# Package: GSE (via r-universe)

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Type Package

<b>Title</b> Robust Estimation in the Presence of Cellwise and Casewise Contamination and Missing Data
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<b>Description</b> Robust Estimation of Multivariate Location and Scatter in the Presence of Cellwise and Casewise Contamination and Missing Data.
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# **Description**

This data set is taken from UCI repository, see reference. Past usage includes price prediction of cars using all numeric and boolean attributes (Kibler et al., 1989).

# Usage

data(auto)

## **Format**

A data frame with 205 observations on the following 26 variables, of which 15 are quantitative and 11 are categorical. The following description is extracted from UCI repository (Frank and Asuncion, 2010):

Normalized-losses the relative average loss payment per insured vehicle year; ranged from 65 to 256

Make Vehicle's make
Fuel-type diesel, gas
Aspiration std, turbo
Num-of-doors four, two

Body-style hardtop, wagon, sedan, hatchback, convertible

Drive-wheels 4wd, fwd, rwd Engine-location front, rear

Wheel-base continuous from 86.6 120.9
Length continuous from 141.1 to 208.1
Width continuous from 60.3 to 72.3
Height continuous from 47.8 to 59.8
Curb-weight continuous from 1488 to 4066

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Engine-type dohc, dohcv, l, ohc, ohcf, ohcv, rotor Num-of-cylinders eight, five, four, six, three, twelve, two

Engine-size continuous from 61 to 326

Fuel-system 1bbl, 2bbl, 4bbl, idi, mfi, mpfi, spdi, spfi

Bore continuous from 2.54 to 3.94 Stroke continuous from 2.07 to 4.17 Compression-ratio continuous from 7 to 23 continuous from 48 to 288 Horsepower Peak-rpm continuous from 4150 to 6600 continuous from 13 to 49 City-mpg Highway-mpg continuous from 16 to 54 continuous from 5118 to 45400 Price

Symboling assigned insurance risk rating: -3, -2, -1, 0, 1, 2, 3

#### Source

The original data have been taken from the UCI Repository Of Machine Learning Databases at

• http://archive.ics.uci.edu/ml/datasets/Automobile.

#### References

Frank, A. & Asuncion, A. (2010). UCI Machine Learning Repository [http://archive.ics.uci.edu/ml]. Irvine, CA: University of California, School of Information and Computer Science.

Kibler, D., Aha, D.W., & Albert,M. (1989). Instance-based prediction of real-valued attributes. Computational Intelligence, Vol 5, 51–57.

boston

Boston Housing Data

# Description

Housing data for 506 census tracts of Boston from the 1970 census. The dataframe boston contains the corrected data by Harrison and Rubinfeld (1979). The data was for a few minor errors and augmented with the latitude and longitude of the observations. The original data can be found in the references below.

# Usage

data(boston)

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## **Format**

The original data are 506 observations on 14 variables, medv being the target variable:

cmedv	corrected median value of owner-occupied homes in USD 1000's
crim	per capita crime rate by town
indus	proportion of non-retail business acres per town
nox	nitric oxides concentration (parts per 10 million)
rm	average number of rooms per dwelling
age	proportion of owner-occupied units built prior to 1940
dis	weighted distances to five Boston employment centres
rad	index of accessibility to radial highways
tax	full-value property-tax rate per USD 10,000
ptratio	pupil-teacher ratio by town
b	$1000(B-0.63)^2$ where B is the proportion of blacks by town
lstat	percentage of lower status of the population

#### Source

The original data have been taken from the UCI Repository Of Machine Learning Databases at

• http://www.ics.uci.edu/~mlearn/MLRepository.html,

the corrected data have been taken from Statlib at

• https://dasl.datadescription.com/ (originally downloaded from lib.stat.cmu.edu/DASL/)

See Statlib and references there for details on the corrections. Both were converted to R format by Friedrich Leisch.

## References

Harrison, D. and Rubinfeld, D.L. (1978). Hedonic prices and the demand for clean air. *Journal of Environmental Economics and Management*, **5**, 81–102.

Gilley, O.W., and R. Kelley Pace (1996). On the Harrison and Rubinfeld Data. *Journal of Environmental Economics and Management*, **31**, 403–405. [Provided corrections and examined censoring.]

Newman, D.J. & Hettich, S. & Blake, C.L. & Merz, C.J. (1998). UCI Repository of machine learning databases [http://www.ics.uci.edu/~mlearn/MLRepository.html]. Irvine, CA: University of California, Department of Information and Computer Science.

Pace, R. Kelley, and O.W. Gilley (1997). Using the Spatial Configuration of the Data to Improve Estimation. *Journal of the Real Estate Finance and Economics*, **14**, 333–340. [Added georeferencing and spatial estimation.]

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calcium	Calcium data	

## **Description**

The Calcium data is from the article by Holcomb and Spalsbury (2005). The dataset used for class was compiled by Boyd, Delost, and Holcomb (1998) for the use of a study to determine if significant gender differences existed between subjects 65 years of age and older with regard to calcium, inorganic phosphorous, and alkaline phosphatase levels. Although the original data from Boyd, Delost, and Holcomb (1998) had observations needing investigation, Holcomb and Spalsbury (2005) further massaged the original data to include data problems and issues that have arisen in other research projects for pedagogical purposes.

## Usage

data(calcium)

#### **Format**

A data frame with 178 observations on the following 8 variables.

obsno	Patient Observation Number
age	Age in years
sex	1=Male, 2=Female
alkphos	Alkaline Phosphatase International Units/Liter
lab	1=Metpath; 2=Deyor; 3=St. Elizabeth's; 4=CB Rouche; 5=YOH; 6=Horizon
cammol	Calcium mmol/L
phosmmol	Inorganic Phosphorus mmol/L
agegroup	Age group 1=65-69: 2=70-74: 3=75-79: 4=80-84: 5=85-89 Years

# Source

The original data have been taken from the Journal of Statistics Education Databases at

• http://jse.amstat.org/datasets/calcium.dat.txt (originally downloaded from www.amstat.org/publications/js

the corrected data have been taken from Statlib at

• http://jse.amstat.org/datasets/calciumgood.dat.txt (originally downloaded from www.amstat.org/publications/jse/datasets/calciumgood.dat.txt)

## References

Boyd, J., Delost, M., and Holcomb, J., (1998). Calcium, phosphorus, and alkaline phosphatase laboratory values of elderly subjects, *Clinical Laboratory Science*, 11, 223-227.

Holcomb, J., and Spalsbury, A. (2005), Teaching Students to Use Summary Statistics and Graphics to Clean and Analyze Data. *Journal of Statistics Education*, 13, Number 3.

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## **Examples**

```
## Not run:
data(calcium)
## remove the categorical variables
calcium.cts <- subset(calcium, select=-c(obsno, sex, lab, agegroup) )
res <- GSE(calcium.cts)
getOutliers(res)
## able to identify majority of the contaminated cases identified
## in the reference
## End(Not run)</pre>
```

CovEM

Gaussian MLE of mean and covariance

# Description

Computes the Gaussian MLE via EM-algorithm for missing data.

## Usage

```
CovEM(x, tol=0.001, maxiter=1000)
```

# **Arguments**

x a matrix or data frame. May contain missing values, but cannot contain columns with completely missing entries.

with completely imaging entires.

tol tolerance level for the maximum relative change of the estimates. Default is

0.001.

maxiter maximum iteration for the EM algorithm. Default is 1000.

#### Value

An S4 object of class CovRobMiss-class. The output S4 object contains the following slots:

mu	Estimated location. Can be accessed via getLocation.
S	Estimated scatter matrix. Can be accessed via getScatter.
pmd	Squared partial Mahalanobis distances. Can be accessed via getDist.
pmd.adj	Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj.
pu	Dimension of the observed entries for each case. Can be accessed via getDim.
call	Object of class "language". Not meant to be accessed.
X	Input data matrix. Not meant to be accessed.
р	Column dimension of input data matrix. Not meant to be accessed.
estimator	Character string of the name of the estimator used. Not meant to be accessed.

## Author(s)

Mike Danilov, Andy Leung <andy.leung@stat.ubc.ca>

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CovRobMiss-class	Class "CovRobMiss" – a superclass for the robust estimates of location and scatter for missing data
	tion and scatter for missing data

#### **Description**

The Superclass of all the objects output from the various robust estimators of location and scatter for missing data, which includes Generalized S-estimator GSE, Extended Minimum Volumn Ellipsoid emve, and Huberized Pairwise HuberPairwise. It can also be constructed using the code partial.mahalanobis.

## **Objects from the Class**

Objects can be created by calls of the form new("CovRobMiss", ...), but the best way of creating CovRobMiss objects is a call to either of the following functions:GSE, emve, HuberPairwise, and partial.mahalanobis, which all serve as a constructor.

#### Slots

```
mu Estimated location. Can be accessed via getLocation.

S Estimated scatter matrix. Can be accessed via getScatter.

pmd Square partial Mahalanobis distances. Can be accessed via getDist.

pmd.adj Adjusted square partial Mahalanobis distances. Can be accessed via getDistAdj.

pu Dimension of the observed entries for each case. Can be accessed via getDim.

call Object of class "language". Not meant to be accessed.
```

- x Input data matrix. Not meant to be accessed.
- p Column dimension of input data matrix. Not meant to be accessed. estimator Character string of the name of the estimator used. Not meant to be accessed.

## Methods

```
show signature(object = "CovRobMiss"): display the object
summary signature(object = "CovRobMiss"): calculate summary information
plot signature(object = "CovRobMiss", cutoff = "numeric"): plot the object. See plot
getDist signature(object = "CovRobMiss"): return the squared partial Mahalanobis distances
getDistAdj signature(object = "CovRobMiss"): return the adjusted squared partial Mahalanobis
    distances
getDim signature(object = "CovRobMiss"): return the dimension of observed entries for each
    case
getLocation signature(object = "CovRobMiss"): return the estimated location vector
getScatter signature(object = "CovRobMiss", cutoff = "numeric"): return the estimated scat-
ter matrix
```

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getMissing signature(object = "CovRobMiss"): return the case number with completely missing data, if any

getOutliers signature(object = "CovRobMiss", cutoff = "numeric"): return the case number(s) adjusted squared distances above (1 - cutoff)th quantile of chi-square p-degrees of freedom.

#### Author(s)

Andy Leung <andy.leung@stat.ubc.ca>

#### See Also

GSE, emve, HuberPairwise, partial.mahalanobis

CovRobMissSc-class

Class "CovRobMissSc" – a subclass of "CovRobMiss" with scale estimate

# **Description**

The Superclass of the GSE-class and emve-class objects.

# **Objects from the Class**

Objects can be created by calls of the form new("CovRobMissSc", ...), but the best way of creating CovRobMissSc objects is a call to either of the following functions:GSE or emve.

## Slots

- mu Estimated location. Can be accessed via getLocation.
- S Estimated scatter matrix. Can be accessed via getScatter.
- sc Estimated M-scale (either GS-scale or MVE-scale). Can be accessed via getScale.
- pmd Square partial Mahalanobis distances. Can be accessed via getDist.
- pmd.adj Adjusted square partial Mahalanobis distances. Can be accessed via getDistAdj.
- pu Dimension of the observed entries for each case. Can be accessed via getDim.
- call Object of class "language". Not meant to be accessed.
- x Input data matrix. Not meant to be accessed.
- p Column dimension of input data matrix. Not meant to be accessed.
- estimator Character string of the name of the estimator used. Not meant to be accessed.

## Extends

Class "CovRobMiss", directly.

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# Methods

In addition to methods inheritedfrom the class "CovRobMiss":

signature(object = "CovRobMissSc"): return the GS-scale or MVE-scale of the best candidate.

# Author(s)

getScale Andy Leung <andy.leung@stat.ubc.ca>

#### See Also

GSE, CovRobMiss-class

emve Extended Minimum Volume Ellipsoid (EMVE) in the presence of missing data	emve	1 / 1
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# Description

Computes the Extended S-Estimate (ESE) version of the minimum volume ellipsoid (EMVE), which is used as an initial estimator in Generlized S-Estimator (GSE) for missing data by default.

# Usage

```
emve(x, maxits=5, sampling=c("uniform","cluster"), n.resample, n.sub.size, seed)
```

# **Arguments**

X	a matrix or data frame. May contain missing values, but cannot contain columns with completely missing entries.
maxits	integer indicating the maximum number of iterations of Gaussian MLE calculation for each subsample. Default is 5.
sampling	which sampling scheme is to use: 'uniform' or 'cluster' (see Leung and Zamar, 2016). Default is 'uniform'.
n.resample	integer indicating the number of subsamples. Default is 15 for clustering-based subsampling and 500 for uniform subsampling.
n.sub.size	integer indicating the sizes of each subsample. Default is $2(p+1)/a$ for clustering-based subsampling and $(p+1)/a$ for uniform subsampling, where a is proportion of non-missing cells.
seed	optional starting value for random generator. Default is seed = 1000.

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#### **Details**

This function computes EMVE as described in Danilov et al. (2012). Two subsampling schemes can be used for computing EMVE: uniform subsampling and the clustering-based subsampling as described in Leung and Zamar (2016). For uniform subsampling, the number of subsamples must be large to ensure high breakdown point. For clustering-based subsampling, the number of subsamples can be smaller. The subsample size  $n_0$  must be chosen to be larger than p to avoid singularity.

In the algorithm, there exists a concentration step in which Gaussian MLE is computed for 50% of the data points using the classical EM-algorithm multiplied by a scalar factor. This step is repeated for each subsample. As the computation can be heavy as the number of subsample increases, we set by default the maximum number of iteration of classical EM-algorithm (i.e. maxits) as 5. Users are encouraged to refer to Danilov et al. (2012) for details about the algorithm and Rubin and Little (2002) for the classical EM-algorithm for missing data.

#### Value

An S4 object of class emve-class which is a subclass of the virtual class CovRobMissSc-class. The output S4 object contains the following slots:

mu	Estimated location. Can be accessed via getLocation.
S	Estimated scatter matrix. Can be accessed via getScatter.
sc	Estimated EMVE scale. Can be accessed via getScale.
pmd	Squared partial Mahalanobis distances. Can be accessed via getDist.
pmd.adj	Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj.
pu	Dimension of the observed entries for each case. Can be accessed via getDim.
call	Object of class "language". Not meant to be accessed.
X	Input data matrix. Not meant to be accessed.
р	Column dimension of input data matrix. Not meant to be accessed.
estimator	Character string of the name of the estimator used. Not meant to be accessed.

#### Author(s)

Andy Leung <andy.leung@stat.ubc.ca>, Ruben H. Zamar, Mike Danilov, Victor J. Yohai

#### References

Danilov, M., Yohai, V.J., Zamar, R.H. (2012). Robust Esimation of Multivariate Location and Scatter in the Presence of Missing Data. *Journal of the American Statistical Association* **107**, 1178–1186

Leung, A. and Zamar, R.H. (2016). Multivariate Location and Scatter Matrix Estimation Under Cellwise and Casewise Contamination. Submitted.

Rubin, D.B. and Little, R.J.A. (2002). *Statistical analysis with missing data* (2nd ed.). New York: Wiley.

#### See Also

```
GSE, emve-class
```

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emve-class	Extended Minimum Volume Ellipsoid (EMVE) in the presence of missing data.

#### Description

Class of Extended Minimum Volume Ellipsoid. It has the superclass of CovRobMissSc.

#### **Objects from the Class**

Objects can be created by calls of the form new("emve", ...), but the best way of creating emve objects is a call to the function emve which serves as a constructor.

#### Slots

```
mu Estimated location. Can be accessed via getLocation.

S Estimated scatter matrix. Can be accessed via getScatter.

sc Estimated EMVE scale. Can be accessed via getScale.

pmd Squared partial Mahalanobis distances. Can be accessed via getDist.

pmd.adj Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj.

pu Dimension of the observed entries for each case. Can be accessed via getDim.

call Object of class "language". Not meant to be accessed.

x Input data matrix. Not meant to be accessed.

p Column dimension of input data matrix. Not meant to be accessed.

estimator Character string of the name of the estimator used. Not meant to be accessed.
```

#### **Extends**

```
Class "CovRobMissSc", directly.
```

## Methods

The following methods are defined with the superclass "CovRobMiss":

```
show signature(object = "CovRobMiss"): display the object
summary signature(object = "CovRobMiss"): calculate summary information
plot signature(object = "CovRobMiss", cutoff = "numeric"): plot the object. See plot
getDist signature(object = "CovRobMiss"): return the squared partial Mahalanobis distances
getDistAdj signature(object = "CovRobMiss"): return the adjusted squared partial Mahalanobis
    distances
getDim signature(object = "CovRobMiss"): return the dimension of observed entries for each
    case
getLocation signature(object = "CovRobMiss"): return the estimated location vector
```

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getScatter signature(object = "CovRobMiss", cutoff = "numeric"): return the estimated scatter matrix

getMissing signature(object = "CovRobMiss"): return the case number(s) with completely missing data, if any

getOutliers signature(object = "CovRobMiss", cutoff = "numeric"): return the case number(s) adjusted squared distances above (1 - cutoff)th quantile of chi-square p-degrees of freedom.

In addition to above, the following methods are defined with the class "CovRobMissSc":

getScale signature(object = "CovRobMissSc"): return the MVE scale of the best candidate

## Author(s)

Andy Leung <andy.leung@stat.ubc.ca>

#### See Also

emve, CovRobMissSc-class, CovRobMiss-class

geochem

Geochemical Data

## Description

Geochemical data analyzed by Smith et al (1984). The variables in the data measures the contents (in parts per million) for 20 chemical elements (e.g., Copper and Zinc) in 53 samples of rocks in Western Australia.

## Usage

data(geochem)

#### **Format**

The data contains 53 observations on 20 variables corresponding to the 20 chemical elements.

#### References

Smith, R.E., Campbell, N.A., Licheld, A. (1984). Multivariate statistical techniques applied to pisolitic laterite geochemistry at Golden Grove, Western Australia. *Journal of Geochemical Exploration*, **22**, 193–216.

Agostinelli, C., Leung, A., Yohai, V.J., and Zamar, R.H. (2014) Robust estimation of multivariate location and scatter in the presence of cellwise and casewise contamination. arXiv:1406.6031[math.ST]

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## **Examples**

```
## Not run:
library(ICSNP)
library(rrcov)
data(geochem)
n <- nrow(geochem)</pre>
p <- ncol(geochem)</pre>
# MLE
res.ML <- list(mu=colMeans(geochem), S=cov(geochem))</pre>
# Tyler's M
geochem.med <- apply(geochem,2,median,na.rm=TRUE)</pre>
res.Tyler <- tyler.shape(geochem, location=geochem.med)</pre>
res.Tyler <- res.Tyler*(median(mahalanobis( geochem, geochem.med, res.Tyler))/qchisq(0.5, df=p))
res.Tyler <- list(mu=geochem.med, S=res.Tyler)</pre>
# Rocke's Covariace
res.Rock <- CovSest(geochem, method="rocke")</pre>
res.Rock <- list(mu=res.Rock@center, S=res.Rock@cov)</pre>
# Fast-MCD
res.FMCD <- CovMcd( geochem)</pre>
res.FMCD <- list(mu=res.FMCD@center, S=res.FMCD@cov)</pre>
# MVE
res.MVE <- CovMve( geochem)</pre>
res.MVE <- list(mu=res.MVE@center, S=res.MVE@cov)</pre>
# S-estimator with bisquare rho function
res.S <- CovSest(geochem, method="bisquare")</pre>
res.S <- list(mu=res.S@center, S=res.S@cov)</pre>
# Fast-S
res.FS <- CovSest(geochem)</pre>
res.FS <- list(mu=res.FS@center, S=res.FS@cov)</pre>
# 2SGS
res.2SGS <- TSGS( geochem, seed=999 )
res.2SGS <- list(mu=res.2SGS@mu, S=res.2SGS@S)</pre>
# Combine all the results
geochem.res <- list(ML=res.ML, Tyler=res.Tyler, Rocke=res.Rock, MCD=res.FMCD,</pre>
 MVE=res.MVE, FS=res.FS, MVES=res.S, TSGS=res.2SGS)
## Compare LRT distances between different estimators
res.tab <- data.frame( LRT.to.2SGS=c(slrt( res.ML$S, res.2SGS$S),
   slrt( res.Tyler$S, res.2SGS$S),
   slrt( res.Rock$S, res.2SGS$S),
   slrt( res.FMCD$S, res.2SGS$S),
   slrt( res.MVE$S, res.2SGS$S),
```

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```
slrt( res.FS$S, res.2SGS$S),
   slrt( res.S$S, res.2SGS$S),
   slrt( res.2SGS$S, res.2SGS$S) ))
row.names(res.tab) <- c("ML","Tyler","Rocke","MCD","MVE","FS","MVES","TSGS")</pre>
# Calculate proportion of outliers cellwise
pairwise.mahalanobis <- function(x, mu, S){</pre>
 # function that computes pairwise mahalanobis distances
p <- ncol(x)
pairs.md <- c()</pre>
 for(i in 1:(p-1)) for(j in (i+1):p)
 pairs.md <- c(pairs.md, mahalanobis( x[,c(i,j)], mu[c(i,j)], S[c(i,j),c(i,j)]))
pairs.md
res.tab$Full <- res.tab$Pairs <- res.tab$Cell <- NA
for(i in names(geochem.res) ){
## Identify cellwise outliers
uni.dist <- sweep(sweep(geochem, 2, geochem.res[[i]]$mu, "-"), 2,</pre>
 sqrt(diag(geochem.res[[i]]$S)), "/")^2
 uni.dist.stat <- mean(uni.dist > qchisq(.99^{(1/(n*p))}, 1))
 res.tab$Cell[ which( row.names(res.tab) == i)] <- round(uni.dist.stat,3)</pre>
 ## Identify pairwise outliers
 pair.dist <- pairwise.mahalanobis( geochem, geochem.res[[i]]$mu, geochem.res[[i]]$S)</pre>
 pair.dist.stat <- mean(pair.dist > qchisq(0.99^{(1/(n*choose(p,2)))}, 2))
 res.tab$Pairs[ which( row.names(res.tab) == i)] <- round(pair.dist.stat,3)</pre>
 ## Identify any large global MD
 full.dist <- mahalanobis( geochem, geochem.res[[i]]$mu, geochem.res[[i]]$S)</pre>
 full.dist.stat \leftarrow mean(full.dist > qchisq(0.99^(1/n), p))
res.tab$Full[ which( row.names(res.tab) == i)] <- round(full.dist.stat,3)</pre>
}
res.tab
## End(Not run)
```

get-methods

Accessor methods to the essential slots of classes CovRobMiss, TSGS, GSE, emve, and HuberPairwise

# Description

Accessor methods to the slots of objects of classes CovRobMiss, TSGS, GSE, emve, and HuberPairwise

# Usage

```
getLocation(object)
getScatter(object)
getDist(object)
getDistAdj(object)
```

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```
getDim(object)
getMissing(object)
getOutliers(object, cutoff)
getScale(obj)
getFiltDat(object)
```

## **Arguments**

obj, object an object of any of the following classes CovRobMiss-class, GSE-class, emve-class,

and HuberPairwise-class. For function getScale the package defines a method

for objects of class GSE-class objects are allowed.

cutoff optional argument for getOutliers - quantiles of chi-square to be used as a

threshold for outliers detection, defaults to 0.99

#### **Details**

```
getLocation signature(object = "CovRobMiss"): return the estimated location vector
```

getScatter signature(object = "CovRobMiss", cutoff = "numeric"): return the estimated scatter matrix

getDist signature(object = "CovRobMiss"): return the squared partial Mahalanobis distances

getDistAdj signature(object = "CovRobMiss"): return the adjusted squared partial Mahalanobis
distances

getDim signature(object = "CovRobMiss"): return the dimension of observed entries for each
 case

getMissing signature(object = "CovRobMiss"): return the case number with completely missing data, if any

getOutliers signature(object = "CovRobMiss", cutoff = "numeric"): return the case number(s) adjusted squared distances above (1 - cutoff)th quantile of chi-square p-degrees of freedom.

**getScale** signature(object = "CovRobMissSc"): return either the estimated generalized S-scale or MVE-scale. See GSE and emve for details.

getFiltDat signature(object = "TSGS"): return filtered data matrix from the first step of 2SGS.

## **Examples**

```
## Not run:
data(boston)
res <- GSE(boston)

## extract estimated location
getLocation(res)

## extract estimated scatter
getScatter(res)

## extract estimated adjusted distances
getDistAdj(res)</pre>
```

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```
## extract outliers
getOutliers(res)
## End(Not run)
```

GSE

Generalized S-Estimator in the presence of missing data

## **Description**

Computes the Generalized S-Estimate (GSE) – a robust estimate of location and scatter for data with contamination and missingness.

# Usage

```
GSE(x, tol=1e-4, maxiter=150, method=c("bisquare","rocke"),
    init=c("emve","qc","huber","imputed","emve_c"), mu0, S0, ...)
```

## **Arguments**

X	a matrix or data frame. May contain missing values, but cannot contain columns with completely missing entries.
tol	tolerance for the convergence criterion. Default is 1e-4.
maxiter	maximum number of iterations for the GSE algorithm. Default is 150.
method	which loss function to use: 'bisquare', 'rocke'.
init	type of initial estimator. Currently this can either be "emve" (EMVE with uniform sampling, see Danilov et al., 2012), "qc" (QC, see Danilov et al., 2012), "huber" (Huber Pairwise, see Danilov et al., 2012), "imputed" (Imputed Sestimator, see the rejoinder in Agostinelli et al., 2015), or "emve_c" (EMVE_C with cluster sampling, see Leung and Zamar, 2016). Default is "emve". If mu0 and S0 are provided, this argument is ignored.
mu0	optional vector of initial location estimate
S0	optional matrix of initial scatter estimate
	optional arguments for computing the initial estimates (see emve, HuberPairwise).

## Details

This function computes GSE (Danilov et al., 2012) and GRE (Leung and Zamar, 2016). The estimator requires a robust positive definite initial estimator. This initial estimator is required to "re-scale" the partial square mahalanobis distance for the different missing pattern, in which a single scale parameter is not enough. This function currently allows two main initial estimators: EMVE (the default; see emve and Huberized Pairwise (see HuberPairwise). GSE using Huberized Pairwise with sign psi function is referred to as QGSE in Danilov et al. (2012). Numerical results have shown that GSE with EMVE as initial has better performance (in both efficiency and robustness), but computing time can be longer.

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#### Value

An S4 object of class GSE-class which is a subclass of the virtual class CovRobMissSc-class. The output S4 object contains the following slots:

mu Estimated location. Can be accessed via getLocation. S Estimated scatter matrix. Can be accessed via getScatter. Generalized S-scale (GS-scale). Can be accessed via getScale. sc Squared partial Mahalanobis distances. Can be accessed via getDist. pmd pmd.adj Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj. Dimension of the observed entries for each case. Can be accessed via getDim. pu Estimated initial location. mu0 Estimated initial scatter matrix. S0 Input data matrix with missing values imputed using best linear predictor. Not meant to be accessed. ximp weights Weights used in the estimation of the location. Not meant to be accessed. First derivative of the weights used in the estimation of the location. Not meant to be accessed. weightsp Number of iterations till convergence. Not meant to be accessed. iter eps relative change of the GS-scale at convergence. Not meant to be accessed. call Object of class "language". Not meant to be accessed. Input data matrix. Not meant to be accessed. х Column dimension of input data matrix. Not meant to be accessed.

#### Author(s)

estimator

Andy Leung <andy.leung@stat.ubc.ca>, Ruben H. Zamar, Mike Danilov, Victor J. Yohai

## References

Agostinelli, C., Leung, A., Yohai, V.J., and Zamar, R.H. (2015) Robust estimation of multivariate location and scatter in the presence of cellwise and casewise contamination. *TEST*.

Character string of the name of the estimator used. Not meant to be accessed.

Danilov, M., Yohai, V.J., Zamar, R.H. (2012). Robust Esimation of Multivariate Location and Scatter in the Presence of Missing Data. *Journal of the American Statistical Association* **107**, 1178–1186.

Leung, A. and Zamar, R.H. (2016). Multivariate Location and Scatter Matrix Estimation Under Cellwise and Casewise Contamination. Submitted.

## See Also

```
emve, HuberPairwise, GSE-class, generate.casecontam
```

# Examples

```
set.seed(12)  
## generate 10-dimensional data with 10% casewise contamination n <-100  
p <-10  
A <- matrix(0.9, p, p)
```

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```
diag(A) <- 1
x <- generate.casecontam(n, p, cond=100, contam.size=10, contam.prop=0.1, A=A)$x

## introduce 5% missingness
pmiss <- 0.05
nmiss <- matrix(rbinom(n*p,1,pmiss), n,p)
x[ which( nmiss == 1 ) ] <- NA

## Using EMVE as initial
res.emve <- GSE(x)
slrt( getScatter(res.emve), A) ## LRT distances to the true covariance

## Using QC as initial
res.qc <- GSE(x, init="qc")
slrt( getScatter(res.qc), A) ## in general performs worse than if EMVE used as initials</pre>
```

GSE-class

Generalized S-Estimator in the presence of missing data

## **Description**

Class of Generalized S-Estimator. It has the superclass of CovRobMissSc.

#### **Objects from the Class**

Objects can be created by calls of the form new("GSE", ...), but the best way of creating GSE objects is a call to the function GSE which serves as a constructor.

#### Slots

- mu Estimated location. Can be accessed via getLocation.
- S Estimated scatter matrix. Can be accessed via getScatter.
- sc Generalized S-scale (GS-scale). Can be accessed via getScale.
- pmd Square partial Mahalanobis distances. Can be accessed via getDist.
- pmd.adj Adjusted square partial Mahalanobis distances. Can be accessed via getDistAdj.
- pu Dimension of the observed entries for each case. Can be accessed via getDim.
- mu0 Estimated initial location.
- S0 Estimated initial scatter matrix.
- ximp Input data matrix with missing values imputed using best linear predictor. Not meant to be accessed.
- weights Weights used in the estimation of the location. Not meant to be accessed.
- weightsp First derivative of the weights used in the estimation of the location. Not meant to be accessed.
- iter Number of iterations till convergence. Not meant to be accessed.

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```
eps relative change of the GS-scale at convergence. Not meant to be accessed.
```

- call Object of class "language". Not meant to be accessed.
- x Input data matrix. Not meant to be accessed.
- p Column dimension of input data matrix. Not meant to be accessed.
- estimator Character string of the name of the estimator used. Not meant to be accessed.

#### **Extends**

```
Class "CovRobMissSc", directly.
```

#### Methods

The following methods are defined with the superclass "CovRobMiss":

```
show signature(object = "CovRobMiss"): display the object
```

summary signature(object = "CovRobMiss"): calculate summary information

plot signature(object = "CovRobMiss", cutoff = "numeric"): plot the object. See plot

getDist signature(object = "CovRobMiss"): return the squared partial Mahalanobis distances

getDistAdj signature(object = "CovRobMiss"): return the adjusted squared partial Mahalanobis
distances

getLocation signature(object = "CovRobMiss"): return the estimated location vector

getScatter signature(object = "CovRobMiss", cutoff = "numeric"): return the estimated scatter matrix

getMissing signature(object = "CovRobMiss"): return the case number(s) with completely missing data, if any

getOutliers signature(object = "CovRobMiss", cutoff = "numeric"): return the case number(s) adjusted squared distances above (1 - cutoff)th quantile of chi-square p-degrees of freedom.

In addition to above, the following methods are defined with the class "CovRobMissSc":

```
getScale signature(object = "CovRobMissSc"): return the GS scale
```

# Author(s)

```
Andy Leung <andy.leung@stat.ubc.ca>
```

#### See Also

```
GSE, CovRobMissSc-class, CovRobMiss-class
```

20 gy.filt

gy.filt	Gervini-Yohai filter for detecting cellwise outliers	

## **Description**

Flags cellwise outliers detected using Gervini-Yohai filter as described in Agostinelli et al. (2015) and Leung and Zamar (2016).

## Usage

```
gy.filt(x, alpha=c(0.95,0.85), bivarQt=0.99, bivarCellPr=0.1, miter=5)
```

#### **Arguments**

X	a matrix or data frame.
alpha	a vector of the quantiles of the univariate and bivariate reference distributions, respectively. Filtering turns off when alpha is 0. For univariate filtering only, $alpha=c(0.95,0)$ . Default value is $c(0.95,0.85)$ .
bivarQt	quantile of the binomial model for the number of flagged pairs in the bivariate filter. Default is 0.99.
bivarCellPr	probability of the binomial model for the number of flagged pairs in the bivariate filter. Default is 0.1.

miter maximum number of iteration of filtering. Default value is 5.

#### **Details**

This function implements the univariate filter and the univariate-plus-bivariate filter as described in Agostinelli et al. (2015) and Leung and Zamar (2016), respectively.

In the univariate filter, outliers are flagged by comparing the empirical tail distribution of each marginal with a reference (normal) distribution using Gervini-Yohai approach.

In the univiarate-plus-bivariate filter, outliers are first flagged by applying the univariate filter. Then, the bivariate filter is applied to flag any additional outliers. In the bivariate filter, outliers are flagged by comparing the empirical tail distribution of each bivariate marginal with a reference (chi-square with 2 d.f.) distribution using Gervini-Yohai approach. The number of flagged pairs associated with each cell approximately follows a binomial model under Independent Cellwise Contamination Model. A cell is additionally flagged if the number of flagged pairs exceeds a large quantile of the binomial model.

## Value

a matrix or data frame of the filtered data.

# Author(s)

Andy Leung <andy.leung@stat.ubc.ca>, Claudio Agostinelli, Ruben H. Zamar, Victor J. Yohai

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#### References

Agostinelli, C., Leung, A., Yohai, V.J., and Zamar, R.H. (2015) Robust estimation of multivariate location and scatter in the presence of cellwise and casewise contamination. TEST.

Leung, A. and Zamar, R.H. (2016). Multivariate Location and Scatter Matrix Estimation Under Cellwise and Casewise Contamination. Submitted.

#### See Also

TSGS, generate.cellcontam

## **Examples**

```
set.seed(12345)

# Generate 5% cell-wise contaminated normal data
x <- generate.cellcontam(n=100, p=10, cond=100, contam.size=5, contam.prop=0.05)$x

## Using univariate filter only
xna <- gy.filt(x, alpha=c(0.95,0))
mean(is.na(xna))

## Using univariate-and-bivariate filter
xna <- gy.filt(x, alpha=c(0.95,0.95))
mean(is.na(xna))</pre>
```

horse

Horse-colic data

## **Description**

This is a modified version of the original data set (taken from UCI repository, see reference), where only quantitative variables are considered. This data set is about horse diseases where the task is to determine if the lesion of the horse was surgical or not. It contains rows with completely missing values except for ID and must be removed by the users. They are kept mainly for pedagogical purposes.

#### Usage

data(horse)

#### **Format**

A data frame with 368 observations on the following 7 variables are quantitative and 1 categorical. The first variable is a numeric id.

Hospital\_Number
Rectal\_temperature
Pulse

numeric id, i.e. the case number assigned to the horse (may not be unique if the horse is to rectal temperature in degree celcius

the heart rate in beats per minute; normal rate is 30-40 for adults

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Respiratory\_rate respiratory rate; normal rate is 8 to 10

Nasogastric\_reflux\_PH scale is from 0 to 14 with 7 being neutral; normal values are in the 3 to 4 range Packed\_cell\_volume the number of red cells by volume in the blood; normal range is 30 to 50

Total\_protein normal values lie in the 6-7.5 (gms/dL) range

Abdomcentesis\_total\_protein Values are in gms/dL

surgical\_leison was the problem (lesion) surgical?; 1 = yes, 2 = no

#### **Source**

The original data have been taken from the Journal of Statistics Education Databases at

• http://archive.ics.uci.edu/ml/datasets/Horse+Colic,

#### References

Frank, A. & Asuncion, A. (2010). UCI Machine Learning Repository [http://archive.ics.uci.edu/ml]. Irvine, CA: University of California, School of Information and Computer Science.

## **Examples**

```
## Not run:
data(horse)
horse.cts <- horse[,-c(1,9)] ## remove the id and categorical variable
res <- GSE(horse.cts)
plot(res, which="dd", xlog10=TRUE, ylog10=TRUE)
getOutliers(res)
## End(Not run)</pre>
```

HuberPairwise

Quadrant Covariance and Huberized Pairwise Scatter

# Description

Computes the Quadrant Covariance (QC) or Huberized Pairwise Scatter as described in Alqallaf et al. (2002).

## Usage

```
HuberPairwise(x, psi=c("huber", "sign"), c0=1.345, computePmd=TRUE)
```

## **Arguments**

Χ	a matrix or data frame. May contain missing values, but cannot contain columns
	with completely missing entries.

loss function to be used in computing pairwise scatter. Default is "huber". If

psi="sign", this yields QC. Other value includes "huber".

psi

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c0 tuning constant for the huber function. c0=0 would yield QC. Default is c0=1.345.

This parameter is unnecessary if psi='sign'.

computePmd logical indicating whether to compute partial Mahalanobis distances (pmd) and

adjusted pmd. Default is TRUE.

#### **Details**

As described in Alqallaf et al. (2002), this estimator requires a robust scale estimate and a location M-estimate, which will be used to transform the data through a loss-function to be outlier-free. Currently, this function takes MADN (normalized MAD) and median as the robust scale and location estimate to save computation time. By default, the loss function psi is a sign function, but users are encouraged to also try Huberized scatter with the loss function as  $\psi_c(x) = \min(\max(-c,x),c), c > 0, c = 1.345$ . The function does not adjust for intrinsic bias as described in Alqallaf et al. (2002). Missing values will be replaced by the corresponding column's median.

## Value

An S4 object of class HuberPairwise-class which is a subclass of the virtual class CovRobMiss-class. The output S4 object contains the following slots:

mu	Estimated location. Can be accessed via getLocation.
S	Estimated scatter matrix. Can be accessed via getScatter.
pmd	Squared partial Mahalanobis distances. Can be accessed via getDist.
pmd.adj	Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj.
pu	Dimension of the observed entries for each case. Can be accessed via getDim.
R	Estimated correlation matrix. Not meant to be accessed.
call	Object of class "language". Not meant to be accessed.
X	Input data matrix. Not meant to be accessed.
р	Column dimension of input data matrix. Not meant to be accessed.
estimator	Character string of the name of the estimator used. Not meant to be accessed.

#### Author(s)

Andy Leung <andy.leung@stat.ubc.ca>

## References

Alqallaf, F.A., Konis, K. P., R. Martin, D., Zamar, R. H. (2002). Scalable Robust Covariance and Correlation Estimates for Data Mining. In Proceedings of the Seventh ACM SIGKDD International Conference on Knowledge Discovery and Data Mining. Edmonton.

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HuberPairwise-class

Quadrant Covariance and Huberized Pairwise Scatter

# **Description**

Class of Quadrant Covariance and Huberized Pairwise Scatter. It has the superclass of CovRobMiss.

# **Objects from the Class**

Objects can be created by calls of the form new("HuberPairwise", ...), but the best way of creating HuberPairwise objects is a call to the function HuberPairwise which serves as a constructor.

## **Slots**

```
mu Estimated location. Can be accessed via getLocation.
```

S Estimated scatter matrix. Can be accessed via getScatter.

pmd Squared partial Mahalanobis distances. Can be accessed via getDist.

pmd.adj Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj.

pu Dimension of the observed entries for each case. Can be accessed via getDim.

R Estimated correlation matrix. Not meant to be accessed.

call Object of class "language". Not meant to be accessed.

x Input data matrix. Not meant to be accessed.

p Column dimension of input data matrix. Not meant to be accessed.

estimator Character string of the name of the estimator used. Not meant to be accessed.

## **Extends**

```
Class "CovRobMiss", directly.
```

# Methods

No methods defined with class "HuberPairwise" in the signature.

## Author(s)

```
Andy Leung <andy.leung@stat.ubc.ca>
```

#### See Also

HuberPairwise, CovRobMiss-class

ImpS 25

|--|

# **Description**

Computes the simple three-step estimator as described in the rejoinder of Agostinelli et al. (2015).

# Usage

```
ImpS(x, alpha=0.95, method=c("bisquare","rocke"), init=c("emve","emve_c"), ...)
```

## Arguments

x	a matrix or data frame.
alpha	quantile of the reference distribution in the univariate filter step (see gy.filt). Default is 0.95.
method	which loss function to use: 'bisquare', 'rocke'.
init	type of initial estimator. Currently this can either be "emve" (EMVE with uniform sampling, see Danilov et al., 2012) or "emve_c" (EMVE_C with cluster sampling, see Leung and Zamar, 2016). Default is "emve".
	optional, additional arguments to be passed to GSE.

## **Details**

This function computes the simple three-step estimator as described in the rejoinder in Agostinelli et al. (2015). The procedure has three steps:

In Step I, the method flags and removes cell-wise outliers using the Gervini-Yohai univariate only filter (see gy.filt).

In Step II, the method imputes the filtered cells using coordinate-wise medians.

In Step III, the method applies MVE-S to the filtered and imputed data from Step II (see GSE).

#### Value

The following gives the major slots in the output S4 object:

- mu Estimated location. Can be accessed via getLocation.
- S Estimated scatter matrix. Can be accessed via getScatter.
- xf Filtered data matrix from the first step of 2SGS. Can be accessed via getFiltDat.

## Author(s)

Andy Leung <andy.leung@stat.ubc.ca>, Claudio Agostinelli, Ruben H. Zamar, Victor J. Yohai

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## References

Agostinelli, C., Leung, A., Yohai, V.J., and Zamar, R.H. (2015) Robust estimation of multivariate location and scatter in the presence of cellwise and casewise contamination. TEST.

#### See Also

```
GSE, gy.filt
```

partial.mahalanobis

Partial Square Mahalanobis Distance

# **Description**

Computes the partial square Mahalanobis distance for all observations in x. Let  $\mathbf{x} = (x_{i1}, ..., x_{ip})'$  be a p-dimensional random vector and  $\mathbf{u} = (u_{i1}, ..., u_{ip})'$  be a p-dimensional vectors of zeros and ones indicating which entry is missing: 0 as missing and 1 as observed. Then partial mahalanobis distance is given by:

$$d(\mathbf{x}, \mathbf{u}, \mathbf{m}, \boldsymbol{\Sigma}) = (\mathbf{x}^{(\mathbf{u})} - \mathbf{m}^{(\mathbf{u})})'(\boldsymbol{\Sigma}^{(\mathbf{u})})^{-1}(\mathbf{x}^{(\mathbf{u})} - \mathbf{m}^{(\mathbf{u})}).$$

With no missing data, this function is equivalent to mahalanobis distance.

## Usage

```
partial.mahalanobis(x, mu, Sigma)
```

## **Arguments**

x a matrix or data frame. May contain missing values, but cannot contain columns

with completely missing entries.

mu location estimate

Sigma scatter estimate. Must be positive definite

## Value

An S4 object of class CovRobMiss-class. The output S4 object contains the following slots:

mu	Estimated location. Can be accessed via getLocation.
S	Estimated scatter matrix. Can be accessed via getScatter.
pmd	Squared partial Mahalanobis distances. Can be accessed via getDist.
pmd.adj	Adjusted squared partial Mahalanobis distances. Can be accessed via getDistAdj.
pu	Dimension of the observed entries for each case. Can be accessed via getDim.
call	Object of class "language". Not meant to be accessed.
X	Input data matrix. Not meant to be accessed.
р	Column dimension of input data matrix. Not meant to be accessed.
estimator	Character string of the name of the estimator used. Not meant to be accessed.

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## Author(s)

```
Andy Leung <andy.leung@stat.ubc.ca>
```

# **Examples**

```
## Not run:
## suppose we would like to compute pmd for an MLE
x <- matrix(rnorm(1000),100,10)
U <- matrix(rbinom(1000,1,0.1),100,10)
x <- x * ifelse(U==1,NA,1)
## compute MLE (i.e. EM in this case)
res <- CovEM(x)
## compute pmd
res.pmd <- partial.mahalanobis(x, mu=getLocation(res), S=getScatter(res))
summary(res.pmd)
plot(res.pmd, which="index")
## End(Not run)</pre>
```

plot-methods

Plot methods for objects of class 'CovRobMiss'

# Description

Plot methods for objects of class 'CovRobMiss'. The following plots are available:

- chi-square qqplot for adjusted square partial Mahalanobis distances
- index plot for adjusted square partial Mahalanobis distances
- distance-distance plot comparing the adjusted distances based on classical MLE and robust estimators

Cases with completely missing data will be dropped out. Outliers are identified using some prespecific cutoff value, for instance 99% quantile of chi-square with p degrees of freedom, where p is the column dimension of the data. Identified outliers can also be retrieved using getOutliers with an optional argument of cutoff, ranged from 0 to 1.

# Usage

```
## S4 method for signature 'CovRobMiss'
plot(x, which = c("all","distance","qqchi2", "dd"),
  which = c("all", "distance", "qqchisq", "dd"),
  ask = (which=="all" && dev.interactive(TRUE)),
  cutoff = 0.99, xlog10 = FALSE, ylog10 = FALSE)
```

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# **Arguments**

Χ	an object of class "CovRobMiss"
which	Which plot to show? Default is which="all".
ask	logical; if 'TRUE', the user is <i>ask</i> ed before each plot, see 'par(ask=.)'. Default is ask = which=="all" && dev.interactive().
cutoff	The quantile cutoff for the distances. Default is 0.99.
xlog10	Base-10 logged x-axis? Default is FALSE.
ylog10	Base-10 logged y-axis? Default is FALSE.

# **Examples**

```
## Not run:
data(boston)
res <- GSE(boston)

## plot all graphs
plot(res)

## plot individuals plots
plot(res, which="qqchisq")
plot(res, which="index")
plot(res, which="dd")

## control the coordinates, e.g. log10 transform the y-axis
plot(res, which="qqchisq", xlog10=TRUE, ylog10=TRUE)
plot(res, which="index", ylog10=TRUE)
plot(res, which="dd", xlog10=TRUE, ylog10=TRUE)
## End(Not run)</pre>
```

simulation-tools Data generator for simulation study on cell- and case-wise contamination

# **Description**

Includes the data generator for the simulation study on cell- and case-wise contamination that appears on Agostinelli et al. (2014).

# Usage

```
generate.randcorr(cond, p, tol=1e-5, maxits=100)
generate.cellcontam(n, p, cond, contam.size, contam.prop, A=NULL)
generate.casecontam(n, p, cond, contam.size, contam.prop, A=NULL)
```

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# Arguments

cond	desired condition number of the random correlation matrix. The correlation matrix will be used to generate multivariate normal samples in generate.cellcontam and generate.cellcontam.
tol	tolerance level for the condition number of the random correlation matrix. Default is 1e-5.
maxits	integer indicating the maximum number of iterations until the condition number of the random correlation matrix is within a tolerance level. Default is 100.
n	integer indicating the number of observations to be generated.
р	integer indicating the number of variables to be generated.
contam.size	size of cell- or case-wise contamination. For cell-wise outliers, random cells in a data matrix are replaced by contam.dist. For case-wise outliers, random cases in a data matrix are replaced by contam.dist times $\boldsymbol{v}$ where $\boldsymbol{v}$
contam.prop	proportion of cell- or case-wise contamination.
A	correlation matrix used for generating data. If A is NULL, a random correlation matrix is generated. Default is NULL.

#### **Details**

Details about how the correlation matrix is randomly generated and how the contaminated data is generated can be found in Agostinelli et al. (2014).

#### Value

generate.randcorr gives the random correlation matrix in dimension p and with condition number cond.

generate.cellcontam and generate.casecontam give the multivariate normal sample that is either cell-wise or case-wise contaminated as described in Agostinelli et al. (2014). The contaminated sample is returned as components of a list with components

- x multivariate normal sample with cell- or case-wise contamination.
- u n by p matrix of 0's and 1's with 1's correspond to an outlier. A row of 1's correspond to a case-wise outlier.
- A random correlation matrix with a specified condition number.

# Author(s)

Andy Leung <andy.leung@stat.ubc.ca>, Claudio Agostinelli, Ruben H. Zamar, Victor J. Yohai

#### References

Agostinelli, C., Leung, A., Yohai, V.J., and Zamar, R.H. (2014) Robust estimation of multivariate location and scatter in the presence of cellwise and casewise contamination. arXiv:1406.6031[math.ST]

## See Also

**TSGS** 

slrt

LRT-based distances between matrices

## **Description**

LRT-distance that we use to evaluate the performance of our covariance estimates.

# Usage

```
slrt(S, trueS)
```

# Arguments

S estimated covariance matrix trueS true covariance matrix.

## **Details**

Note that this is not actually a distance in a sense that slrt(M1,M2) != slrt(M2,M1)

## Value

scalar LRT-distance

## Author(s)

Mike Danilov

## References

Seber, G.A. (2004) Multivariate observations, Wiley

Danilov, M. (2010). Robust Estimation of Multivariate Scatter under Non-Affine Equivariant Scenarios. Ph.D. thesis, Department of Statistics, University of British Columbia.

 ${\it Summary CovGSE"-class} \quad {\it Class~"Summary CovGSE"-displaying~summary~of~"CovRobMiss"~objects}$ 

# **Description**

Displays summary information for CovRobMiss-class objects

# **Objects from the Class**

Objects can be created by calls of the form new("SummaryCovGSE", ...).

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# **Slots**

```
obj: CovRobMiss-class object
evals: Eigenvalues and eigenvectors of the covariance or correlation matrix
```

## Methods

```
show signature(object = "SummaryCovGSE"): display the object
```

# Author(s)

Andy Leung <andy.leung@stat.ubc.ca>

**TSGS** 

Two-Step Generalized S-Estimator for cell- and case-wise outliers

# **Description**

Computes the Two-Step Generalized S-Estimate (2SGS) – a robust estimate of location and scatter for data with cell-wise and case-wise contamination.

# Usage

```
TSGS(x, filter=c("UBF-DDC","UBF","DDC","UF"),
   partial.impute=FALSE, tol=1e-4, maxiter=150, method=c("bisquare","rocke"),
   init=c("emve","qc","huber","imputed","emve_c"), mu0, S0)
```

# **Arguments**

x	a matrix or data frame.
filter	the filter to be used in the first step (see Leung et al. $(2016)$ ). Default is 'UBF-DDC'. For all filters, the default parameters are used.
partial.impute	whether partial imputation is used prior to estimation (see details). The default is $FALSE$ .
tol	tolerance for the convergence criterion. Default is 1e-4.
maxiter	maximum number of iterations for the GSE algorithm. Default is 150.
method	which loss function to use: 'bisquare', 'rocke'.
init	type of initial estimator. Currently this can either be "emve" (EMVE with uniform sampling, see Danilov et al., 2012), "qc" (QC, see Danilov et al., 2012), "huber" (Huber Pairwise, see Danilov et al., 2012), "imputed" (Imputed Sestimator, see the rejoinder in Agostinelli et al., 2015), or "emve_c" (EMVE_C with cluster sampling, see Leung and Zamar, 2016). Default is "emve". If mu0 and S0 are provided, this argument is ignored.
mu0	optional vector of initial location estimate
SØ	optional matrix of initial scatter estimate

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#### **Details**

This function computes 2SGS as described in Agostinelli et al. (2015) and Leung and Zamar (2016). The procedure has two major steps:

In Step I, the method filters (i.e., flags and removes) cell-wise outliers using Gervini-Yohai univariate filter (Agostinelli et al., 2015) or univariate-bivariate filter (Leung et al., 2016) or univariate-bivariate-plus-DDC filter (Leung et al., 2016; Rousseeuw and Van den Bossche, 2016). The filtering step can be called on its own by using the function gy.filt or DDC.

In Step II, the method applies GSE or GRE (GSE with a Rocke-type loss function), which has been specifically designed to deal with incomplete multivariate data with case-wise outliers, to the filted data coming from Step I. The second step can be called on its own by using the function GSE.

The 2SGS method is intended for continuous variables, and requires that the number of observations n be relatively larger than 5 times the number of variables p for desirable performance (see the rejoinder in Agostinelli et al., 2015). In our numerical studies, for n too small relative to p, 2SGS may experience a lack of convergence, especially for filtered data sets with a proportion of complete observations less than 1/2 + (p+1)/n. To overcome this problem, partial imputation prior to estimation is proposed (see the rejoinder in Agostinelli et al., 2015). The procedure is rather ad hoc, but initial numerical experiements show that partial imputation may work. Further research on this topic is still needed. By default, partial imputation is not used, unless specified.

In general, we warn users to use 2SGS with caution for data set with n relatively smaller than 5 times p.

The application to the chemical data set analyzed in Agostinelli et al. (2015) can be found in geochem.

The tools that were used to generate contaminated data in the simulation study in Agostinelli et al. (2015) can be found in generate.cellcontam and generate.casecontam.

#### Value

The following gives the major slots in the output S4 object:

- mu Estimated location. Can be accessed via getLocation.
- S Estimated scatter matrix. Can be accessed via getScatter.
- xf Filtered data matrix from the first step of 2SGS. Can be accessed via getFiltDat.

## Author(s)

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#### References

Agostinelli, C., Leung, A., Yohai, V.J., and Zamar, R.H. (2015) Robust estimation of multivariate location and scatter in the presence of cellwise and casewise contamination. TEST.

Leung, A., Yohai, V.J., Zamar, R.H. (2016). Multivariate Location and Scatter Matrix Estimation Under Cellwise and Casewise Contamination. arXiv:1609.00402.

Rousseeuw P.J., Van den Bossche W. (2016). Detecting deviating data cells. arXiv:1601.07251

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## See Also

```
GSE, gy.filt, DDC, generate.cellcontam, generate.casecontam
```

# **Examples**

```
set.seed(12345)
# Generate 5% cell-wise contaminated normal data
# using a random correlation matrix with condition number 100
x <- generate.cellcontam(n=100, p=10, cond=100, contam.size=5, contam.prop=0.05)
## Using MLE
slrt(cov(x$x), x$A)
## Using Fast-S
slrt( rrcov:::CovSest(x$x)@cov, x$A)
## Using 2SGS
slrt( TSGS(x$x)@S, x$A)
# Generate 5% case-wise contaminated normal data
# using a random correlation matrix with condition number 100
x <- generate.casecontam(n=100, p=10, cond=100, contam.size=15, contam.prop=0.05)
## Using MLE
slrt(cov(x$x), x$A)
## Using Fast-S
slrt( rrcov:::CovSest(x$x)@cov, x$A)
## Using 2SGS
slrt( TSGS(x$x)@S, x$A)
```

TSGS-class

Two-Step Generalized S-Estimator for cell- and case-wise outliers

## **Description**

Class of Two-Step Generalized S-Estimator. It has the superclass of GSE.

# **Objects from the Class**

Objects can be created by calls of the form new("TSGS", ...), but the best way of creating TSGS objects is a call to the function TSGS which serves as a constructor.

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## **Slots**

- mu Estimated location. Can be accessed via getLocation.
- S Estimated scatter matrix. Can be accessed via getScatter.
- xf Filtered data matrix from the first step of 2SGS. Can be accessed via getFiltDat.

#### **Extends**

```
Class "GSE", directly.
```

#### Methods

In addition to the methods defined in the superclass "GSE", the following methods are also defined: **getFiltDat** signature(object = "TSGS"): return the filtered data matrix.

## Author(s)

```
Andy Leung <andy.leung@stat.ubc.ca>
```

#### See Also

```
TSGS, GSE, GSE-class
```

wages

Wages and Hours

# Description

The data are from a national sample of 6000 households with a male head earning less than USD 15,000 annually in 1966. The data were clasified into 39 demographic groups for analysis. The study was undertaken in the context of proposals for a guaranteed annual wage (negative income tax). At issue was the response of labor supply (average hours) to increasing hourly wages. The study was undertaken to estimate this response from available data.

# Usage

```
data(wages)
```

## **Format**

A data frame with 39 observations on the following 10 variables:

HRS	Average hours worked during the year
RATE	Average hourly wage (USD)
ERSP	Average yearly earnings of spouse (USD)
ERNO	Average yearly earnings of other family members (USD)
NEIN	Average yearly non-earned income

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ASSET	Average family asset holdings (Bank account, etc.) (USD)
AGE	Average age of respondent
DEP	Average number of dependents
RACE	Percent of white respondents
SCH00L	Average highest grade of school completed

# Source

DASL library (http://lib.stat.cmu.edu/DASL/Datafiles/wagesdat.html), the dataset is not anymore available at this source.

# References

 $\ensuremath{\text{D.H.}}$  Greenberg and M. Kosters, (1970). Income Guarantees and the Working Poor, The Rand Corporation.

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