)
```

swissbanknotes

Swiss banknote data

Description

Six variables measured on 100 genuine and 100 counterfeit old (printed before the second world war) Swiss 1000-franc bank notes (Flury and Riedwyl, 1988).

Usage

data(swissbanknotes)

Format

A data frame with 200 observations on the following 7 variables.

- `length` Length of bill, mm
- `left` Width of left edge, mm
- `right` Width of right edge, mm
swissheads

bottom  Bottom margin width, mm

top  Top margin width, mm
diagonal  Length of image diagonal, mm
class  1 = genuine, 2 = counterfeit

Source


References


Examples

```r
library(rrcov)
data(swissbanknotes)
head(swissbanknotes)
plot(CovMcd(swissbanknotes[, 1:6]), which="pairs", col=swissbanknotes$class)
```

---

**swissheads**  
**Swiss heads data**

**Description**

Six dimensions in millimetres of the heads of 200 twenty year old Swiss soldiers (Flury and Riedwyl, 1988, p. 218 and also Flury, 1997, p. 6).

The data were collected to determine the variability in size and shape of heads of young men in order to help in the design of a new protection mask for the Swiss army.

**Usage**

```r
data(swissheads)
```

**Format**

A data frame with 200 observations on the following 6 variables.

minimal_frontal_breadth  Minimal frontal breadth, mm

breadth_angulus_mandibulae  Breadth of angulus mandibulae, mm

true_facial_height  True facial height, mm

length_glabella_nasi  Length from glabella to apex nasi, mm

length_tragion_nasion  Length from tragion to nasion, mm

length_tragion_gnathion  Length from tragion to gnathion, mm
tclusteda.object

Source

References

Examples
library(rrcov)
data(swissheads)
head(swissheads)
plot(CovMcd(swissheads), which="pairs")

---

tclusteda.object

Objects returned by the function tclustfsda with the option monitoring=TRUE

Description
An object of class tclusteda.object holds information about the result of a call to tclustfsda with the option monitoring=TRUE.

Value
The functions print() and summary() are used to obtain and print a summary of the results. An object of class tclusteda is a list containing at least the following components:
call the matched call
k number of groups
alpha trimming level
restrfactor restriction factor
IDX an n-by-length(alpha) vector containing assignment of each unit to each of the k groups. Cluster names are integer numbers from 1 to k. 0 indicates trimmed observations. The first column refers to alpha[1], the second column refers to alpha[2], ..., the last column refers to alpha[length(alpha)].

MU a 3 dimensional array of size k-by-p-by-length(alpha) containing the monitoring of the centroid for each value of alpha. MU[, ,1] refers to alpha[1] ..., MU[, ,length(alpha)] refers to alpha[length(alpha)]. The first row in each slice refers to group 1, second row refers to group 2, etc.

SIGMA A list of length length(alpha) containing in element j, with j=1, 2, ..., length(alpha), the 3D array of size p-by-p-by-k containing the k (constrained) estimated covariance matrices associated with alpha[j].
Amon stands for alpha monitoring. Matrix of size \((\text{length}(\alpha)-1)\times 4\) which contains for two consecutive values of \(\alpha\) the monitoring of three quantities (change in classification, change in centroid location, change in covariance location).

- 1st col = value of \(\alpha\).
- 2nd col = ARI index.
- 3rd col = squared Euclidean distance between centroids.
- 4th col = squared Euclidean distance between covariance matrices.

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- tclustfsda(hbk[, 1:3], k=2, monitoring=TRUE))
class(out)
summary(out)
## End(Not run)
```

tclustfsda

Computes trimmed clustering with scatter restrictions

Description

Partitions the points in the \(n\times v\) data matrix \(Y\) into \(k\) clusters. This partition minimizes the trimmed sum, over all clusters, of the within-cluster sums of point-to-cluster-centroid distances. Rows of \(Y\) correspond to points, columns correspond to variables. Returns in the output object of class \texttt{tclustfsda.object} an \(n\times 1\) vector \(\text{idx}\) containing the cluster indices of each point. By default, \texttt{tclustfsda()} uses (squared), possibly constrained, Mahalanobis distances.

Usage

```r
tclustfsda(
  x,
  k,
  alpha,
  restrfactor = 12,
  monitoring = FALSE,
  plot = FALSE,
  nsamp,
  refsteps = 15,
  reftol = 1e-13,
  equalweights = FALSE,
  mixt = 0,
  msg = FALSE,
  nocheck = FALSE,
)```

Arguments

\(x\) An \(n \times p\) data matrix (\(n\) observations and \(p\) variables). Rows of \(x\) represent observations, and columns represent variables. Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

\(k\) Number of groups.

\(\alpha\) A scalar between 0 and 0.5 or an integer specifying the number of observations which have to be trimmed. If \(\alpha=0\), \(\text{tclust}\) reduces to traditional model based or mixture clustering (mclust): see for example the Matlab function \(\text{gmdistribution}\). More in detail, if \(0 < \alpha < 1\) clustering is based on \(h = \text{floor}(n \times (1-\alpha))\) observations, else if \(\alpha\) is an integer greater than 1 clustering is based on \(h = n - \text{floor}(\alpha)\). If \(\text{monitoring=TRUE}\), \(\alpha\) is a vector which specifies the values of trimming levels which have to be considered - contains decreating elements which lie in the interval 0 and 0.5. For example if \(\alpha=c(0.1, 0.05, 0)\), \(\text{tclust()}\) considers these 3 values of trimming level. The default for \(\alpha\) is vector \(\alpha=c(0.1, 0.05, 0)\). The sequence is forced to be monotonically decreasing.

\(\text{restrfactor}\) Positive scalar which constrains the allowed differences among group scatters. Larger values imply larger differences of group scatters. On the other hand a value of 1 specifies the strongest restriction forcing all eigenvalues/determinants to be equal and so the method looks for similarly scattered (respectively spherical) clusters. The default is to apply \(\text{restrfactor}\) to eigenvalues. In order to apply \(\text{restrfactor}\) to determinants it is necessary to use the optional input argument \(\text{restrtype}\).

\(\text{monitoring}\) If \(\text{monitoring=TRUE}\) TCLUST is performed for a series of values of the trimming factor \(\alpha\) given \(k\) (number of groups) and given \(c\) (restriction factor).

\(\text{plot}\) If \(\text{plot=}\text{FALSE}\) (default) or \(\text{plot=}0\) no plot is produced. If \(\text{plot=}\text{TRUE}\) and \(\text{monitoring=}\text{FALSE}\) a plot with the classification is shown (using the \(\text{spmplot}\) function). The plot can be:

- for \(p = 1\), a histogram of the univariate data,
- for \(p = 2\), a bivariate scatterplot,
- for \(p > 2\), a scatterplot matrix generated by the MATLAB function \(\text{spmplot()}\).

When \(p \geq 2\) the following additional features are offered (for \(p = 1\) the behaviour is forced to be as for \(\text{plots=}\text{TRUE}\):
• plot = 'contourf' adds in the background of the bivariate scatterplots a filled contour plot. The colormap of the filled contour is based on grey levels as default. This argument may also be inserted in a field named 'type' of a list. In the latter case it is possible to specify the additional field 'cmap', which changes the default colors of the color map used. The field 'cmap' may be a three-column matrix of values in the range [0,1] where each row is an RGB triplet that defines one color. Check the colormap function for additional informations.

• plot = 'contour' adds in the background of the bivariate scatterplots a contour plot. The colormap of the contour is based on grey levels as default. This argument may also be inserted in a field named type of a list. In the latter case it is possible to specify the additional field cmap, which changes the default colors of the color map used. The field cmap may be a three-column matrix of values in the range [0,1] where each row is an RGB triplet that defines one color. Check the colormap() (MATLAB) function for additional information.

• plot = 'ellipse' superimposes confidence ellipses to each group in the bivariate scatterplots. The size of the ellipse is \( q \chi^2(0.95, 2) \), i.e. the confidence level used by default is 95 percent. This argument may also be inserted in a field named type of a list. In the latter case it is possible to specify the additional field conflev, which specifies the confidence level to use and it is a value between 0 and 1.

• plot = 'boxplotb' superimposes on the bivariate scatterplots the bivariate boxplots for each group, using the boxplotb function. This argument may also be inserted in a field named type of a list.

The parameter plot can be also a list and in this case its elements are:

• type - specifies the type of plot as when plot option is a character. Therefore, plot$type can be one of 'contourf', 'contour', 'ellipse' or 'boxplotb'.

• cmap - used to set a colormap for the plot type (MATLAB style). For example, plot$cmap = 'autumn'. See the MATLAB help of colormap for a list of colormap possibilities.

• conflev - this is the confidence level for the confidence ellipses. It must be a scalar between 0 and 1.

If plot=TRUE and monitoring=TRUE two plots are shown. The first plot (monitor plot) shows three panels with the monitoring between two consecutive values of alpha: (i) the change in classification using ARI index (top panel), (ii) the change in centroids using squared euclidean distances (central panel) and (iii) the change in covariance matrices using squared euclidean distance (bottom panel).

The second plot (gscatter plot) shows a series of subplots which monitor the classification for each value of alpha. In order to make sure that consistent labels are used for the groups, between two consecutive values of alpha, we assign label r to a group if this group shows the smallest distance with group r for the previous value of alpha. The type of plot which is used to monitor the stability of the classification depends on the data dimensionality p.

• for p = 1, a histogram of the univariate data (the MATLAB function histFS() is called),
• for \( p = 2 \), a bivariate scatterplot (the MATLAB function gscatter() is called),
• for \( p > 2 \), a scatterplot of the first two principal components (function gscatter() is called and we show on the axes titles the percentage of variance explained by the first two principal components).

Also in the case of monitoring=TRUE the parameter plot can be a list and its elements are:

• name: character vector which enables to specify which plot to display. name = "gscatter" produces a figure with a series of subplots which show the classification for each value of alpha. name = "monitor" shows a figure with three panels which monitor between two consecutive values of alpha the change in classification using ARI index (top panel), the change in centroids using squared euclidean distances (central panel), the change in covariance matrices using squared euclidean distance (bottom panel). If this field is not specified, by default name=c("gscatter", "monitor") and both figures will be shown.

• alphasel: a numeric vector which specifies for which values of alpha it is possible to see the classification. For example if alphasel = c(0.05, 0.02), the classification will be shown just for alpha=0.05 and alpha=0.02. If this field is not specified alphasel=alpha and therefore the classification is shown for each value of alpha.

\[ \text{nsamp} \]
If a scalar, it contains the number of subsamples which will be extracted. If \( \text{nsamp} = 0 \) all subsets will be extracted. Remark - if the number of all possible subset is greater than 300 the default is to extract all subsets, otherwise just 300. If \( \text{nsamp} \) is a matrix it contains in the rows the indexes of the subsets which have to be extracted. \( \text{nsamp} \) in this case can be conveniently generated by function subsets(). \( \text{nsamp} \) can have k columns or \( k \times (p + 1) \) columns. If \( \text{nsamp} \) has k columns the k initial centroids each iteration i are given by \( X[\text{nsamp}[i, \ ]] \) and the covariance matrices are equal to the identity.

If \( \text{nsamp} \) has \( k \times (p + 1) \) columns, the initial centroids and covariance matrices in iteration \( i \) are computed as follows:

• \( X1 \leftarrow X[\text{nsamp}[i, \ ]] \)
  • mean(\( X1[1:p + 1, \] \)) contains the initial centroid for group 1
  • cov(\( X1[1:p + 1, \] \)) contains the initial cov matrix for group 1
  • mean(\( X1[(p + 2):(2*p + 2), \] \)) contains the initial centroid for group 2
  • cov(\( X1[(p + 2):(2*p + 2), \] \)) contains the initial cov matrix for group 2
  • ...
  • mean(\( X1[(k-1)*p+1):(k*(p+1)), \] \)) contains the initial centroids for group k
  • cov(\( X1[(k-1)*p+1):(k*(p+1)), \] \)) contains the initial cov matrix for group k.

REMARK: If \( \text{nsamp} \) is not a scalar, the option startv1 given below is ignored. More precisely, if \( \text{nsamp} \) has k columns startv1 = 0 else if \( \text{nsamp} \) has \( k \times (p + 1) \) columns option startv1=1.

\[ \text{refsteps} \]
Number of refining iterations in each subsample. Default is \( \text{refsteps}=15 \).
\( \text{refsteps} = 0 \) means "raw-subsampling" without iterations.

\[ \text{reftol} \]
Tolerance of the refining steps. The default value is \( 1e-14 \)
equalweights  A logical specifying whether cluster weights in the concentration and assignment steps shall be considered. If equalweights=TRUE we are (ideally) assuming equally sized groups, else if equalweights = false (default) we allow for different group weights. Please, check in the given references which functions are maximized in both cases.

mixt  Specifies whether mixture modelling or crisp assignment approach to model based clustering must be used. In the case of mixture modelling parameter mixt also controls which is the criterion to find the untrimmed units in each step of the maximization. If mixt >=1 mixture modelling is assumed else crisp assignment. The default value is mixt=0, i.e. crisp assignment. Please see for details the provided references. The parameter mixt also controls the criterion to select the units to trim. If mixt = 2 the h units are those which give the largest contribution to the likelihood, else if mixt=1 the criterion to select the h units is exactly the same as the one which is used in crisp assignment.

msg  Controls whether to display or not messages on the screen. If msg==TRUE messages are displayed on the screen. If msg=2, detailed messages are displayed, for example the information at iteration level.

nocheck  Check input arguments. If nocheck=TRUE no check is performed on matrix X. The default nocheck=FALSE.

startv1  How to initialize centroids and covariance matrices. Scalar. If startv1=1 then initial centroids and covariance matrices are based on (p+1) observations randomly chosen, else each centroid is initialized taking a random row of input data matrix and covariance matrices are initialized with identity matrices. The default value is startv1=1.

Remark 1: in order to start with a routine which is in the required parameter space, eigenvalue restrictions are immediately applied.

Remark 2 - option startv1 is used just if nsamp is a scalar (see for more details the help associated with nsamp).

RandNumbForNini  pre-extracted random numbers to initialize proportions. Matrix of size k-by-nrow(nsamp) containing the random numbers which are used to initialize the proportions of the groups. This option is effective just if nsamp is a matrix which contains pre-extracted subsamples. The purpose of this option is to enable to user to replicate the results in case routine tclustreg*() is called using a parfor instruction (as it happens for example in routine IC, where tclustreg() is called through a parfor for different values of the restriction factor). The default is that RandNumbForNini is empty - then uniform random numbers are used.

restrtype  Type of restriction to be applied on the cluster scatter matrices. Valid values are 'eigen' (default), or 'deter'. "eigen" implies restriction on the eigenvalues while "deter" implies restriction on the determinants.

UnitsSameGroup  List of the units which must (whenever possible) have a particular label. For example UnitsSameGroup=c(20, 26), means that group which contains unit 20 is always labelled with number 1. Similarly, the group which contains unit 26 is always labelled with number 2, (unless it is found that unit 26 already belongs to group 1). In general, group which contains unit UnitsSameGroup(r)
where \( r=2, \ldots, \text{length}(kk)-1 \) is labelled with number \( r \) (unless it is found that unit \( \text{UnitsSameGroup}(r) \) has already been assigned to groups 1, 2, \ldots, \( r-1 \)).

**numpool** The number of parallel sessions to open. If numpool is not defined, then it is set equal to the number of physical cores in the computer.

**cleanpool** Logical, indicating if the open pool must be closed or not. It is useful to leave it open if there are subsequent parallel sessions to execute, so that to save the time required to open a new pool.

**trace** Whether to print intermediate results. Default is \( \text{trace}=\text{FALSE} \).

... potential further arguments passed to lower level functions.

**Details**

This iterative algorithm initializes \( k \) clusters randomly and performs concentration steps in order to improve the current cluster assignment. The number of maximum concentration steps to be performed is given by input parameter refsteps. For approximately obtaining the global optimum, the system is initialized \( nsamp \) times and concentration steps are performed until convergence or refsteps is reached. When processing more complex data sets higher values of \( nsamp \) and refsteps have to be specified (obviously implying extra computation time). However, if more than 10 per cent of the iterations do not converge, a warning message is issued, indicating that \( nsamp \) has to be increased.

**Value**

Depending on the input parameter monitoring, one of the following objects will be returned:

1. **tclustfsda.object**
2. **tclusteda.object**

**Author(s)**

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**References**


**Examples**

```r
## Not run:
data(hbk, package="robustbase")
(out <- tclustfsda(hbk[, 1:3], k=2))
class(out)
summary(out)

## TCLUST of Gayser data with three groups (k=3), 10% trimming (alpha=0.1)
## and restriction factor (c=10000)
```
data(geyser2)
(out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000))

## Use the plot options to produce more complex plots

## Plot with all default options
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000, plot=TRUE)

## Default confidence ellipses.
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000, plot="ellipse")

## Confidence ellipses specified by the user: confidence ellipses set to 0.5
plots <- list(type="ellipse", conflev=0.5)
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000, plot=plots)

## Confidence ellipses set to 0.9
plots <- list(type="ellipse", conflev=0.9)
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000, plot=plots)

## Contour plots
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000, plot="contour")

## Filled contour plots with additional options: contourf plot with a named colormap.
## Here we define four MATLAB-like colormaps, but the user can define anything else,
## presented by a matrix with three columns which are the RGB triplets.

summer <- as.matrix(data.frame(x1=seq(from=0, to=1, length=65),
                x2=seq(from=0.5, to=1, length=65),
                x3=rep(0.4, 65)))

spring <- as.matrix(data.frame(x1=rep(1, 65),
                x2=seq(from=0, to=1, length=65),
                x3=seq(from=1, to=0, length=65)))

winter <- as.matrix(data.frame(x1=rep(0, 65),
                x2=seq(from=0, to=1, length=65),
                x3=seq(from=1, to=0, length=65)))

autumn <- as.matrix(data.frame(x1=rep(1, 65),
                x2=seq(from=0, to=1, length=65),
                x3=rep(0, 65)))

out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000,
      plot=list(type="contourf", cmap=autumn))
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000,
      plot=list(type="contourf", cmap=winter))
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000,
      plot=list(type="contourf", cmap=summer))

## We compare the output using three different values of restriction factor
## nsamp is the number of subsamples which will be extracted
data(geyser2)
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10000, nsamp=500, plot="ellipse")
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=10, nsamp=500, refsteps=10, plot="ellipse")
out <- tclustfsda(geyser2, k=3, alpha=0.1, restrfactor=1, nsamp=500, refsteps=10, plot="ellipse")

### TCLUS applied to M5 data: A bivariate data set obtained from three normal
### bivariate distributions with different scales and proportions 1:2:2. One of the
### components is very overlapped with another one. A 10 per cent background noise is
### added uniformly distributed in a rectangle containing the three normal components
### and not very overlapped with the three mixture components. A precise description
### of the M5 data set can be found in Garcia-Escudero et al. (2008).
###
### data(M5data)
plot(M5data[, 1:2])
### Scatter plot matrix
library(rrcov)
plot(CovClassic(M5data[,1:2]), which="pairs")

out <- tclustfsda(M5data[,1:2], k=3, alpha=0, restrfactor=1000, nsamp=100, plot=TRUE)
out <- tclustfsda(M5data[,1:2], k=3, alpha=0, restrfactor=10, nsamp=100, plot=TRUE)
out <- tclustfsda(M5data[,1:2], k=3, alpha=0.1, restrfactor=1, nsamp=1000, plot=TRUE, equalweights=TRUE)
out <- tclustfsda(M5data[,1:2], k=3, alpha=0.1, restrfactor=1000, nsamp=100, plot=TRUE)

### TCLUS with simulated data: 5 groups and 5 variables
###
n1 <- 100
n2 <- 80
n3 <- 50
n4 <- 80
n5 <- 70
p <- 5
Y1 <- matrix(rnorm(n1 * p) + 5, ncol=p)
Y2 <- matrix(rnorm(n2 * p) + 3, ncol=p)
Y3 <- matrix(rnorm(n3 * p) - 2, ncol=p)
Y4 <- matrix(rnorm(n4 * p) + 2, ncol=p)
Y5 <- matrix(rnorm(n5 * p), ncol=p)

group <- c(rep(1, n1), rep(2, n2), rep(3, n3), rep(4, n4), rep(5, n5))
Y <- Y1
Y <- rbind(Y, Y2)
Y <- rbind(Y, Y3)
Y <- rbind(Y, Y4)
Y <- rbind(Y, Y5)
dim(Y)
table(group)
out <- tclustfsda(Y, k=5, alpha=0.05, restrfactor=1.3, refsteps=20, plot=TRUE)

### Automatic choice of best number of groups for Geyser data ------------------------
###
data(geyser2)
maxk <- 6
CLACLA <- matrix(0, nrow=maxk, ncol=2)
CLACLA[,1] <- 1:maxk
MIXCLA <- MIXMIX <- CLACLA

for(j in 1:maxk) {
  out <- tclustfsda(geyser2, k=j, alpha=0.1, restrfactor=5)
  CLACLA[j, 2] <- out$CLACLA
}

for(j in 1:maxk) {
  out <- tclustfsda(geyser2, k=j, alpha=0.1, restrfactor=5, mixt=2)
  MIXMIX[j, 2] <- out$MIXMIX
  MIXCLA[j, 2] <- out$MIXCLA
}

oldpar <- par(mfrow=c(1,3))
plot(CLACLA[,1:2], type="l", xlim=c(1, maxk), xlab="Number of groups", ylab="CLACLA")
plot(MIXMIX[,1:2], type="l", xlim=c(1, maxk), xlab="Number of groups", ylab="MIXMIX")
plot(MIXCLA[,1:2], type="l", xlim=c(1, maxk), xlab="Number of groups", ylab="MIXCLA")
par(oldpar)

## Monitoring examples ------------------------------------------

## Monitoring using Geyser data

## Monitoring using Geyser data (all default options)
## alpha and restriction factor are not specified therefore alpha=c(0.10, 0.05, 0)
## is used while the restriction factor is set to c=12
out <- tclustfsda(geyser2, k=3, monitoring=TRUE)

## Monitoring using Geyser data with alpha and c specified.
## out <- tclustfsda(geyser2, k=3, restrfac=100, alpha=seq(0.10, 0, by=-0.01), monitoring=TRUE)

## Monitoring using Geyser data with plot argument specified as a list.
## The trimming levels to consider in this case are 31 values of alpha
## out <- tclustfsda(geyser2, k=3, restrfac=100, alpha=seq(0.30, 0, by=-0.01), monitoring=TRUE,
##plot=list(alphasel=c(0.2, 0.10, 0.05, 0.01)), trace=TRUE)

## Monitoring using Geyser data with argument UnitsSameGroup
## Make sure that group containing unit 10 is in a group which is labelled "group 1"
## and group containing unit 12 is in group which is labelled "group 2"
## Mixture model is used (mixt=2)
## out <- tclustfsda(geyser2, k=3, restrfac=100, alpha=seq(0.30, 0, by=-0.01), monitoring=TRUE,
mixt=2, UnitsSameGroup=c(10, 12), trace=TRUE)

## Monitoring using M5 data
data(M5data)

## alphavec=vector which contains the trimming levels to consider
## in this case 31 values of alpha are considered
alphavec <- seq(0.10, 0, by=-0.02)
out <- tclustfsda(M5data[, 1:2], 3, alpha=alphavec, restrfac=1000, monitoring=TRUE,
                  nsamp=1000, plots=TRUE)

## End(Not run)

tclustfsda.object

Objects returned by the function tclustfsda

Description
An object of class tclustfsda.object holds information about the result of a call to tclustfsda.

Value
The functions print() and summary() are used to obtain and print a summary of the results. An object of class tclustfsda is a list containing at least the following components:

call the matched call
muopt a k-by-p matrix containing cluster centroid locations. Robust estimate of final centroids of the groups
sigmaopt a p-by-p-by-k array ray containing estimated constrained covariance for the k groups
idx a vector of length n containing assignment of each unit to each of the k groups. Cluster names are integer numbers from 1 to k. 0 indicates trimmed observations.
size a matrix of size (k+1)-by-3. The 1st col is sequence from 0 to k (cluster name); the 2nd col is the number of observations in each cluster; the 3rd col is the percentage of observations in each cluster. Remark: 0 denotes unassigned units.
postprob n-by-k matrix containing posterior probabilities. postprob[i, j] contains posterior probability of unit i from component (cluster) j. For the trimmed units posterior probabilities are 0.
emp "Empirical" statistics computed on final classification. When convergence is reached, emp=0. When convergence is not obtained, this field is a list which contains the statistics of interest: idxemp (ordered from 0 to k*, k* being the number of groups with at least one observation and 0 representing the possible group of outliers), muemp, sigmaemp and sizemp, which are the empirical counterparts of idx, muopt, sigmaopt and size.
MIXMIX BIC which uses parameters estimated using the mixture loglikelihood and the maximized mixture likelihood as goodness of fit measure. Remark: this output is present just if mixt > 0.
MIXCLA BIC which uses parameters estimated using the mixture loglikelihood and the maximized mixture likelihood as goodness of fit measure. Remark: this output is present just if mixt > 0.
tclustIC

Perform cluster analysis by calling tclustfsda for different number of groups k and restriction factors c

Description

Computes the values of BIC (MIXMIX), ICL (MIXCLA) or CLA (CLACLA), for different values of k (number of groups) and different values of c (restriction factor), for a prespecified level of trimming (the last two letters in the name stand for 'Information Criterion'). In order to minimize randomness, given k, the same subsets are used for each value of c.

Usage

tclustIC(
  x,
  kk = 1:5,
  cc = c(1, 2, 4, 8, 16, 32, 64, 128),
)
alpha = 0,
whichIC = c("ALL", "MIXMIX", "MIXCLA", "CLACLA"),
nsamp,
refsteps = 15,
reftol = 1e-14,
equalweights = FALSE,
msg = TRUE,
nocheck = FALSE,
plot = FALSE,
startv1 = 1,
restrtype = c("eigen", "deter"),
UnitsSameGroup,
numpool,
cleanpool,
trace = FALSE,
...
)

Arguments

x
An n x p data matrix (n observations and p variables). Rows of x represent observations, and columns represent variables. Missing values (NA's) and infinite values (Inf’s) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

kk
an integer vector specifying the number of mixture components (clusters) for which the BIC is to be calculated. By default kk=1:5.

cc
an vector specifying the values of the restriction factor which have to be considered. By default cc=c(1, 2, 4, 8, 16, 32, 64, 128).

alpha
Global trimming level. A scalar between 0 and 0.5 or an integer specifying the number of observations which have to be trimmed. If alpha=0 all observations are considered. By default alpha=0.

More in detail, if 0 < alpha < 1 clustering is based on h = fix(n * (1-alpha)) observations, else if alpha is an integer greater than 1 clustering is based on h = n - floor(alpha).

whichIC
A character value which specifies which information criteria must be computed for each k (number of groups) and each value of the restriction factor c. Possible values for whichIC are:

- "MIXMIX": a mixture model is fitted and for computing the information criterion the mixture likelihood is used. This option corresponds to the use of the Bayesian Information criterion (BIC). In output just the matrix MIXMIX is given.
- "MIXCLA": a mixture model is fitted but to compute the information criterion the classification likelihood is used. This option corresponds to the use of the Integrated Complete Likelihood (ICL). In the output just the matrix MIXCLA is given.
"CLACLA": everything is based on the classification likelihood. This information criterion will be called CLA. In the output just the matrix CLACLA is given.

"ALL": both classification and mixture likelihood are used. In this case all three information criteria CLA, ICL and BIC are computed. In the output all three matrices MIXMIX, MIXCLA and CLACLA are given.

**nsamp**
If a scalar, it contains the number of subsamples which will be extracted. If nsamp = 0 all subsets will be extracted. Remark - if the number of all possible subset is greater than 300 the default is to extract all subsets, otherwise just 300. If nsamp is a matrix it contains in the rows the indexes of the subsets which have to be extracted. nsamp in this case can be conveniently generated by function subsets(). nsamp can have k columns or k * (p + 1) columns. If nsamp has k columns the k initial centroids each iteration i are given by X[nsamp[i,] ,] and the covariance matrices are equal to the identity.

If nsamp has k * (p + 1) columns, the initial centroids and covariance matrices in iteration i are computed as follows:

- X1 <- X[nsamp[i ,] ,]
- mean(X1[1:p + 1, ]) contains the initial centroid for group 1
- cov(X1[1:p + 1, ]) contains the initial cov matrix for group 1
- mean(X1[(p + 2):(2*p + 2), ]) contains the initial centroid for group 2
- cov(X1[(p + 2):(2*p + 2), ]) contains the initial cov matrix for group 2
- ...
- mean(X1[(k-1)*p+1):(k*(p+1), ]) contains the initial centroids for group k
- cov(X1[(k-1)*p+1):(k*(p+1), ]) contains the initial cov matrix for group k.

REMARK: If nsamp is not a scalar, the option startv1 given below is ignored. More precisely, if nsamp has k columns startv1 = 0 else if nsamp has k*(p+1) columns option startv1=1.

**refsteps**
Number of refining iterations in each subsample. Default is refsteps=15. refsteps = 0 means "raw-subsampling" without iterations.

**reftol**
Tolerance of the refining steps. The default value is 1e-14

**equalweights**
A logical specifying whether cluster weights in the concentration and assignment steps shall be considered. If equalweights=TRUE we are (ideally) assuming equally sized groups, else if equalweights = false (default) we allow for different group weights. Please, check in the given references which functions are maximized in both cases.

**msg**
Controls whether to display or not messages on the screen. If msg==TRUE (default) messages are displayed on the screen. If msg=2, detailed messages are displayed, for example the information at iteration level.

**nocheck**
Check input arguments. If nocheck=TRUE no check is performed on matrix X. The default nocheck=FALSE.

**plot**
If plot=TRUE, a plot of the BIC (MIXMIX), ICL (MIXCLA) curve and CLA-CLA is shown on the screen. The plots which are shown depend on the input option whichIC.
**startv1**
How to initialize centroids and covariance matrices. Scalar. If startv1=1 then initial centroids and covariance matrices are based on \((p+1)\) observations randomly chosen, else each centroid is initialized taking a random row of input data matrix and covariance matrices are initialized with identity matrices. The default value is startv1=1.

Remark 1: in order to start with a routine which is in the required parameter space, eigenvalue restrictions are immediately applied.

Remark 2: option startv1 is used just if nsamp is a scalar (see for more details the help associated with nsamp).

**restrtype**
Type of restriction to be applied on the cluster scatter matrices. Valid values are 'eigen' (default), or 'deter'. "eigen" implies restriction on the eigenvalues while "deter" implies restriction on the determinants.

**UnitsSameGroup**
List of the units which must (whenever possible) have a particular label. For example UnitsSameGroup=c(20, 26), means that group which contains unit 20 is always labelled with number 1. Similarly, the group which contains unit 26 is always labelled with number 2, (unless it is found that unit 26 already belongs to group 1). In general, group which contains unit UnitsSameGroup(r) where \(r=2, \ldots, \text{length(kk)}-1\) is labelled with number \(r\) (unless it is found that unit UnitsSameGroup(r) has already been assigned to groups 1, 2, \ldots, \(r-1\)).

**numpool**
The number of parallel sessions to open. If numpool is not defined, then it is set equal to the number of physical cores in the computer.

**cleanpool**
Logical, indicating if the open pool must be closed or not. It is useful to leave it open if there are subsequent parallel sessions to execute, so that to save the time required to open a new pool.

**trace**
Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

**Value**
An S3 object of class `tclustic.object`

**Author(s)**
FSDA team, <valentin.todorov@chello.at>

**References**

**See Also**
`tclustfsda, tclustICplot, tclustICsol, carbikeplot`
Examples

```r
## Not run:
data(geyser2)
(out <- tclustIC(geyser2, whichIC="MIXMIX", plot=FALSE, alpha=0.1))
summary(out)

## End(Not run)

## Not run:
data(flea)
Y <- as.matrix(flea[, 1:(ncol(flea)-1)])  # select only the numeric variables
rownames(Y) <- 1:nrow(Y)
head(Y)

(out <- tclustIC(Y, whichIC="CLACLA", plot=FALSE, alpha=0.1, nsamp=100, numpool=1))
summary(out)

## End(Not run)
```

tclustic.object  

Objects returned by the function tclustIC

Description

An object of class `tclustic.object` holds information about the result of a call to `tclustIC`.

Value

The functions `print()` and `summary()` are used to obtain and print a summary of the results. An object of class `tclustic` is a list containing at least the following components:

- **call**: the matched call
- **kk**: a vector containing the values of k (number of components) which have been considered. This vector is identical to the optional argument `kk` (default is `kk=1:5`).
- **cc**: a vector containing the values of c (values of the restriction factor) which have been considered. This vector is identical to the optional argument `cc` (default is `cc=c(1, 2, 4, 8, 16, 32, 64, 128)`).
- **alpha**: trimming level
- **whichIC**: Information criteria used
- **CLACLA**: a matrix of size `length(kk)`-times-`length(cc)` containing the value of the penalized classification likelihood. This output is present only if `whichIC="CLACLA"` or `whichIC="ALL"`.
- **IDXCLA**: a matrix of lists of size `length(kk)`-times-`length(cc)` containing the assignment of each unit using the classification model. This output is present only if `whichIC="CLACLA"` or `whichIC="ALL"`. 
MIXMIX  a matrix of size length(kk)-times-length(cc) containing the value of the penalized mixture likelihood. This output is present only if whichIC="MIXMIX" or whichIC="ALL".

IDXMIX  a matrix of lists of size length(kk)-times-length(cc) containing the assignment of each unit using the classification model. This output is present only if whichIC="MIXMIX" or whichIC="ALL".

MIXCLA  a matrix of size length(kk)-times-length(cc) containing the value of the ICL criterion. This output is present only if whichIC="MIXCLA" or whichIC="ALL".

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- tclustIC(hbk[, 1:3]))
class(out)
summary(out)
## End(Not run)
```

tclustICplot  

Plots information criterion as a function of c and k, based on the solutions obtained by tclustIC

Description

The function tclustICplot() takes as input an object of class tclustic.object, the output of function tclustIC (that is a series of matrices which contain the values of the information criteria BIC/ICL/CLA for different values of k and c) and plots them as function of c or of k. The plot enables interaction in the sense that, if option databrush has been activated, it is possible to click on a point in the plot and to see the associated classification in the scatter plot matrix.

Usage

tclustICplot(
  out,
  whichIC = c("ALL", "MIXMIX", "MIXCLA", "CLAACL"),
  tag,
  datatooltip,
  databrush,
  nameY,
  trace = FALSE,
  ...
)
Arguments

out
An S3 object of class `tclustic.object` (output of `tclustIC`) containing the values of the information criteria BIC (MIXMIX), ICL (MIXCLA) or CLA (CLACLA), for different values of k (number of groups) and different values of c (restriction factor), for a prespecified level of trimming.

whichIC
Specifies the information criterion to use for the plot. See `tclustIC()` for the possible values of whichIC.

tag
plot handle. String which identifies the handle of the plot which is about to be created. The default is to use tag 'pl_IC'. Notice that if the program finds a plot which has a tag equal to the one specified by the user, then the output of the new plot overwrites the existing one in the same window else a new window is created.

datatooltip
Interactive clicking. It is inactive if this parameter is set to FALSE. If datatooltip=TRUE, the user can select with the mouse a solution in order to have the following information:

• 1) value of k which has been selected
• 2) value of c which has been selected
• 3) values of the information criterion
• 4) frequency distribution of the associated classification.

If datatooltip is a list it may contain the following fields:

1. DisplayStyle determines how the data cursor displays. Possible values are 'datatip' and 'window' (default). 'datatip' displays data cursor information in a small yellow text box attached to a black square marker at a data point you interactively select. 'window' displays data cursor information for the data point you interactively select in a floating window within the figure.

2. SnapToDataVertex: specifies whether the data cursor snaps to the nearest data value or is located at the actual pointer position. Possible values are SnapToDataVertex='on' (default) and SnapToDataVertex='off'.

databrush
Interactive mouse brushing. If databrush is missing or empty (default), no brushing is done. The activation of this option (databrush is TRUE or a list) enables the user to select a set of values of IC in the current plot and to see the corresponding classification highlighted in the scatterplot matrix. If the scatterplot matrix does not exist it is automatically created. Note that the window style of the other figures is set equal to that which contains the IC plot. In other words, if the IC plot is docked all the other figures will be docked too.

If databrush=TRUE the default selection tool is a rectangular brush and it is possible to brush only once (that is persist=""). If databrush=list(...) it is possible to use all optional arguments of the MATLAB function `selectdataFS()` and the following optional arguments:

• persist: Persist is an empty value or a character containing 'on' or 'off'. The default value is persist="", that is brushing is allowed only once. If persist="on" or persist="off" brushing can be done as many time as the user requires. If persist="on" then the unit(s) currently brushed are added to those previously brushed. It is possible, every time a new brushing
is done, to use a different color for the brushed units. If persist='off' every time a new brush is performed units previously brushed are removed.

- label: add labels of brushed units in the monitoring plot.
- dispopt: controls how to fill the diagonals in the scatterplot matrix of the brushed solutions. Set dispopt="hist" (default) to plot histograms, or dispopt="box" to plot boxplots.

nameY: Add variable labels in plot. A vector of strings of length p containing the labels of the variables in the dataset. If it is empty (default) the sequence X1, ..., Xp will be created automatically

trace: Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

Author(s)

FSDA team, <valentin.todorov@chello.at>

References


See Also

tclustIC, tclustfsda

Examples

```r
## Not run:
data(geyser2)
out <- tclustIC(geyser2, whichIC="MIXMIX", plot=FALSE, alpha=0.1)
tclustICplot(out, whichIC="MIXMIX")
```

```r
## End(Not run)
```

---

**tclustICsol**

Extracts a set of best relevant solutions obtained by *tclustIC*
Description

The function tclustICsol() takes as input an object of class tclustic.object, the output of function tclustIC (that is a series of matrices which contain the values of the information criteria BIC/ICL/CLA for different values of k and c) and extracts the first best solutions. Two solutions are considered equivalent if the value of the adjusted Rand index (or the adjusted Fowlkes and Mallows index) is above a certain threshold. For each tentative solution the program checks the adjacent values of c for which the solution is stable. A matrix with adjusted Rand indexes is given for the extracted solutions.

Usage

```
tclustICsol(
  out,
  NumberOfBestSolutions = 5,
  ThreshRandIndex = 0.7,
  whichIC = c("ALL", "CLACLA", "MIXMIX", "MIXCLA"),
  Rand = TRUE,
  msg = TRUE,
  plot = FALSE,
  trace = FALSE,
  ...
)
```

Arguments

- `out` An S3 object of class tclustic.object (output of tclustIC) containing the values of the information criteria BIC (MIXMIX), ICL (MIXCLA) or CLA (CLACLA), for different values of k (number of groups) and different values of c (restriction factor), for a prespecified level of trimming.
- `NumberOfBestSolutions` Number of best solutions to extract from BIC/ICL matrix. The default value of NumberOfBestSolutions is 5
- `ThreshRandIndex` Threshold to identify spurious solutions - the threshold of the adjusted Rand index to use to consider two solutions as equivalent. The default value of ThreshRandIndex is 0.7
- `whichIC` Specifies the information criterion to use to extract best solutions. Possible values for whichIC are:
  - CLACLA = in this case best solutions are referred to the classification likelihood.
  - MIXMIX = in this case in case best solutions are referred to the mixture likelihood (BIC).
  - MIXCLA = in this case in this case best solutions are referred to ICL.
  - ALL = in this case best solutions both three solutions using classification and mixture likelihood are produced. In the output class out all the three matrices MIXMIXbs, CLACLABs and MIXCLAb are given.

The default value is whichIC="ALL".
Index to use to compare partitions. If Rand=TRUE (default) the adjusted Rand index is used, else the adjusted Fowlkes and Mallows index is used.

msg

It controls whether to display or not messages (from MATLAB) on the screen. If msg=TRUE (default) messages about the progression of the search are displayed on the screen otherwise only error messages will be displayed.

plot

If plot=TRUE, the best solutions which have been found are shown on the screen.

trace

Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

Value

An S3 object of class tclusticsol.object

Author(s)

FSDA team, <valentin.todorov@chello.at>

References


See Also

tclustIC, tclustfsda, carbikeplot

Examples

```r
## Not run:
data(geyser2)
out <- tclustIC(geyser2, whichIC="MIXMIX", plot=FALSE, alpha=0.1)

## Plot first two best solutions using as Information criterion MIXMIX
print("Best solutions using MIXMIX")
outMIXMIX <- tclustICsol(out, whichIC="MIXMIX", plot=TRUE, NumberOfBestSolutions=2)
print(outMIXMIX$MIXMIXbs)

## End(Not run)
```
**tclusticsol.object**

**Objects returned by the function tclustICsol**

**Description**

An object of class `tclusticsol.object` holds information about the result of a call to `tclustICsol`.

**Value**

The functions `print()` and `summary()` are used to obtain and print a summary of the results. An object of class `tclusticsol` is a list containing at least the following components:

- **call**
  - the matched call

- **kk**
  - a vector containing the values of k (number of components) which have been considered. This vector is identical to the optional argument `kk` (default is `kk=1:5`).

- **cc**
  - a vector containing the values of c (values of the restriction factor) which have been considered. This vector is identical to the optional argument `cc` (default is `cc=c(1, 2, 4, 8, 16, 32, 64, 128)`).

- **alpha**
  - trimming level

- **whichIC**
  - Information criteria used

- **MIXMIXbs**
  - a matrix of lists of size `NumberOfBestSolutions`-times-`5` which contains the details of the best solutions for MIXMIX (BIC). Each row refers to a solution. The information which is stored in the columns is as follows.
    - 1st col = value of k for which solution takes place
    - 2nd col = value of c for which solution takes place;
    - 3rd col = a vector of length `d` which contains the values of `c` for which the solution is uniformly better.
    - 4th col = a vector of length `d + r` which contains the values of `c` for which the solution is considered stable (i.e. for which the value of the adjusted Rand index (or the adjusted Fowlkes and Mallows index) does not go below the threshold defined in input option `ThreshRandIndex`).
    - 5th col = string which contains 'true' or 'spurious'. The solution is labelled spurious if the value of the adjusted Rand index with the previous solutions is greater than `ThreshRandIndex`.

Remark: the field `MIXMIXbs` is present only if `whichIC=ALL` or `whichIC="MIXMIX"`.

- **MIXMIXbsari**
  - a matrix of adjusted Rand indexes (or Fowlkes and Mallows indexes) associated with the best solutions for MIXMIX. A matrix of size `NumberOfBestSolutions`-times-`NumberOfBestSolutions` whose i,j-th entry contains the adjusted Rand index between classification produced by solution i and solution j, i,j=1,2, ..., `NumberOfBestSolutions`.

Remark: the field `MIXMIXbsari` is present only if `whichIC=ALL` or `whichIC="MIXMIX"`. 
tclustreg


Remark: the field ARIMIX is present only if whichIC=ALL or whichIC="MIXMIX" or whichIC="MIXCLA".

MIXCLAbs has the same structure as MIXMIXbs but refers to MIXCLA.

Remark: the field MIXCLAbs is present only if whichIC=ALL or whichIC="MIXCLA".

MIXCLAbsari has the same structure as MIXMIXbsari but refers to MIXCLA.

Remark: the field MIXCLAbsari is present only if whichIC=ALL or whichIC="MIXCLA".

CLACLAbs has the same structure as MIXMIXbs but refers to CLACLA.

Remark: the field CLACLAbs is present only if whichIC=ALL or whichIC="CLACLA".

CLACLAbsari has the same structure as MIXMIXbsari but refers to CLACLA.

Remark: the field CLACLAbsari is present only if whichIC=ALL or whichIC="CLACLA".


Remark: the field ARICLA is present only if whichIC=ALL or whichIC="CLACLA".

See Also
tclustICsol, carbikeplot

Examples

```r
## Not run:
data(hbk, package="robustbase")
(out <- tclustIC(hbk[,1:3]))

## Plot first two best solutions using as Information criterion MIXMIX
print("Best solutions using MIXMIX")
outMIXMIX <- tclustICsol(out, whichIC="MIXMIX", plot=TRUE, NumberOfBestSolutions=2)
class(outMIXMIX)
summary(outMIXMIX)
print(outMIXMIX$MIXMIXbs)

## End(Not run)
```

---

Computes robust linear grouping analysis

**Description**

Performs robust linear grouping analysis.
Usage

tclustreg(
  y,
  x,
  k,
  alphaLik,
  alphaX,
  restrfactor = 12,
  intercept = TRUE,
  plot = FALSE,
  nsamp,
  refsteps = 10,
  reftol = 1e-13,
  equalweights = FALSE,
  mixt = 0,
  wtrim = 0,
  we,
  msg = TRUE,
  RandNumbForNini,
  trace = FALSE,
  ...
)

Arguments

y Response variable. A vector with n elements that contains the response variable.

x An n x p data matrix (n observations and p variables). Rows of x represent
observations, and columns represent variables.
Missing values (NA’s) and infinite values (Inf’s) are allowed, since observations
(rows) with missing or infinite values will automatically be excluded from the
computations.

k Number of groups.

alphaLik Trimming level, a scalar between 0 and 0.5 or an integer specifying the number
of observations which have to be trimmed. If alphaLik=0, there is no trimming.
More in detail, if 0 < alphaLik < 1 clustering is based on h = floor(n * (1 -
alphaLik)) observations. If alphaLik is an integer greater than 1 clustering is
based on h = n - floor(alphaLik). More in detail, likelihood contributions are
sorted and the units associated with the smallest n - h contributions are trimmed.

alphaX Second-level trimming or constrained weighted model for x.

restrfactor Restriction factor for regression residuals and covariance matrices of the ex-
planatory variables. Scalar or vector with two elements. If restrfactor is
a scalar it controls the differences among group scatters of the residuals. The
value 1 is the strongest restriction. If restrfactor is a vector with two el-
ments the first element controls the differences among group scatters of the
residuals and the second the differences among covariance matrices of the ex-
planatory variables. Note that restrfactor[2] is used just if alphaX=1, that is
if constrained weighted model for x is assumed.
intercept  wheather to use constant term (default is intercept=TRUE
plot    If plot=FALSE (default) or plot=0 no plot is produced. If plot=TRUE a plot with
        the final allocation is shown (using the spmplot function). If X is 2-dimensional,
        the lines associated to the groups are shown too.
nsamp    If a scalar, it contains the number of subsamples which will be extracted. If
         nsamp = 0 all subsets will be extracted. Remark - if the number of all possible
         subset is greater than 300 the default is to extract all subsets, otherwise just 300.
         If nsamp is a matrix it contains in the rows the indexes of the subsets which have
         to be extracted. nsamp in this case can be conveniently generated by function
         subsets(). nsamp must have k * p columns. The first p columns are used to
         estimate the regression coefficient of group 1, ..., the last p columns are used to
         estimate the regression coefficient of group k.
refsteps  Number of refining iterations in each subsample. Default is refsteps=10. refsteps = 0
         means "raw-subsampling" without iterations.
reftol    Tolerance of the refining steps. The default value is 1e-14
equalweights A logical specifying wheather cluster weights in the concentration and assign-
       ment steps shall be considered. If equalweights=TRUE we are (ideally) assum-
       ing equally sized groups, else if equalweights = false (default) we allow for
       different group weights. Please, check in the given references which functions
       are maximized in both cases.
mixt    Specifies whether mixture modelling or crisp assignment approach to model
         based clustering must be used. In the case of mixture modelling parameter mixt
         also controls which is the criterion to find the untrimmed units in each step of the
         maximization. If mixt>=1 mixture modelling is assumed else crisp assignment.
         The default value is mixt=0, i.e. crisp assignment. Please see for details the
         provided references. The parameter mixt also controls the criterion to select the
         units to trim. If mixt = 2 the h units are those which give the largest contribution
         to the likelihood, else if mixt=1 the criterion to select the h units is exactly the
         same as the one which is used in crisp assignment.
wtrim    How to apply the weights on the observations - a flag taking values in c(0, 1, 2,
         3, 4).
         • If wtrim==0 (no weights), the algorithm reduces to the standard tclustreg
           algorithm.
         • If wtrim==1, trimming is done by weighting the observations using values
           specified in vector we. In this case, vector we must be supplied by the user.
         • If wtrim==2, trimming is again done by weighting the observations using
           values specified in vector we. In this case, vector we is computed from the
           data as a function of the density estimate pdfe. Specifically, the weight of
           each observation is the probability of retaining the observation, computed
           as
           \[ \text{pretain}_{ig} = 1 - \frac{\text{pdfe}_{ig}}{\text{max}_{ig}(\text{pdfe}_{ig})} \]
         • If wtrim==3, trimming is again done by weighting the observations using
           values specified in vector we. In this case, each element wei of vector we is
           a Bernoulli random variable with probability of success pdfe_{ig}. In the clus-
           tering framework this is done under the constraint that no group is empty.
• If wtrim==4, trimming is done with the tandem approach of Cerioli and Perrotta (2014).

\texttt{w}

Weights. A vector of size n-by-1 containing application-specific weights. Default is a vector of ones.

\texttt{msg}

Controls whether to display or not messages on the screen. If msg==TRUE (default), messages are displayed on the screen. If msg=2, detailed messages are displayed, for example the information at iteration level.

\texttt{RandNumbForNini}

pre-extracted random numbers to initialize proportions. Matrix of size k-by-nrow(nsamp) containing the random numbers which are used to initialize the proportions of the groups. This option is effective only if nsamp is a matrix which contains pre-extracted subsamples. The purpose of this option is to enable the user to replicate the results when the function \texttt{tclustreg()} is called using a \texttt{parfor} instruction (as it happens for example in routine IC, where \texttt{tclustreg()} is called through a \texttt{parfor} for different values of the restriction factor). The default is that \texttt{RandNumbForNini} is empty - then uniform random numbers are used.

\texttt{trace}

Whether to print intermediate results. Default is trace=FALSE.

... potential further arguments passed to lower level functions.

\textbf{Value}

An S3 object of class \texttt{tclustreg.object}

\textbf{Author(s)}

FSDA team, <valentin.todorov@chello.at>

\textbf{References}


Examples

### Not run:
### The X data have been introduced by Gordaliza, Garcia-Escudero & Mayo-Iscar (2013).
### The dataset presents two parallel components without contamination.

data(X)
y1 = X[, ncol(X)]
X1 = X[, -ncol(X), drop=FALSE]

(out <- tclustreg(y1, X1, k=2, alphaLik=0.05, alphaX=0.01, restrfactor=5, plot=TRUE, trace=TRUE))

(out <- tclustreg(y1, X1, k=2, alphaLik=0.05, alphaX=0.01, restrfactor=2,
mixt=2, plot=TRUE, trace=TRUE))

### Examples with fishery data

data(fishery)
X <- fishery

### some jittering is necessary because duplicated units are not treated:
### this needs to be addressed
X <- X + 10^(-8) * abs(matrix(rnorm(nrow(X)*ncol(X)), ncol=2))

y1 <- X[, ncol(X)]
X1 <- X[, -ncol(X), drop=FALSE]

(out <- tclustreg(y1, X1, k=3, restrfact=50, alphaLik=0.04, alphaX=0.01, trace=TRUE))

### Example 2:
### Define some arbitrary weightssome arbitrary weights for the units
we <- sqrt(X1)/sum(sqrt(X1))

### tclustreg required parameters
k <- 2; restrfact <- 50; alpha1 <- 0.04; alpha2 <- 0.01

### Now tclust is run on each combination of mixt and wtrim options

cat("\nmixt=0; wtrim=0",
  "\nStandard tclustreg, with classification likelihood and without thinning\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=0, wtrim=0, trace=TRUE))

cat("\nmixt=2; wtrim=0",
  "\nMixture likelihood, no thinning\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=2, wtrim=0, trace=TRUE))

cat("\nmixt=0; wtrim=1",
  "\nClassification likelihood, thinning based on user weights\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=0, we=we, wtrim=1, trace=TRUE))
cat("\nmixt=2; wtrim=1",
"Mixture likelihood, thinning based on user weights\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=2, we=we, wtrim=1, trace=TRUE))

cat("\nmixt=0; wtrim=2",
"Classification likelihood, thinning based on retention probabilities\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=0, wtrim=2, trace=TRUE))

cat("\nmixt=2; wtrim=2",
"Mixture likelihood, thinning based on retention probabilities\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=2, wtrim=2, trace=TRUE))

cat("\nmixt=0; wtrim=3",
"Classification likelihood, thinning based on bernoulli weights\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=0, wtrim=3, trace=TRUE))

cat("\nmixt=2; wtrim=3",
"Mixture likelihood, thinning based on bernoulli weights\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=2, wtrim=3, trace=TRUE))

cat("\nmixt=0; wtrim=4",
"Classification likelihood, tandem thinning based on bernoulli weights\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=0, wtrim=4, trace=TRUE))

cat("\nmixt=2; wtrim=4",
"Mixture likelihood, tandem thinning based on bernoulli weights\n")
(out <- tclustreg(y1, X1, k=k, restrfact=restrfact, alphaLik=alpha1, alphaX=alpha2,
mixt=2, wtrim=4, trace=TRUE))

## End(Not run)

**tclustreg.object**  
*Objects returned by the function tclustreg*

**Description**

An object of class `tclustreg.object` holds information about the result of a call to `tclustreg`.

**Value**

The functions `print()` and `summary()` are used to obtain and print a summary of the results. An object of class `tclustreg` is a list containing at least the following components:

call the matched call
tclustregIC

See Also

\texttt{tclustreg}

Examples

```r
## Not run:

## The X data have been introduced by Gordaliza, Garcia-Escudero & Mayo-Iscar (2013).
## The dataset presents two parallel components without contamination.

data(X)
y1 = X[, ncol(X)]
X1 = X[,-ncol(X), drop=FALSE]

out <- tclustreg(y1, X1, k=2, alphaLik=0.05, alphaX=0.01, restrfactor=5, trace=TRUE)
class(out)
str(out)

## End(Not run)
```

tclustregIC

\textit{Computes} \texttt{tclustreg} \textit{for different number of groups} \texttt{k} \textit{and restriction factors} \texttt{c}.

Description

(\texttt{the last two letters stand for ‘Information Criterion’}) \textit{computes the values of BIC (MIXMIX), ICL (MIXCLA) or CLA (CLACLA), for different values of k (number of groups) and different values of c (restriction factor for the variances of the residuals), for a prespecified level of trimming. In order to minimize randomness, given k, the same subsets are used for each value of c.}

Usage

\begin{verbatim}
tclustregIC(  y,  x,  alphaLik,  alphaX,  intercept = TRUE,  plot = FALSE,  nsamp,  refsteps = 10,  reftol = 1e-13,  equalweights = FALSE,  wtrim = 0,  we,  msg = TRUE,  c)
\end{verbatim}
RandNumbForNini,
trace = FALSE,
...
)

Arguments

y
Response variable. A vector with \( n \) elements that contains the response variable.

x
An \( n \times p \) data matrix (\( n \) observations and \( p \) variables). Rows of \( x \) represent observations, and columns represent variables. Missing values (NA's) and infinite values (Inf's) are allowed, since observations (rows) with missing or infinite values will automatically be excluded from the computations.

alphaLik
Trimming level, a scalar between 0 and 0.5 or an integer specifying the number of observations which have to be trimmed. If \( \alpha_{\text{Lik}} = 0 \), there is no trimming. More in detail, if \( 0 < \alpha_{\text{Lik}} < 1 \) clustering is based on \( h = \text{floor}(n \times (1 - \alpha_{\text{Lik}})) \) observations. If \( \alpha_{\text{Lik}} \) is an integer greater than 1 clustering is based on \( h = n - \text{floor}(\alpha_{\text{Lik}}) \). More in detail, likelihood contributions are sorted and the units associated with the smallest \( n - h \) contributions are trimmed.

alphaX
Second-level trimming or constrained weighted model for \( x \).

intercept
Whether to use constant term (default is \( \text{intercept=TRUE} \)).

plot
If plot=FALSE (default) or plot=0 no plot is produced. If plot=TRUE a plot with the final allocation is shown (using the spmplot function). If \( X \) is 2-dimensional, the lines associated to the groups are shown too.

dsamp
If a scalar, it contains the number of subsamples which will be extracted. If \( \text{nsamp} = 0 \) all subsets will be extracted. Remark - if the number of all possible subset is greater than 300 the default is to extract all subsets, otherwise just 300. If \( \text{nsamp} \) is a matrix it contains in the rows the indexes of the subsets which have to be extracted. \( \text{nsamp} \) in this case can be conveniently generated by function \( \text{subsets()} \). \( \text{nsamp} \) must have \( k \times p \) columns. The first \( p \) columns are used to estimate the regression coefficient of group 1, ..., the last \( p \) columns are used to estimate the regression coefficient of group \( k \).

refsteps
Number of refining iterations in each subsample. Default is \( \text{refsteps}=10 \). \( \text{refsteps} = 0 \) means "raw-subsampling" without iterations.

reftol
Tolerance of the refining steps. The default value is \( 1 \times 10^{-14} \).

equalweights
A logical specifying whether cluster weights in the concentration and assignment steps shall be considered. If \( \text{equalweights=TRUE} \) we are (ideally) assuming equally sized groups, else if \( \text{equalweights = false} \) (default) we allow for different group weights. Please, check in the given references which functions are maximized in both cases.

wtrim
How to apply the weights on the observations - a flag taking values in \( \{0, 1, 2, 3, 4\} \).

- If \( \text{wtrim}=0 \) (no weights), the algorithm reduces to the standard tclustreg algorithm.
- If \( \text{wtrim}=1 \), trimming is done by weighting the observations using values specified in vector \( \text{we} \). In this case, vector \( \text{we} \) must be supplied by the user.
• If \( wtr\text{im}=2 \), trimming is again done by weighting the observations using values specified in vector \( \text{we} \). In this case, vector \( \text{we} \) is computed from the data as a function of the density estimate \( \text{pdf}e_{ig} \). Specifically, the weight of each observation is the probability of retaining the observation, computed as

\[
\text{pretain}_{ig} = 1 - \frac{\text{pdf}e_{ig}}{\text{max}_{ig}(\text{pdf}e_{ig})}
\]

• If \( wtr\text{im}=3 \), trimming is again done by weighting the observations using values specified in vector \( \text{we} \). In this case, each element \( \text{wei} \) of vector \( \text{we} \) is a Bernoulli random variable with probability of success \( \text{pdf}e_{ig} \). In the clustering framework this is done under the constraint that no group is empty.

• If \( wtr\text{im}=4 \), trimming is done with the tandem approach of Cerioli and Perrotta (2014).

\( \text{we} \) Weights. A vector of size n-by-1 containing application-specific weights Default is a vector of ones.

\( \text{msg} \) Controls whether to display or not messages on the screen If \( \text{msg}==\text{TRUE} \) (default) messages are displayed on the screen. If \( \text{msg}=2 \), detailed messages are displayed, for example the information at iteration level.

\( \text{RandNumbForNini} \) pre-extracted random numbers to initialize proportions. Matrix of size k-by-nrow(\( \text{nsamp} \)) containing the random numbers which are used to initialize the proportions of the groups. This option is effective only if \( \text{nsamp} \) is a matrix which contains pre-extracted subsamples. The purpose of this option is to enable the user to replicate the results when the function \( \text{tclustreg()} \) is called using a parfor instruction (as it happens for example in routine \( \text{IC} \), where \( \text{tclustreg()} \) is called through a parfor for different values of the restriction factor). The default is that \( \text{RandNumbForNini} \) is empty - then uniform random numbers are used.

\( \text{trace} \) Whether to print intermediate results. Default is \( \text{trace}=\text{FALSE} \).

... potential further arguments passed to lower level functions.

Value

An S3 object of class \( \text{tclustreg.object} \)

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References

wool  

Wool data.

Description

The wool data give the number of cycles to failure of a worsted yarn under cycles of repeated loading. The variables are: length of test specimen; amplitude of loading cycle; load.

Usage

data(wool)

Format

A data frame with 27 rows and 4 variables

X  

Simulated data X.

Description

The X dataset has been simulated by Gordaliza, Garcia-Escudero and Mayo-Iscar during the Workshop ADVANCES IN ROBUST DATA ANALYSIS AND CLUSTERING held in Ispra on October 21st-25th 2013. It is a bivariate dataset of 200 observations. It presents two parallel components without contamination.

Usage

data(X)

Format

A data frame with 200 rows and 2 variables
Description
Simulated data to test tclustIC() and tclustICsol(), carbike() functions

Usage
data(z1)

Format
A data frame with 150 rows and 2 variables. The variables are as follows:

- X1
- X2

References

Examples
data(z1)
head(z1)
## Not run:
(out <- tclustIC(z1, plots=FALSE, whichIC="CLACLA"))
(outCLACLA <- tclustICsol(out, whichIC="CLACLA", plot=FALSE))
carbikeplot(outCLACLA)

## End(Not run)
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