

# Package ‘rebmix’

July 10, 2024

**Version** 2.16.0

**Title** Finite Mixture Modeling, Clustering & Classification

**Description** Random univariate and multivariate finite mixture model generation, estimation, clustering, latent class analysis and classification. Variables can be continuous, discrete, independent or dependent and may follow normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or circular von Mises parametric families.

**Depends** R (>= 3.1.0)

**Imports** methods, stats, utils, graphics, grDevices

**License** GPL (>= 2)

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**NeedsCompilation** yes

**Repository** CRAN

**Date/Publication** 2024-07-10 14:00:02 UTC

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**adult****Adult Dataset**

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## Description

The adult dataset containing 48842 instances with 16 continuous, binary and discrete variables was extracted from the census bureau database. Extraction was done by Barry Becker from the 1994 census bureau database.

## Usage

```
data(adult)
```

## Format

adult is a data frame with 48842 cases (rows) and 16 variables (columns) named:

1. Type binary train or test.
2. Age continuous.
3. Workclass one of the 8 discrete values private, self-emp-not-inc, self-emp-inc, federal-gov, local-gov, state-gov, without-pay or never-worked.
4. Fnlwgt stands for continuous final weight.
5. Education one of the 16 discrete values bachelors, some-college, 11th, hs-grad, prof-school, assoc-acdm, assoc-voc, 9th, 7th-8th, 12th, masters, 1st-4th, 10th, doctorate, 5th-6th or preschool.
6. Education.Num continuous.
7. Marital.Status one of the 7 discrete values married-civ-spouse, divorced, never-married, separated, widowed, married-spouse-absent or married-af-spouse.
8. Occupation one of the 14 discrete values tech-support, craft-repair, other-service, sales, exec-managerial, prof-specialty, handlers-cleaners, machine-op-inspct, adm-clerical, farming-fishing, transport-moving, priv-house-serv, protective-serv or armed-forces.
9. Relationship one of the 6 discrete values wife, own-child, husband, not-in-family, other-relative or unmarried.
10. Race one of the 5 discrete values white, asian-pac-islander, amer-indian-eskimo, other or black.
11. Sex binary female or male.
12. Capital.Gain continuous.
13. Capital.Loss continuous.
14. Hours.Per.Week continuous.
15. Native.Country one of the 41 discrete values united-states, cambodia, england, puerto-rico, canada, germany, outlying-us(guam-usvi-etc), india, japan, greece, south, china, cuba, iran, honduras, philippines, italy, poland, jamaica, vietnam, mexico, portugal, ireland, france, dominican-republic, laos, ecuador, taiwan, haiti, columbia, hungary, guatemala, nicaragua, scotland, thailand, yugoslavia, el-salvador, trinidad&tobago, peru, hong or holand-netherlands.
16. Income binary <=50k or >50k.

## Source

A. Asuncion and D. J. Newman. Uci machine learning repository, 2007. <http://archive.ics.uci.edu/ml/>.

## References

A. Asuncion and D. J. Newman. Uci machine learning repository, 2007. <http://archive.ics.uci.edu/ml/>.

## Examples

```
data(adult)

# Find complete cases.

adult <- adult[complete.cases(adult),]

# Show level attributes for binary and discrete variables.

levels(adult[["Type"]])
levels(adult[["Workclass"]])
levels(adult[["Education"]])
levels(adult[["Marital.Status"]])
levels(adult[["Occupation"]])
levels(adult[["Relationship"]])
levels(adult[["Race"]])
levels(adult[["Sex"]])
levels(adult[["Native.Country"]])
levels(adult[["Income"]])
```

## Description

Returns the Akaike information criterion at pos.

## Usage

```
## S4 method for signature 'REBMIX'
AIC(x = NULL, pos = 1, ...)
## S4 method for signature 'REBMIX'
AIC3(x = NULL, pos = 1, ...)
## S4 method for signature 'REBMIX'
AIC4(x = NULL, pos = 1, ...)
## S4 method for signature 'REBMIX'
AICc(x = NULL, pos = 1, ...)
## S4 method for signature 'REBMIX'
CAIC(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- x see Methods section below.
- pos a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
- ... currently not used.

## Methods

- `signature(x = "REBMIX")` an object of class REBMIX.
- `signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- H. Akaike. A new look at the statistical model identification. *IEEE Transactions on Automatic Control*, 19(51):716-723, 1974.
- A. F. M. Smith and D. J. Spiegelhalter. Bayes factors and choice criteria for linear models. *Journal of the Royal Statistical Society. Series B*, 42(2):213-220, 1980. <https://www.jstor.org/stable/2984964>.
- H. Bozdogan. Model selection and akaike's information criterion (aic): The general theory and its analytical extensions. *Psychometrika*, 52(3):345-370, 1987. doi:[10.1007/BF02294361](https://doi.org/10.1007/BF02294361).
- C. M. Hurvich and C.-L. Tsai. Regression and time series model selection in small samples. *Biometrika*, 76(2):297-307, 1989. <https://www.jstor.org/stable/2336663>.

AWE-methods

*Approximate Weight of Evidence Criterion*

## Description

Returns the approximate weight of evidence criterion at pos.

## Usage

```
## S4 method for signature 'REBMIX'
AWE(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- x see Methods section below.
- pos a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
- ... currently not used.

## Methods

`signature(x = "REBMIX")` an object of class REBMIX.  
`signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

J. D. Banfield and A. E. Raftery. Model-based gaussian and non-gaussian clustering. *Biometrics*, 49(3):803-821, 1993. [doi:10.2307/2532201](https://doi.org/10.2307/2532201).

`bearings`

*Bearings Faults Detection Data*

## Description

These data are the results of the extraction process from the vibrational data of healthy and faulty bearings. Different faults are considered: faultless (1), defect on outer race (2), defect on inner race (3) and defect on ball (4). The extracted features are: root mean square (RMS), square root of the amplitude (SRA), kurtosis value (KV), skewness value (SV), peak to peak value (PPV), crest factor (CF), impulse factor (IF), margin factor (MF), shape factor (SF), kurtosis factor (KF), frequency centre (FC), root mean square frequency (RMSF) and root variance frequency (RVF).

## Usage

`data(bearings)`

## Format

`bearings` is a data frame with 1906 cases (rows) and 14 variables (columns) named:

1. RMS continuous.
2. SRA continuous.
3. KV continuous.
4. SV continuous.
5. PPV continuous.
6. CF continuous.
7. IF continuous.
8. MF continuous.
9. SF continuous.
10. KF continuous.
11. FC continuous.
12. RMSF continuous.
13. RVF continuous.
14. Class discrete 1, 2, 3 or 4.

## Source

Case Western Reserve University Bearing Data Center Website <https://engineering.case.edu/bearingdatacenter/welcome>.

## References

B. Panic, J. Klemenc and M. Nagode. Gaussian mixture model based classification revisited: Application to the bearing fault classification. Journal of Mechanical Engineering, 66(4):215-226, 2020.  
[doi:10.5545/svjmme.2020.6563](https://doi.org/10.5545/svjmme.2020.6563).

## Examples

```
## Not run:
data(bearings)

# Split dataset into train (75

set.seed(3)

Bearings <- split(p = 0.75, Dataset = bearings, class = 14)

# Estimate number of components, component weights and component
# parameters for train subsets.

bearingsest <- REBMIX(model = "REBMVNORM",
Dataset = a.train(Bearings),
Preprocessing = "histogram",
cmax = 15,
Criterion = "BIC")

# Classification.

bearingscla <- RCLSMIX(model = "RCLSMVNORM",
x = list(bearingsest),
Dataset = a.test(Bearings),
Zt = a.Zt(Bearings))

bearingscla

summary(bearingscla)

## End(Not run)
```

## Description

Returns as default the optimized RCLSMIX algorithm output for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities. If `model` equals "RCLSMVNORM" optimized output for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices is returned.

## Usage

```
## S4 method for signature 'RCLSMIX'
BFSMIX(model = "RCLSMIX", x = list(), Dataset = data.frame(),
        Zt = factor(), ...)
## ... and for other signatures
```

## Arguments

<code>model</code>	see Methods section below.
<code>x</code>	a list of objects of class REBMIX of length $o$ obtained by running <code>REBMIX</code> on $g = 1, \dots, s$ train datasets $Y_{\text{train}_g}$ all of length $n_{\text{train}_g}$ . For the train datasets the corresponding class membership $\Omega_g$ is known. This yields $n_{\text{train}} = \sum_{g=1}^s n_{\text{train}_g}$ , while $Y_{\text{train}_q} \cap Y_{\text{train}_g} = \emptyset$ for all $q \neq g$ . Each object in the list corresponds to one chunk, e.g., $(y_{1j}, y_{3j})^\top$ . The default value is <code>list()</code> .
<code>Dataset</code>	a data frame containing test dataset $Y_{\text{test}}$ of length $n_{\text{test}}$ . For the test dataset the corresponding class membership $\Omega_g$ is not known. The default value is <code>data.frame()</code> .
<code>Zt</code>	a factor of true class membership $\Omega_g$ for the test dataset. The default value is <code>factor()</code> .
<code>...</code>	currently not used.

## Value

Returns an optimized object of class RCLSMIX or RCLSMVNORM.

## Methods

```
signature(model = "RCLSMIX") a character giving the default class name "RCLSMIX" for mix-
tures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial,
Poisson, Dirac, uniform or von Mises component densities.

signature(model = "RCLSMVNORM") a character giving the class name "RCLSMVNORM" for mix-
tures of multivariate normal component densities with unrestricted variance-covariance matri-
ces.
```

## Author(s)

Marko Nagode

## References

R. Kohavi and G. H. John. Wrappers for feature subset selection, Artificial Intelligence, 97(1-2):273-324, 1997. doi:[10.1016/S00043702\(97\)00043X](https://doi.org/10.1016/S00043702(97)00043X).

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BIC-methods

*Bayesian Information Criterion*

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## Description

Returns the Bayesian information criterion at pos.

## Usage

```
## S4 method for signature 'REBMIX'  
BIC(x = NULL, pos = 1, ...)  
## ... and for other signatures
```

## Arguments

- |     |  |
|-----|--|
| x   | see Methods section below.   |
| pos | a desired row number in x@summary for which the information criterion is calculated. The default value is 1. |
| ... | currently not used.  |

## Methods

- signature(x = "REBMIX") an object of class REBMIX.
- signature(x = "REBMVNORM") an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

G. Schwarz. Estimating the dimension of the model. The Annals of Statistics, 6(2):461-464, 1978.

## Description

Returns the list of data frames containing bin means  $\bar{y}_1, \dots, \bar{y}_v$  and frequencies  $k_1, \dots, k_v$  for the histogram preprocessing.

## Usage

```
## S4 method for signature 'list'
bins(Dataset = list(), K = matrix(),
      ymin = numeric(), ymax = numeric(), ...)
## ... and for other signatures
```

## Arguments

Dataset	a list of length $n_D$ of data frames of size $n \times d$ containing $d$ -dimensional datasets. Each of the $d$ columns represents one random variable. Numbers of observations $n$ equal the number of rows in the datasets.
K	a matrix of size $n_D \times d$ containing numbers of bins $v_1, \dots, v_d$ for the histogram. If, e.g., $K = \text{matrix}(c(10, 15, 18, 5, 7, 9), \text{byrow} = \text{TRUE}, \text{ncol} = 3)$ than $d = 3$ and the list Dataset contains $n_D = 2$ data frames. Hence, different numbers of bins can be assigned to $y_1, \dots, y_d$ . The default value is $\text{matrix}()$ .
ymin	a vector of length $d$ containing minimum observations. The default value is $\text{numeric}()$ .
ymax	a vector of length $d$ containing maximum observations. The default value is $\text{numeric}()$ .
...	currently not used.

## Methods

`signature(x = "list")` a list of data frames.

## Author(s)

Branislav Panic, Marko Nagode

## References

M. Nagode. Finite mixture modeling via REBMIX. Journal of Algorithms and Optimization, 3(2):14-28, 2015. <https://repozitorij.uni-lj.si/Dokument.php?id=127674&lang=eng>.

## Examples

```
# Generate multivariate normal datasets.

n <- c(7, 10)

Theta <- new("RNGMVNORM.Theta", c = 2, d = 2)

a.theta1(Theta, 1) <- c(8, 6)
a.theta1(Theta, 2) <- c(6, 8)
a.theta2(Theta, 1) <- c(8, 2, 2, 4)
a.theta2(Theta, 2) <- c(2, 1, 1, 4)

sim2d <- RNGMIX(model = "RNGMVNORM",
Dataset.name = paste("sim2d_", 1:2, sep = ""),
rseed = -1,
n = n,
Theta = a.Theta(Theta))

# Calculate optimal numbers of bins.

opt.k <- optbins(Dataset = sim2d@Dataset,
Rule = "Knuth equal",
kmin = 1,
kmax = 20)

opt.k

Y <- bins(Dataset = sim2d@Dataset, K = opt.k)

Y

opt.k <- optbins(Dataset = sim2d@Dataset,
Rule = "Knuth unequal",
kmin = 1,
kmax = 20)

opt.k

Y <- bins(Dataset = sim2d@Dataset, K = opt.k)

Y
```

## Description

Returns as default the boot output for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities. If

`x` is of class `RNGMVNORM` the boot output for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices is returned.

## Usage

```
## S4 method for signature 'REBMIX'
boot(x = NULL, rseed = -1, pos = 1, Bootstrap = "parametric",
      B = 100, n = numeric(), replace = TRUE, prob = numeric(), ...)
## ... and for other signatures
## S4 method for signature 'REBMIX.boot'
summary(object, ...)
## ... and for other signatures
```

## Arguments

<code>x</code>	see Methods section below.
<code>rseed</code>	set the random seed to any negative integer value to initialize the sequence. The first bootstrap dataset corresponds to it. For each next bootstrap dataset the random seed is decremented $r_{seed} = r_{seed} - 1$ . The default value is <code>-1</code> .
<code>pos</code>	a desired row number in <code>x@summary</code> to be bootstrapped. The default value is <code>1</code> .
<code>Bootstrap</code>	a character giving the bootstrap type. One of default <code>"parametric"</code> or <code>"nonparametric"</code> .
<code>B</code>	number of bootstrap datasets. The default value is <code>100</code> .
<code>n</code>	number of observations. The default value is <code>numeric()</code> .
<code>replace</code>	logical. The sampling is with replacement if <code>TRUE</code> , see also <code>sample</code> . The default value is <code>TRUE</code> .
<code>prob</code>	a vector of length <code>n</code> containing probability weights, see also <code>sample</code> . The default value is <code>numeric()</code> .
<code>...</code>	maximum number of components <code>cmax</code> , minimum number of components <code>cmin</code> and further arguments to <code>sample</code> ; additional arguments affecting the summary produced.
<code>object</code>	see Methods section below.

## Value

Returns an object of class `REBMIX.boot` or `REBMVNORM.boot`.

## Methods

```
signature(x = "REBMIX") an object of class REBMIX for mixtures of conditionally independent
normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises
component densities.

signature(x = "REBMVNORM") an object of class REBMVNORM for mixtures of multivariate normal
component densities with unrestricted variance-covariance matrices.

signature(object = "REBMIX") an object of class REBMIX.

signature(object = "REBMVNORM") an object of class REBMVNORM.
```

**Author(s)**

Marko Nagode

**References**

G. McLachlan and D. Peel. Finite Mixture Models. John Wiley & Sons, New York, 2000.

**Examples**

```
## Not run:
data(weibull)

# Create object of class EM.Control.

EM <- new("EM.Control", strategy = "single", variant = "EM",
acceleration = "fixed", acceleration.multiplier = 1.0, tolerance = 1.0E-4,
maximum.iterations = 1000)

# Estimate number of components, component weights and component parameters.

weibullest <- REBMIX(Dataset = list(weibull),
Preprocessing = "kernel density estimation",
cmin = 2,
cmax = 4,
Criterion = "BIC",
pdf = "Weibull",
EMcontrol = EM)

# Plot finite mixture.

plot(weibullest, what = c("pdf", "marginal cdf", "IC", "logL", "D"),
nrow = 3, ncol = 2, npts = 1000)

# Bootstrap finite mixture.

weibullboot <- boot(x = weibullest, Bootstrap = "nonparametric", B = 10)

weibullboot

## End(Not run)
```

**Description**

Returns an object of class `Histogram`. The method can be called recursively. This way more than one dataset can be binned into one histogram. The method is time consuming.

## Usage

```
## S4 method for signature 'Histogram'
histogram(x = NULL, Dataset = data.frame(),
           K = numeric(), ymin = numeric(), ymax = numeric(), ...)
## ... and for other signatures
```

## Arguments

<code>x</code>	an object of class <code>Histogram</code> .
<code>Dataset</code>	a data frame of size $n \times d$ containing $d$ -dimensional dataset. Each of the $d$ columns represents one random variable. Number of observations $n$ equals the number of rows in the dataset.
<code>K</code>	an integer or a vector of length $d$ containing numbers of bins $v$ .
<code>ymin</code>	a vector of length $d$ containing minimum observations.
<code>ymax</code>	a vector of length $d$ containing maximum observations.
<code>...</code>	currently not used.

## Methods

`signature(x = "Histogram")` an object of class `Histogram`.

## Author(s)

Marko Nagode

## Examples

```
# Create three datasets.

set.seed(1)

n <- 15

Dataset1 <- as.data.frame(cbind(rnorm(n, 157, 8), rnorm(n, 71, 10)))
Dataset2 <- as.data.frame(cbind(rnorm(n, 244, 14), rnorm(n, 61, 29)))
Dataset3 <- as.data.frame(cbind(rnorm(n, 198, 8), rnorm(n, 252, 13)))

apply(Dataset1, 2, range)
apply(Dataset2, 2, range)
apply(Dataset3, 2, range)

# Bin the first dataset.

hist <- histogram(Dataset = Dataset1, K = c(4, 5), ymin = c(100.0, 0.0), ymax = c(300.0, 300.0))

# Bin the second dataset.

hist <- histogram(x = hist, Dataset = Dataset2)
```

```
# Bin the third dataset.

hist <- histogram(x = hist, Dataset = Dataset3)

hist
```

chunk-methods

*Extracts Chunk from Train and Test Datasets*

## Description

Returns (invisibly) the object containing train and test observations  $x_1, \dots, x_n$  as well as true class membership  $\Omega_g$  for the test dataset. Vectors  $x$  are subvectors of  $y = (y_1, \dots, y_d)^\top$ .

## Usage

```
## S4 method for signature 'RCLS.chunk'
chunk(x = NULL, variables = expression(1:d))
## ... and for other signatures
```

## Arguments

<code>x</code>	see Methods section below.
<code>variables</code>	a vector containing indices of variables in subvectors $x$ . The default value is <code>1:d</code> .

## Value

Returns an object of class `RCLS.chunk`.

## Methods

`signature(x = "RCLS.chunk")` an object of class `RCLS.chunk`.

## Author(s)

Marko Nagode

## Examples

```
data(iris)

# Split dataset into train (75%) and test (25%) subsets.

set.seed(5)

Iris <- split(p = 0.75, Dataset = iris, class = 5)

# Extract chunk from train and test datasets.
```

```
Iris14 <- chunk(x = Iris, variables = c(1,4))

Iris14
```

## Description

Returns the classification likelihood criterion at pos.

## Usage

```
## S4 method for signature 'REBMIX'
CLC(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- |     |  |
|-----|--|
| x   | see Methods section below.   |
| pos | a desired row number in x@summary for which the information criterion is calculated. The default value is 1. |
| ... | currently not used.  |

## Methods

- signature(x = "REBMIX") an object of class REBMIX.
- signature(x = "REBMVNORM") an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- C. Biernacki and G. Govaert. Using the classification likelihood to choose the number of clusters.  
In E. J. Wegman and S. P. Azen, editors, Computing Science and Statistics, 1997.

## Description

Returns the data frame containing observations  $\mathbf{x}_1, \dots, \mathbf{x}_n$  and empirical densities  $f_1, \dots, f_n$  for the kernel density estimation or  $k$ -nearest neighbour or bin means  $\bar{\mathbf{x}}_1, \dots, \bar{\mathbf{x}}_v$  and empirical densities  $f_1, \dots, f_v$  for the histogram preprocessing. Vectors  $\mathbf{x}$  and  $\bar{\mathbf{x}}$  are subvectors of  $\mathbf{y} = (y_1, \dots, y_d)^\top$  and  $\bar{\mathbf{y}} = (\bar{y}_1, \dots, \bar{y}_d)^\top$ .

## Usage

```
## S4 method for signature 'REBMIX'
demix(x = NULL, pos = 1, variables = expression(1:d), ...)
## ... and for other signatures
```

## Arguments

<code>x</code>	see Methods section below.
<code>pos</code>	a desired row number in <code>x@summary</code> for which the empirical densities are calculated. The default value is 1.
<code>variables</code>	a vector containing indices of variables in subvectors $\mathbf{x}$ or $\bar{\mathbf{x}}$ . The default value is <code>1:d</code> .
<code>...</code>	currently not used.

## Methods

`signature(x = "REBMIX")` an object of class REBMIX.  
`signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- M. Nagode and M. Fajdiga. The rebmix algorithm for the univariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(5):876-892, 2011a. doi:[10.1080/03610920903480890](https://doi.org/10.1080/03610920903480890).
- M. Nagode and M. Fajdiga. The rebmix algorithm for the multivariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(11):2022-2034, 2011b. doi:[10.1080/03610921003725788](https://doi.org/10.1080/03610921003725788).
- M. Nagode. Finite mixture modeling via REBMIX. Journal of Algorithms and Optimization, 3(2):14-28, 2015. <https://repozitorij.uni-lj.si/Dokument.php?id=127674&lang=eng>.

## Examples

```
# Generate simulated dataset.

n <- c(15, 15)

Theta <- new("RNGMIX.Theta", c = 2, pdf = rep("normal", 3))

a.theta1(Theta, 1) <- c(10, 20, 30)
a.theta1(Theta, 2) <- c(3, 4, 5)
a.theta2(Theta, 1) <- c(3, 2, 1)
a.theta2(Theta, 2) <- c(15, 10, 5)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1:4, sep = ""),
rseed = -1,
n = n,
Theta = a.Theta(Theta))

# Create object of class EM.Control.

EM <- new("EM.Control", strategy = "best")

# Estimate number of components, component weights and component parameters.

simulatedest <- REBMIX(model = "REBMVNORM",
Dataset = a.Dataset(simulated),
Preprocessing = "h",
cmax = 8,
Criterion = "BIC",
EMcontrol = NULL)

# Preprocess simulated dataset.

f <- demix(simulatedest, pos = 3, variables = c(1, 3))

f

# Plot finite mixture.

opar <- plot(simulatedest, pos = 3, nrow = 3, ncol = 1)

par(usr = opar[[2]]$usr, mfg = c(2, 1))

text(x = f[, 1], y = f[, 2], labels = format(f[, 3], digits = 3), cex = 0.8, pos = 1)
```

## Description

Returns the data frame containing observations  $\mathbf{x}_1, \dots, \mathbf{x}_n$  and predictive marginal densities  $f(\mathbf{x}|c, \mathbf{w}, \Theta)$ . Vectors  $\mathbf{x}$  are subvectors of  $\mathbf{y} = (y_1, \dots, y_d)^\top$ . If  $\mathbf{x} = \mathbf{y}$  the method returns the data frame containing observations  $\mathbf{y}_1, \dots, \mathbf{y}_n$  and the corresponding predictive mixture densities  $f(\mathbf{y}|c, \mathbf{w}, \Theta)$ .

## Usage

```
## S4 method for signature 'REBMIX'
dfmix(x = NULL, Dataset = data.frame(), pos = 1, variables = expression(1:d), ...)
## ... and for other signatures
```

## Arguments

x	see Methods section below.
Dataset	a data frame containing observations $\mathbf{y} = (y_1, \dots, y_d)^\top$ for which the predictive marginal densities are calculated. The default value is <code>data.frame()</code> .
pos	a desired row number in <code>x@summary</code> for which the predictive marginal densities are calculated. The default value is 1.
variables	a vector containing indices of variables in subvectors $\mathbf{x}$ . The default value is <code>1:d</code> .
...	currently not used.

## Methods

`signature(x = "REBMIX")` an object of class REBMIX.

`signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

M. Nagode and M. Fajdiga. The rebmix algorithm for the univariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(5):876-892, 2011a. doi:[10.1080/03610920903480890](https://doi.org/10.1080/03610920903480890).

M. Nagode and M. Fajdiga. The rebmix algorithm for the multivariate finite estimation. Communications in Statistics - Theory and Methods, 40(11):2022-2034, 2011b. doi:[10.1080/03610921003725788](https://doi.org/10.1080/03610921003725788).

M. Nagode. Finite mixture modeling via REBMIX. Journal of Algorithms and Optimization, 3(2):14-28, 2015. <https://repozitorij.uni-lj.si/Dokument.php?id=127674&lang=eng>.

## Examples

```
# Generate simulated dataset.

n <- c(15, 15)

Theta <- new("RNGMIX.Theta", c = 2, pdf = rep("normal", 3))

a.theta1(Theta, 1) <- c(10, 20, 30)
a.theta1(Theta, 2) <- c(3, 4, 5)
a.theta2(Theta, 1) <- c(3, 2, 1)
a.theta2(Theta, 2) <- c(15, 10, 5)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1:4, sep = ""),
rseed = -1,
n = n,
Theta = a.Theta(Theta))

# Number of classes or nearest neighbours to be processed.

K <- c(as.integer(1 + log2(sum(n))), # Minimum v follows Sturges rule.
as.integer(10 * log10(sum(n)))) # Maximum v follows log10 rule.

# Estimate number of components, component weights and component parameters.

simulatedest <- REBMIX(model = "REBMVNORM",
Dataset = a.Dataset(simulated),
Preprocessing = "h",
cmax = 4,
Criterion = "BIC")

# Preprocess simulated dataset.

Dataset <- data.frame(c(-7, 1), NA, c(3, 7))

f <- dfmix(simulatedest, Dataset = Dataset, pos = 3, variables = c(1, 3))

f

# Plot finite mixture.

opar <- plot(simulatedest, pos = 3, nrow = 3, ncol = 1,
contour.drawlabels = TRUE, contour.labcex = 0.6)

par(usr = opar[[2]]$usr, mfg = c(2, 1))

points(x = f[, 1], y = f[, 2])

text(x = f[, 1], y = f[, 2], labels = format(f[, 3], digits = 3), cex = 0.8, pos = 4)
```

## Description

Object of class EM.Control.

## Objects from the Class

Objects can be created by calls of the form new("EM.Control", ...). Accessor methods for the slots are a.strategy(x = NULL), a.variant(x = NULL), a.acceleration(x = NULL), a.tolerance(x = NULL), a.acceleration.multiplier(x = NULL), a.maximum.iterations(x = NULL), a.K(x = NULL) and a.eliminate.zero.components(x = NULL), where x stands for an object of class EM.Control. Setter methods a.strategy(x = NULL), a.variant(x = NULL), a.acceleration(x = NULL), a.tolerance(x = NULL), a.acceleration.multiplier(x = NULL), a.maximum.iterations(x = NULL), a.K(x = NULL) and eliminate.zero.components are provided to write to strategy, variant, acceleration, tolerance, acceleration.multiplier, maximum.iterations and eliminate.zero.components slot respectively.

## Slots

**strategy:** a character containing the EM and REBMIX strategy. One of "none", "exhaustive", "best" and "single". The default value is "none".

**variant:** a character containing the type of the EM algorithm to be used. One of "EM" or "ECM". The default value is "EM".

**acceleration:** a character containing the type of acceleration of the EM iteration increment. One of "fixed", "line" or "golden". The default value is "fixed".

**tolerance:** tolerance value for the EM convergence criteria. The default value is 1.0E-4.

**acceleration.multiplier:** acceleration.multiplier  $a_{EM}$ ,  $1.0 \leq a_{EM} \leq 2.0$ . acceleration.multiplier for the EM step increment. The default value is 1.0.

**maximum.iterations:** a positive integer containing the maximum allowed number of iterations of the EM algorithm. The default value is 1000.

**K:** an integer containing the number of bins for the histogram based EM algorithm. This option can reduce computational time drastically if the datasets contain a large number of observations  $n$  and K is set to the value  $\ll n$ . The default value of 0 means that the EM algorithm runs over all  $n$ .

**eliminate.zero.components:** a logical indicating if the components with  $w_l = 0$  should be eliminated from output. Only used with EMMIX-methods.

## Author(s)

Branislav Panic

## References

B. Panic, J. Klemenc, M. Nagode. Improved initialization of the EM algorithm for mixture model parameter estimation. Mathematics, 8(3):373, 2020. doi:[10.3390/math8030373](https://doi.org/10.3390/math8030373).

A. P. Dempster et al. Maximum likelihood from incomplete data via the EM algorithm. Journal of the Royal Statistical Society. Series B, 39(1):1-38, 1977. <https://www.jstor.org/stable/2984875>.

G. Celeux and G. Govaert. A classification EM algorithm for clustering and two stochastic versions, Computational Statistics & Data Analysis, 14(3):315:332, 1992. doi:[10.1016/01679473\(92\)90042-E](https://doi.org/10.1016/01679473(92)90042-E).

## Examples

```
# Inline creation by new call.

EM <- new("EM.Control", strategy = "exhaustive",
           variant = "EM", acceleration = "fixed",
           tolerance = 1e-4, acceleration.multiplier = 1.0,
           maximum.iterations = 1000, K = 0)

EM

# Creation of EM object with setter method.

EM <- new("EM.Control")

a.strategy(EM) <- "exhaustive"
a.variant(EM) <- "EM"
a.acceleration(EM) <- "fixed"
a.tolerance(EM) <- 1e-4
a.acceleration.multiplier(EM) <- 1.0
a.maximum.iterations(EM) <- 1000
a.K(EM) <- 256

EM
```

## Description

Returns as default the EM algorithm output for mixtures of conditionally independent normal, log-normal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac or von Mises component densities. If model equals "REBMVNORM" output for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices is returned.

## Usage

```
## S4 method for signature 'REBMIX'
EMMIX(model = "REBMIX", Dataset = list(),
      Theta = NULL, EMcontrol = NULL, ...)
## ... and for other signatures
```

## Arguments

model	see Methods section below.
Dataset	a list of length $n_D$ of data frames of size $n \times d$ containing $d$ -dimensional datasets. Each of the $d$ columns represents one random variable. Numbers of observations $n$ equal the number of rows in the datasets.
Theta	an object of class EMMIX.Theta or EMMVNORM.Theta.
EMcontrol	an object of class EM.Control.
...	currently not used.

## Value

Returns an object of class REBMIX or REBMVNORM.

## Methods

```
signature(model = "REBMIX") a character giving the default class name "REBMIX" for mixtures
of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Pois-
son, Dirac or von Mises component densities.

signature(model = "REBMVNORM") a character giving the class name "REBMVNORM" for mixtures
of multivariate normal component densities with unrestricted variance-covariance matrices.
```

## Author(s)

Branislav Panic

## References

B. Panic, J. Klemenc, M. Nagode. Improved initialization of the EM algorithm for mixture model parameter estimation. Mathematics, 8(3):373, 2020. doi:[10.3390/math8030373](https://doi.org/10.3390/math8030373).

## Examples

```
## Not run:
devAskNewPage(ask = TRUE)

# Load faithful dataset.

data(faithful)

# Plot faithfull dataset.

plot(faithful)

# Number of dimensions.

d <- ncol(faithful)

# Obtain 2 component solution with Gaussian mixtures.
```

```

c <- 2

# Create EMMVNORM.Theta object with new call.

Theta <- new("EMMVNORM.Theta", d = d, c = c)

# Set parameters of Theta.
# Weights.

a.w(Theta) <- c(0.5, 0.5)

# Means.

a.theta1.all(Theta) <- c(2.0, 55.0, 4.5, 80.0)

# Covariances.

a.theta2.all(Theta) <- c(1, 0, 0, 1, 1, 0, 0, 1)

# Run EMMIX method.

model <- EMMIX(model = "REBMVNORM", Dataset = list(faithful), Theta = Theta)

# show.

model

# summary.

summary(model)

# plot.

plot(model, nrow = 3, ncol = 2, what = c("pdf", "marginal pdf", "marginal cdf"))

# Create EMMIX.Theta object with new call.

Theta <- new("EMMIX.Theta", c = c, pdf = c("normal", "normal"))

# Set parameters of Theta.
# Weights.

a.w(Theta) <- c(0.5, 0.5)

# Means.

a.theta1.all(Theta) <- c(2.0, 55.0, 4.5, 80.0)

# Covariances.

a.theta2.all(Theta) <- c(1, 1, 1, 1)

# Run EMMIX method.

```

```

model <- EMMIX(Dataset = list(faithful), Theta = Theta)

# show.

model

# summary.

summary(model)

# plot.

plot(model, nrow = 3, ncol = 2, what = c("pdf", "marginal pdf", "marginal cdf"))

## End(Not run)

```

EMMIX.Theta-class      *Class "EMMIX.Theta"*

## Description

Object of class EMMIX.Theta.

## Objects from the Class

Objects can be created by calls of the form `new("EMMIX.Theta", ...)`. Accessor methods for the slots are `a.c(x = NULL)`, `a.d(x = NULL)`, `a.pdf(x = NULL)` and `a.Theta(x = NULL)`, where `x` stands for an object of class EMMIX.Theta. Setter methods `a.theta1(x = NULL, l = numeric())`, `a.theta2(x = NULL, l = numeric())`, `a.theta3(x = NULL, l = numeric())`, `a.theta1.all(x = NULL)`, `a.theta2.all(x = NULL)`, `a.theta3.all(x = NULL)` and `a.w(x = NULL)` are provided to write to Theta slot, where  $l = 1, \dots, c$ .

## Slots

- c:** number of components  $c > 0$ . The default value is 1.
- d:** number of dimensions.
- pdf:** a character vector of length  $d$  containing continuous or discrete parametric family types. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac" or "vonMises".
- Theta:** a list containing  $c$  parametric family types `pdfl`. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac" or circular "vonMises" defined for  $0 \leq y_i \leq 2\pi$ . Component parameters `theta1.l` follow the parametric family types. One of  $\mu_{il}$  for normal, lognormal, Gumbel and von Mises distributions and  $\theta_{il}$  for Weibull, gamma, binomial, Poisson and Dirac distributions. Component parameters `theta2.l` follow `theta1.l`. One of  $\sigma_{il}$  for normal, lognormal and Gumbel distributions,  $\beta_{il}$  for Weibull and gamma distributions,  $p_{il}$  for binomial distribution,  $\kappa_{il}$  for von Mises distribution. Component parameters `theta3.l` follow `theta2.l`. One of  $\xi_{il} \in \{-1, 1\}$  for Gumbel distribution.
- w:** a vector of length  $c$  containing component weights  $w_l$  summing to 1.

**Author(s)**

Branislav Panic

**Examples**

```

Theta <- new("EMMIX.Theta", c = 2, pdf = c("normal", "Gumbel"))

a.w(Theta) <- c(0.4, 0.6)

a.theta1(Theta, l = 1) <- c(2, 10)
a.theta2(Theta, l = 1) <- c(0.5, 2.3)
a.theta3(Theta, l = 1) <- c(NA, 1.0)
a.theta1(Theta, l = 2) <- c(20, 50)
a.theta2(Theta, l = 2) <- c(3, 4.2)
a.theta3(Theta, l = 2) <- c(NA, -1.0)

Theta

Theta <- new("EMMIX.Theta", c = 2, pdf = c("normal", "Gumbel", "Poisson"))

a.w(Theta) <- c(0.4, 0.6)

a.theta1.all(Theta) <- c(2, 10, 30, 20, 50, 60)
a.theta2.all(Theta) <- c(0.5, 2.3, NA, 3, 4.2, NA)
a.theta3.all(Theta) <- c(NA, 1.0, NA, NA, -1.0, NA)

Theta

Theta <- new("EMMVNORM.Theta", c = 2, d = 3)

a.w(Theta) <- c(0.4, 0.6)

a.theta1(Theta, l = 1) <- c(2, 10, -20)
a.theta2(Theta, l = 1) <- c(9, 0, 0, 0, 4, 0, 0, 0, 1)
a.theta1(Theta, l = 2) <- c(-2.4, -15.1, 30)
a.theta2(Theta, l = 2) <- c(4, -3.2, -0.2, -3.2, 4, 0, -0.2, 0, 1)

Theta

Theta <- new("EMMVNORM.Theta", c = 2, d = 3)

a.w(Theta) <- c(0.4, 0.6)

a.theta1.all(Theta) <- c(2, 10, -20, -2.4, -15.1, 30)

a.theta2.all(Theta) <- c(9, 0, 0, 0, 4, 0, 0, 0, 1,
4, -3.2, -0.2, -3.2, 4, 0, -0.2, 0, 1)

Theta

```

---

 fhistogram-methods      *Fast Histogram Calculation*


---

**Description**

Returns an object of class `Histogram`. The method can be called recursively. This way more than one dataset can be binned into one histogram. Set `shrink` to `TRUE` only when the method is called for the last time to optimize the size of the object. The method is memory consuming.

**Usage**

```
## S4 method for signature 'Histogram'
fhistogram(x = NULL, Dataset = data.frame(),
           K = numeric(), ymin = numeric(), ymax = numeric(),
           shrink = FALSE, ...)
## ... and for other signatures
```

**Arguments**

<code>x</code>	an object of class <code>Histogram</code> .
<code>Dataset</code>	a data frame of size $n \times d$ containing $d$ -dimensional dataset. Each of the $d$ columns represents one random variable. Number of observations $n$ equals the number of rows in the dataset.
<code>K</code>	an integer or a vector of length $d$ containing numbers of bins $v$ .
<code>ymin</code>	a vector of length $d$ containing minimum observations.
<code>ymax</code>	a vector of length $d$ containing maximum observations.
<code>shrink</code>	logical. If <code>TRUE</code> the output is shrunk to its optimal size. The default value is <code>FALSE</code> .
<code>...</code>	currently not used.

**Methods**

`signature(x = "Histogram")` an object of class `Histogram`.

**Author(s)**

Marko Nagode

**Examples**

```
# Create three datasets.

set.seed(1)

n <- 15
```

```

Dataset1 <- as.data.frame(cbind(rnorm(n, 157, 8), rnorm(n, 71, 10)))
Dataset2 <- as.data.frame(cbind(rnorm(n, 244, 14), rnorm(n, 61, 29)))
Dataset3 <- as.data.frame(cbind(rnorm(n, 198, 8), rnorm(n, 252, 13)))

apply(Dataset1, 2, range)
apply(Dataset2, 2, range)
apply(Dataset3, 2, range)

# Bin the first dataset.

hist <- fhistogram(Dataset = Dataset1, K = c(4, 5), ymin = c(100.0, 0.0), ymax = c(300.0, 300.0))

# Bin the second dataset.

hist <- fhistogram(x = hist, Dataset = Dataset2)

# Bin the third dataset and shrink the hist object.

hist <- fhistogram(x = hist, Dataset = Dataset3, shrink = TRUE)

hist

```

galaxy

*Galaxy Dataset*

## Description

The unfilled survey of the Corona Borealis region contains the velocities of 82 galaxies from 6 well separated conic sections of space.

## Usage

```
data(galaxy)
```

## Format

galaxy is a data frame with 82 cases (rows) and 1 continuous variable (columns) called Velocity.

## Source

K. Roeder. Density estimation with confidence sets exemplified by superclusters and voids in the galaxies. *Journal of American Statistical Association*, 85(411):617-624, 1990. <https://www.jstor.org/stable/2289993>.

## References

S. Richardson and P. J. Green. On bayesian analysis of mixtures with an unknown number of components. *Journal of the Royal Statistical Society B*, 59(4):731-792, 1997. <https://www.jstor.org/stable/2985194>.

G. McLachlan and D. Peel. Contribution to the discussion of paper by s. richardson and p.j. green.

Journal of the Royal Statistical Society B, 59(4):779-780, 1997. <https://www.jstor.org/stable/2985194>.

M. Stephens. Bayesian analysis of mixture models with an unknown number of components - an alternative to reversible jump methods. The Annals of Statistics, 28(1):40-74, 2000. <https://www.jstor.org/stable/2673981>.

Histogram-class	Class "Histogram"
-----------------	-------------------

## Description

Object of class `Histogram`.

## Objects from the Class

Objects can be created by calls of the form `new("Histogram", ...)`. Accessor methods for the slots are `a.Y(x = NULL)`, `a.K(x = NULL)`, `a.ymin(x = NULL)`, `a.ymax(x = NULL)`, `a.y0(x = NULL)`, `a.h(x = NULL)`, `a.n(x = NULL)` and `a.ns(x = NULL)`.

## Slots

- Y:** a data frame of size  $v \times (d + 1)$  containing  $d$ -dimensional histogram. Each of the first  $d$  columns represents one random variable and contains bin means  $\bar{y}_1, \dots, \bar{y}_v$ . Column  $d + 1$  contains frequencies  $k_1, \dots, k_v$ .
- K:** an integer or a vector of length  $d$  containing numbers of bins  $v$ .
- ymin:** a vector of length  $d$  containing minimum observations.
- ymax:** a vector of length  $d$  containing maximum observations.
- y0:** a vector of length  $d$  containing origins.
- h:** a vector of length  $d$  containing bin widths.
- n:** an integer containing total number  $n$  of observations.
- ns:** an integer containing number  $n_s$  of samples.

## Author(s)

Marko Nagode

## Examples

```

Y <- as.data.frame(matrix(1.0, nrow = 8, ncol = 3))

hist <- new("Histogram", Y = Y, K = c(4, 2), ymin = c(2, 1), ymax = c(10, 8))

a.Y(hist)
a.K(hist)
a.ymin(hist)
a.ymax(hist)
a.y0(hist)
a.h(hist)
a.n(hist)
a.ns(hist)

```

```

a.h(hist)
a.n(hist)
a.ns(hist)

# Multiplay Y[ , d + 1] by 0.1.

a.Y(hist) <- 0.1

```

## Description

Returns the Hannan-Quinn information criterion at pos.

## Usage

```

## S4 method for signature 'REBMIX'
HQC(x = NULL, pos = 1, ...)
## ... and for other signatures

```

## Arguments

- |     |  |
|-----|--|
| x   | see Methods section below.   |
| pos | a desired row number in x@summary for which the information criterion is calculated. The default value is 1. |
| ... | currently not used.  |

## Methods

- signature(x = "REBMIX") an object of class REBMIX.
- signature(x = "REBMVNORM") an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- E. J. Hannan and B. G. Quinn. The determination of the order of an autoregression. Journal of the Royal Statistical Society. Series B, 41(2):190-195, 1979. <https://www.jstor.org/stable/2985032>.

---

ICL-methods*Integrated Classification Likelihood Criterion*

---

**Description**

Returns the integrated classification likelihood criterion at pos.

**Usage**

```
## S4 method for signature 'REBMIX'
ICL(x = NULL, pos = 1, ...)
## ... and for other signatures
```

**Arguments**

x	see Methods section below.
pos	a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
...	currently not used.

**Methods**

```
signature(x = "REBMIX") an object of class REBMIX.
signature(x = "REBMVNORM") an object of class REBMVNORM.
```

**Author(s)**

Marko Nagode

**References**

C. Biernacki, G. Celeux and G. Govaert. Assessing a mixture model for clustering with the integrated classification likelihood. Technical Report 3521, INRIA, Rhone-Alpes, 1998.

---

ICLBIC-methods*Approximate Integrated Classification Likelihood Criterion*

---

**Description**

Returns the approximate integrated classification likelihood criterion at pos.

**Usage**

```
## S4 method for signature 'REBMIX'
ICLBIC(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- x see Methods section below.
- pos a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
- ... currently not used.

## Methods

- `signature(x = "REBMIX")` an object of class REBMIX.
- `signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- C. Biernacki, G. Celeux and G. Govaert. Assessing a mixture model for clustering with the integrated classification likelihood. Technical Report 3521, INRIA, Rhone-Alpes, 1998.

**iris**

*Iris Data Set*

## Description

This is perhaps the best known database to be found in the pattern recognition literature. Fisher's paper is a classic in the field and is referenced frequently to this day. The data set contains 3 classes of 50 instances each, where each class refers to a type of iris plant. One class is linearly separable from the other 2; the latter are NOT linearly separable from each other.

## Usage

```
data(iris)
```

## Format

`iris` is a data frame with 150 cases (rows) and 5 variables (columns) named:

1. Sepal.Length continuous.
2. Sepal.Width continuous.
3. Petal.Length continuous.
4. Petal.Width continuous.
5. Class discrete `iris-setosa`, `iris-versicolour` or `iris-virginica`.

## Source

A. Asuncion and D. J. Newman. Uci machine learning repository, 2007. <http://archive.ics.uci.edu/ml/>.

## References

R. A. Fisher. The use of multiple measurements in taxonomic problems. Annals of Eugenics, 7(2):179-188, 1936.

## Examples

```
## Not run:
devAskNewPage(ask = TRUE)

data(iris)

# Show level attributes.

levels(iris[["Class"]])

# Split dataset into train (75

set.seed(5)

Iris <- split(p = 0.6, Dataset = iris, class = 5)

# Estimate number of components, component weights and component
# parameters for train subsets.

n <- range(a.ntrain(Iris))

irisest <- REBMIX(model = "REBMVNORM",
Dataset = a.train(Iris),
Preprocessing = "histogram",
cmax = 10,
Criterion = "ICL-BIC",
EMcontrol = new("EM.Control", strategy = "single"))

plot(irisest, pos = 1, nrow = 3, ncol = 2, what = c("pdf"))
plot(irisest, pos = 2, nrow = 3, ncol = 2, what = c("pdf"))
plot(irisest, pos = 3, nrow = 3, ncol = 2, what = c("pdf"))

# Selected chunks.

iriscla <- RCLSMIX(model = "RCLSMVNORM",
x = list(irisest),
Dataset = a.test(Iris),
Zt = a.Zt(Iris))

iriscla

summary(iriscla)
```

```
# Plot selected chunks.

plot(iriscla, nrow = 3, ncol = 2)

## End(Not run)
```

kseq

*Sequence of Bins or Nearest Neighbours Generation***Description**

Returns (invisibly) a vector containing numbers of bins  $v$  for the histogram and the kernel density estimation or numbers of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**Usage**

```
kseq(from = NULL, to = NULL, f = 0.05, ...)
```

**Arguments**

- |      |   |
|------|---|
| from | starting value of the sequence. The default value is NULL.  |
| to   | end value of the sequence. The default value is NULL.   |
| f    | number specifying the fraction by which the bins or nearest neighbours should be separated $0.0 < f < 1.0$ . The default value is 0.05. |
| ...  | currently not used.   |

**Author(s)**

Marko Nagode

**Examples**

```
# Generate numbers of bins.

n <- 10000

Sturges <- as.integer(1 + log2(n)) # Minimum v follows Sturges rule.
Log10 <- as.integer(10 * log10(n)) # Maximum v follows Log10 rule.
RootN <- as.integer(2 * n^0.5) # Maximum v follows RootN rule.

K <- kseq(from = Sturges, to = Log10, f = 0.05)

K

K <- kseq(from = Sturges, to = RootN, f = 0.03)

K
```

---

 labelmoments-methods *Label Image Moments*


---

## Description

Returns the list with the data frame  $M_{ij}$  containing the cluster levels  $l$ , the numbers of pixels  $n$  and the cluster moments  $\mathbf{M} = (M_{10}, M_{01}, M_{11})^\top$  for 2D images or the data frame  $M_{ijk}$  containing the cluster levels  $l$ , the numbers of voxels  $n$  and the cluster moments  $\mathbf{M} = (M_{100}, M_{010}, M_{001}, M_{111})^\top$  for 3D images and the adjacency matrix  $A$  of size  $c_{\max} \times c_{\max}$ . It may have some NA rows and columns. To calculate the adjacency matrix  $A(i, j) = \exp(-\|\mathbf{M}_i - \mathbf{M}_j\|^2/2\sigma^2)$ , the raw cluster moments are first converted into z-scores.

## Usage

```
## S4 method for signature 'array'
labelmoments(Zp = array(), cmax = integer(), Sigma = 1.0, ...)
## ... and for other signatures
```

## Arguments

Zp	a 2D array of size $width \times height$ or a 3D array of size $width \times height \times depth$ containing the predictive cluster membership $\Omega_l$ , where $l \in \{0, 1, \dots, c\}$ . The cluster $l = 0$ may contain the pixels for 2D images or the voxels for 3D images, which are ignored by this method.
cmax	maximum number of clusters $c_{\max} \geq c$ .
Sigma	scale parameter $\sigma$ for the Gaussian kernel function $\exp(-\ \mathbf{M}_i - \mathbf{M}_j\ ^2/2\sigma^2)$ . The default value is 1.0.
...	currently not used.

## Methods

signature(Zp = "array") an array.

## Author(s)

Marko Nagode, Branislav Panic

## References

A. Ng, M. Jordan and Y. Weiss. On spectral clustering: Analysis and an algorithm. Advances in Neural Information Processing Systems 14 (NIPS 2001).

## Examples

```
Zp <- matrix(rep(0, 100), nrow = 10, ncol = 10)

Zp[2, 2:4] <- 1;
Zp[2:4, 5] <- 2;
Zp[8, 7:10] <- 3;
Zp[9, 6] <- 4; Zp[10, 5] <- 4
Zp[10, 1:4] <- 5
Zp[6:9, 1] <- 6

labelmoments <- labelmoments(Zp, cmax = 6, Sigma = 1.0)

set.seed(12)

mergelabels <- mergelabels(list(labelmoments$A), w = 1.0, k = 2, nstart = 3)

Zp

mergelabels
```

logL

*Log Likelihood*

## Description

Returns the log likelihood at pos.

## Usage

```
## S4 method for signature 'REBMIX'
logL(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- x see Methods section below.
- pos a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
- ... currently not used.

## Methods

- signature(x = "REBMIX") an object of class REBMIX.
- signature(x = "REBMVNORM") an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

G. McLachlan and D. Peel. Finite Mixture Models. John Wiley & Sons, New York, 2000.

mapclusters-methods    *Map Clusters*

## Description

Returns a factor of predictive cluster membership for dataset.

## Usage

```
## S4 method for signature 'RCLRMIX'
mapclusters(x = NULL, Dataset = data.frame(),
            s = expression(c), ...)
## ... and for other signatures
```

## Arguments

- |         |  |
|---------|--|
| x       | see Methods section below.   |
| Dataset | a data frame of size $n \times d$ containing $d$ -dimensional dataset. Each of the $d$ columns represents one random variable. Number of observations $n$ equal the number of rows in the dataset. |
| s       | a desired number of clusters to be created. The default value is <code>expression(c)</code> .  |
| ...     | currently not used.  |

## Methods

- `signature(x = "RCLRMIX")` an object of class RCLRMIX.
- `signature(x = "RCLRMVNORM")` an object of class RCLRMVNORM.

## Author(s)

Marko Nagode, Branislav Panic

## Examples

```
devAskNewPage(ask = TRUE)

# Generate normal dataset.

n <- c(50, 20, 40)

Theta <- new("RNGMVNORM.Theta", c = 3, d = 2)

a.theta1(Theta, 1) <- c(3, 10)
a.theta1(Theta, 2) <- c(8, 6)
```

```

a.theta1(Theta, 3) <- c(12, 11)
a.theta2(Theta, 1) <- c(3, 0.3, 0.3, 2)
a.theta2(Theta, 2) <- c(5.7, -2.3, -2.3, 3.5)
a.theta2(Theta, 3) <- c(2, 1, 1, 2)

normal <- RNGMIX(model = "RNGMVNORM", Dataset.name = paste("normal_", 1:10, sep = ""),
n = n, Theta = a.Theta(Theta))

# Convert all datasets to single histogram.

hist <- NULL

n <- length(normal@Dataset)

hist <- fhistogram(Dataset = normal@Dataset[[1]], K = c(10, 10),
ymin = a.ymin(normal), ymax = a.ymax(normal))

for (i in 2:n) {
  hist <- fhistogram(x = hist, Dataset = normal@Dataset[[i]], shrink = i == n)
}

# Estimate number of components, component weights and component parameters.

normalest <- REBMIX(model = "REBMVNORM",
Dataset = list(hist),
Preprocessing = "histogram",
cmax = 6,
Criterion = "BIC")

summary(normalest)

# Plot finite mixture.

plot(normalest)

# Cluster dataset.

normalclu <- RCLRMIX(model = "RCLRMVNORM", x = normalest)

# Plot clusters.

plot(normalclu)

summary(normalclu)

# Map clusters.

Zp <- mapclusters(x = normalclu, Dataset = a.Dataset(normal, 4))

Zt <- a.Zt(normal)

Zp

```

`zt`

MDL-methods	<i>Minimum Description Length</i>
-------------	-----------------------------------

## Description

Returns the minimum description length at pos.

## Usage

```
## S4 method for signature 'REBMIX'
MDL2(x = NULL, pos = 1, ...)
## S4 method for signature 'REBMIX'
MDL5(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- `x` see Methods section below.
- `pos` a desired row number in `x@summary` for which the information criterion is calculated. The default value is 1.
- `...` currently not used.

## Methods

- `signature(x = "REBMIX")` an object of class REBMIX.
- `signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- M. H. Hansen and B. Yu. Model selection and the principle of minimum description length. *Journal of the American Statistical Association*, 96(454):746-774, 2001. <https://www.jstor.org/stable/2670311>.

---

 mergelabels-methods      *Merge Labels Based on Probability Adjacency Matrix*


---

## Description

Returns the list with the normalised adjacency matrix  $L$  of size  $c \times c$ . The normalised adjacency matrix  $L = D^{-1/2}PD^{-1/2}$  depends on the probability adjacency matrix  $P(i, j) = \sum_{l=1}^n p_l A_l(i, j)$ , where  $p_l = w_l / \sum_{i=1}^c \sum_{j=i+1}^c A_l(i, j)$  and the degree matrix  $D(i, i) = \sum_{j=1}^c P(i, j)$ . The  $A_l$  matrices may contain some NA rows and columns, which are eliminated by the method. The list also contains the vector of integers `cluster` of length  $k$ , which indicates the cluster to which each label is assigned.

## Usage

```
## S4 method for signature 'list'
mergelabels(A = list(), w = numeric(), k = 2, ...)
## ... and for other signatures
```

## Arguments

- `A` a list of length  $n$  of adjacency matrices  $A_l$  of size  $c_{\max} \times c_{\max}$ , where  $c_{\max} \geq c$ .
- `w` vector of length  $n$  containing weights  $w_l$  summing to 1.
- `k` number of clusters  $k$  for `kmeans`. The default value is 2.
- `...` further arguments to `kmeans`.

## Methods

`signature(A = "list")` a list.

## Author(s)

Marko Nagode, Branislav Panic

## References

A. Ng, M. Jordan and Y. Weiss. On spectral clustering: Analysis and an algorithm. Advances in Neural Information Processing Systems 14 (NIPS 2001).

## Examples

```
Zp <- array(0, dim = c(10, 10, 2))

Zp[ , ,1][10, 1:4] <- 1
Zp[ , ,1][1:4, 10] <- 2

Zp[ , ,2][9, 1:5] <- 3
Zp[ , ,2][1:6, 9] <- 4
```

```

labelmoments <- labelmoments(Zp, cmax = 4, Sigma = 1.0)

labelmoments

set.seed(3)

mergelabels <- mergelabels(list(labelmoments$A), w = 1.0, k = 2, nstart = 5)

Zp

mergelabels

```

## Description

Returns the matrix of size  $n_D \times d$  containing optimal numbers of bins  $v_1, \dots, v_d$  for all processed datasets.

## Usage

```

## S4 method for signature 'list'
optbins(Dataset = list(), Rule = "Knuth equal",
        ymin = numeric(), ymax = numeric(), kmin = numeric(),
        kmax = numeric(), ...)
## ... and for other signatures

```

## Arguments

Dataset	a list of length $n_D$ of data frames of size $n \times d$ containing $d$ -dimensional datasets. Each of the $d$ columns represents one random variable. Numbers of observations $n$ equal the number of rows in the datasets.
Rule	a character giving the histogram binning rule. One of "Sturges", "Log10", "RootN", default "Knuth equal" or "Knuth unequal".
ymin	a vector of length $d$ containing minimum observations. The default value is <code>numeric()</code> .
ymax	a vector of length $d$ containing maximum observations. The default value is <code>numeric()</code> .
kmin	lower limit of the number of bins. The default value is <code>numeric()</code> .
kmax	upper limit of the number of bins. The default value is <code>numeric()</code> .
...	currently not used.

## Methods

`signature(x = "list")` a list of data frames.

## Author(s)

Branislav Panic, Marko Nagode

## References

K. K. Knuth. Optimal data-based binning for histograms and histogram-based probability density models. Digital Signal Processing, 95:102581, 2019. [doi:10.1016/j.dsp.2019.102581](https://doi.org/10.1016/j.dsp.2019.102581).

B. Panic, J. Klemenc, M. Nagode. Improved initialization of the EM algorithm for mixture model parameter estimation. Mathematics, 8(3):373, 2020. [doi:10.3390/math8030373](https://doi.org/10.3390/math8030373).

## Examples

```
# Generate multivariate normal datasets.

n <- c(750, 1000)

Theta <- new("RNGMVNORM.Theta", c = 2, d = 2)

a.theta1(Theta, 1) <- c(8, 6)
a.theta1(Theta, 2) <- c(6, 8)
a.theta2(Theta, 1) <- c(8, 2, 2, 4)
a.theta2(Theta, 2) <- c(2, 1, 1, 4)

sim2d <- RNGMIX(model = "RNGMVNORM",
  Dataset.name = paste("sim2d_", 1:5, sep = ""),
  rseed = -1,
  n = n,
  Theta = a.Theta(Theta))

# Calculate optimal numbers of bins.

opt.k <- optbins(Dataset = sim2d@Dataset,
  Rule = "Knuth equal",
  ymin = sim2d@ymin,
  ymax = sim2d@ymax,
  kmin = 2,
  kmax = 20)

opt.k

# Create object of class EM.Control.

EM <- new("EM.Control", strategy = "exhaustive", variant = "EM",
  acceleration = "fixed", acceleration.multiplier = 1.0, tolerance = 1.0E-4,
  maximum.iterations = 1000)

# Estimate number of components, component weights and component parameters.

sim2dest <- REBMIX(model = "REBMVNORM",
  Dataset = a.Dataset(sim2d),
  Preprocessing = "h",
```

```

cmax = 10,
ymin = a.ymin(sim2d),
ymax = a.ymax(sim2d),
K = opt.k,
Criterion = "BIC",
EMcontrol = EM)

# Plot finite mixture.

plot(sim2dest, pos = 3, nrow = 4, what = c("pdf", "marginal pdf", "IC"))

# Estimate number of components, component weights and component
# parameters for well known Iris dataset.

Dataset <- list(iris[, c(1:4)])

# Calculate optimal numbers of bins using non-equal number of bins in each dimension.

opt.k <- optbins(Dataset = Dataset,
  Rule = "Knuth unequal",
  kmin = 2,
  kmax = 20)

opt.k

# Estimate number of components, component weights and component parameters.

irisest <- REBMIX(model = "REBMVNORM",
  Dataset = Dataset,
  Preprocessing = "h",
  cmax = 10,
  K = opt.k,
  Criterion = "BIC",
  EMcontrol = EM)

irisest

```

**Description**

Returns the partition coefficient of Bezdek at pos.

**Usage**

```

## S4 method for signature 'REBMIX'
PC(x = NULL, pos = 1, ...)
## ... and for other signatures

```

## Arguments

- x see Methods section below.
- pos a desired row number in `x@summary` for which the information criterion is calculated. The default value is 1.
- ... currently not used.

## Methods

`signature(x = "REBMIX")` an object of class REBMIX.  
`signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

G. McLachlan and D. Peel. Finite Mixture Models. John Wiley & Sons, New York, 2000.

## Description

Returns the data frame containing observations  $\mathbf{x}_1, \dots, \mathbf{x}_n$  and empirical distribution functions  $F_1, \dots, F_n$ . Vectors  $\mathbf{x}$  are subvectors of  $\mathbf{y} = (y_1, \dots, y_d)^\top$ .

## Usage

```
## S4 method for signature 'REBMIX'
pemix(x = NULL, pos = 1, variables = expression(1:d),
      lower.tail = TRUE, log.p = FALSE, ...)
## ... and for other signatures
```

## Arguments

- x see Methods section below.
- pos a desired row number in `x@summary` for which the empirical distribution functions are calculated. The default value is 1.
- variables a vector containing indices of variables in subvectors  $\mathbf{x}$ . The default value is `1:d`.
- lower.tail logical. If TRUE, probabilities are  $P[X \leq x]$ , otherwise,  $P[X > x]$ . The default value is TRUE.
- log.p logical. if TRUE, probabilities  $p$  are given as  $\log(p)$ . The default value is FALSE.
- ... currently not used.

## Methods

`signature(x = "REBMIX")` an object of class REBMIX.  
`signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- M. Nagode and M. Fajdiga. The rebmix algorithm for the univariate finite mixture estimation. *Communications in Statistics - Theory and Methods*, 40(5):876-892, 2011a. doi:[10.1080/03610920903480890](https://doi.org/10.1080/03610920903480890).
- M. Nagode and M. Fajdiga. The rebmix algorithm for the multivariate finite mixture estimation. *Communications in Statistics - Theory and Methods*, 40(11):2022-2034, 2011b. doi:[10.1080/03610921003725788](https://doi.org/10.1080/03610921003725788).
- M. Nagode. Finite mixture modeling via REBMIX. *Journal of Algorithms and Optimization*, 3(2):14-28, 2015. <https://repozitorij.uni-lj.si/Dokument.php?id=127674&lang=eng>.

## Examples

```
# Generate simulated dataset.

n <- c(15, 15)

Theta <- new("RNGMIX.Theta", c = 2, pdf = rep("normal", 3))

a.theta1(Theta, 1) <- c(10, 20, 30)
a.theta1(Theta, 2) <- c(3, 4, 5)
a.theta2(Theta, 1) <- c(3, 2, 1)
a.theta2(Theta, 2) <- c(15, 10, 5)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1:4, sep = ""),
  rseed = -1,
  n = n,
  Theta = a.Theta(Theta))

# Create object of class EM.Control.

EM <- new("EM.Control", strategy = "exhaustive", variant = "ECM",
  acceleration = "fixed", acceleration.multiplier = 1.0, tolerance = 1.0E-4,
  maximum.iterations = 1000)

# Estimate number of components, component weights and component parameters.

simulatedest <- REBMIX(Dataset = a.Dataset(simulated),
  Preprocessing = "kernel density estimation",
  cmax = 4,
  pdf = c("n", "n", "n"),
  EMcontrol = EM)
```

```
# Preprocess simulated dataset.

f <- pemix(simulatedest, pos = 3, variables = c(1))

f
```

## Description

Returns the data frame containing observations  $\mathbf{x}_1, \dots, \mathbf{x}_n$  and predictive marginal distribution functions  $F(\mathbf{x}|c, \mathbf{w}, \Theta)$ . Vectors  $\mathbf{x}$  are subvectors of  $\mathbf{y} = (y_1, \dots, y_d)^\top$ . If  $\mathbf{x} = \mathbf{y}$  the method returns the data frame containing observations  $y_1, \dots, y_n$  and the corresponding predictive mixture distribution function  $F(\mathbf{y}|c, \mathbf{w}, \Theta)$ .

## Usage

```
## S4 method for signature 'REBMIX'
pfmix(x = NULL, Dataset = data.frame(), pos = 1,
       variables = expression(1:d), lower.tail = TRUE, log.p = FALSE, ...)
## ... and for other signatures
```

## Arguments

<code>x</code>	see Methods section below.
<code>Dataset</code>	a data frame containing observations $\mathbf{y} = (y_1, \dots, y_d)^\top$ for which the predictive marginal distribution functions are calculated. The default value is <code>data.frame()</code> .
<code>pos</code>	a desired row number in <code>x@summary</code> for which the predictive marginal distribution functions are calculated. The default value is 1.
<code>variables</code>	a vector containing indices of variables in subvectors $\mathbf{x}$ . The default value is <code>1:d</code> .
<code>lower.tail</code>	logical. If TRUE, probabilities are $P[X \leq x]$ , otherwise, $P[X > x]$ . The default value is TRUE.
<code>log.p</code>	logical. if TRUE, probabilities $p$ are given as $\log(p)$ . The default value is FALSE.
...	currently not used.

## Methods

`signature(x = "REBMIX")` an object of class REBMIX.  
`signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- M. Nagode and M. Fajdiga. The rebmix algorithm for the univariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(5):876-892, 2011a. doi:[10.1080/03610920903480890](https://doi.org/10.1080/03610920903480890).
- M. Nagode and M. Fajdiga. The rebmix algorithm for the multivariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(11):2022-2034, 2011b. doi:[10.1080/03610921003725788](https://doi.org/10.1080/03610921003725788).
- M. Nagode. Finite mixture modeling via REBMIX. Journal of Algorithms and Optimization, 3(2):14-28, 2015. <https://repozitorij.uni-lj.si/Dokument.php?id=127674&lang=eng>.

## Examples

```
# Generate simulated dataset.

n <- c(15, 15)

Theta <- new("RNGMIX.Theta", c = 2, pdf = rep("normal", 3))

a.theta1(Theta, 1) <- c(10, 20, 30)
a.theta1(Theta, 2) <- c(3, 4, 5)
a.theta2(Theta, 1) <- c(3, 2, 1)
a.theta2(Theta, 2) <- c(15, 10, 5)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1:4, sep = ""),
  rseed = -1,
  n = n,
  Theta = a.Theta(Theta))

# Number of classes or nearest neighbours to be processed.

K <- c(as.integer(1 + log2(sum(n))), # Minimum v follows Sturges rule.
       as.integer(10 * log10(sum(n)))) # Maximum v follows log10 rule.

# Estimate number of components, component weights and component parameters.

simulatedest <- REBMIX(Dataset = a.Dataset(simulated),
  Preprocessing = "h",
  cmax = 4,
  Criterion = "BIC",
  pdf = c("n", "n", "n"))

# Preprocess simulated dataset.

Dataset <- data.frame(c(25, 5, -20), NA, c(31, 20, 20))

f <- pfmix(simulatedest, Dataset = Dataset, pos = 3, variables = c(1, 3))

f

# Plot finite mixture.
```

```

opar <- plot(simulatedest, pos = 3, nrow = 3, ncol = 1,
             what = "pdf", contour.drawlabels = TRUE, contour.labcex = 0.6)

par(usr = opar[[2]]$usr, mfg = c(2, 1))

points(x = f[, 1], y = f[, 2])

text(x = f[, 1], y = f[, 2], labels = format(f[, 3], digits = 3), cex = 0.8, pos = 4)

```

**plot-methods***Plots RNGMIX, REBMIX, RCLRMIX and RCLSMIX Output***Description**

Plots true clusters if  $x$  equals "RNGMIX". Plots the REBMIX output depending on what argument if  $x$  equals "REBMIX". Plots predictive clusters if  $x$  equals "RCLRMIX". Wrongly clustered observations are plotted only if  $x@Zt$  is available. Plots predictive classes and wrongly classified observations if  $x$  equals "RCLSMIX".

**Usage**

```

## S4 method for signature 'RNGMIX,missing'
plot(x, y, pos = 1, nrow = 1, ncol = 1, cex = 0.8,
      fg = "black", lty = "solid", lwd = 1, pty = "m", tcl = 0.5,
      plot.cex = 0.8, plot.pch = 19, ...)
## S4 method for signature 'REBMIX,missing'
plot(x, y, pos = 1, what = c("pdf"),
      nrow = 1, ncol = 1, npts = 200, n = 200, cex = 0.8, fg = "black",
      lty = "solid", lwd = 1, pty = "m", tcl = 0.5,
      plot.cex = 0.8, plot.pch = 19, contour.drawlabels = FALSE,
      contour.labcex = 0.8, contour.method = "flattest",
      contour.nlevels = 12, log = "", ...)
## S4 method for signature 'RCLRMIX,missing'
plot(x, y, s = expression(c), nrow = 1, ncol = 1, cex = 0.8,
      fg = "black", lty = "solid", lwd = 1, pty = "m", tcl = 0.5,
      plot.cex = 0.8, plot.pch = 19, ...)
## S4 method for signature 'RCLSMIX,missing'
plot(x, y, nrow = 1, ncol = 1, cex = 0.8,
      fg = "black", lty = "solid", lwd = 1, pty = "m", tcl = 0.5,
      plot.cex = 0.8, plot.pch = 19, ...)
## ... and for other signatures

```

**Arguments**

- $x$  see Methods section below.
- $y$  currently not used.

pos	a desired row number in <code>x@summary</code> to be plotted. The default value is 1.
s	a desired number of clusters to be plotted. The default value is <code>expression(c)</code> .
what	a character vector giving the plot types. One of "pdf" for probability density function, "marginal pdf" for marginal probability density function, "IC" for information criterion depending on numbers of components $c$ , "logL" for log likelihood, "D" for total of positive relative deviations, "K" for information criterion depending on bins $v$ or numbers of nearest neighbours $k$ , "cdf" for univariate distribution function or "marginal cdf" for marginal distribution function. The default value is "pdf".
nrow	a desired number of rows in which the empirical and predictive densities are to be plotted. The default value is 1.
ncol	a desired number of columns in which the empirical and predictive densities are to be plotted. The default value is 1.
npts	a number of points at which the predictive densities are to be plotted. The default value is 200.
n	a number of observations to be plotted. The default value is 200.
cex	a numerical value giving the amount by which the plotting text and symbols should be magnified relative to the default, see also <code>par</code> . The default value is 0.8.
fg	a colour used for things like axes and boxes around plots, see also <code>par</code> . The default value is "black".
lty	a line type, see also <code>par</code> . The default value is "solid".
lwd	a line width, see also <code>par</code> . The default value is 1.
pty	a character specifying the type of the plot region to be used. One of "s" generating a square plotting region or "m" generating the maximal plotting region. The default value is "m".
tcl	a length of tick marks as a fraction of the height of a line of the text, see also <code>par</code> . The default value is 0.5.
plot.cex	a numerical vector giving the amount by which plotting characters and symbols should be scaled relative to the default. It works as a multiple of <code>par("cex")</code> . NULL and NA are equivalent to 1.0. Note that this does not affect annotation, see also <code>plot.default</code> . The default value is 0.8.
plot.pch	a vector of plotting characters or symbols, see also <code>points</code> . The default value is 19.
contour.drawlabels	logical. The contours are labelled if TRUE. The default value is FALSE.
contour.labcex	cex for contour labelling. The default value is 0.8. This is an absolute size, not a multiple of <code>par("cex")</code> .
contour.method	a character specifying where the labels will be located. The possible values are "simple", "edge" and default "flattest", see also <code>contour</code> .
contour.nlevels	a number of desired contour levels. The default value is 12.

- `log` a character which contains "x" if the x axis is to be logarithmic, "y" if the y axis is to be logarithmic and "xy" or "yx" if both axes are to be logarithmic. The default value is "".
- `...` further arguments to `par`.

### Value

Returns (invisibly) a list containing graphical parameters `par`. Such a list can be passed as an argument to `par` to restore the parameter values.

### Methods

```
signature(x = "RNGMIX", y = "missing") an object of class RNGMIX.
signature(x = "RNGMVNORM", y = "missing") an object of class RNGMVNORM.
signature(x = "REBMIX", y = "missing") an object of class REBMIX.
signature(x = "REBMVNORM", y = "missing") an object of class REBMVNORM.
signature(x = "RCLRMIX", y = "missing") an object of class RCLRMIX.
signature(x = "RCLRMVNORM", y = "missing") an object of class RCLRMVNORM.
signature(x = "RCLSMIX", y = "missing") an object of class RCLSMIX.
signature(x = "RCLSMVNORM", y = "missing") an object of class RCLSMVNORM.
```

### Author(s)

Marko Nagode

### References

C. M. Bishop. Neural Networks for Pattern Recognition. Clarendon Press, Oxford, 1995.

### Examples

```
## Not run:
devAskNewPage(ask = TRUE)

data(wine)

colnames(wine)

# Remove Cultivar column from wine dataset.

winecolnames <- !(colnames(wine))

wine <- wine[, winecolnames]

# Determine number of dimensions d and wine dataset size n.

d <- ncol(wine)
n <- nrow(wine)
```

```
wineest <- REBMIX(model = "REBMVNORM",
  Dataset = list(wine = wine),
  Preprocessing = "kernel density estimation",
  Criterion = "ICL-BIC",
  EMcontrol = new("EM.Control", strategy = "best"))

# Plot finite mixture.

plot(wineest, what = c("pdf", "IC", "logL", "D"),
  nrow = 2, ncol = 2, pty = "s")

## End(Not run)
```

**PRD-methods***Total of Positive Relative Deviations***Description**

Returns the total of positive relative deviations D at pos.

**Usage**

```
## S4 method for signature 'REBMIX'
PRD(x = NULL, pos = 1, ...)
## ... and for other signatures
```

**Arguments**

- x see Methods section below.
- pos a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
- ... currently not used.

**Methods**

- `signature(x = "REBMIX")` an object of class REBMIX.
- `signature(x = "REBMVNORM")` an object of class REBMVNORM.

**Author(s)**

Marko Nagode

## References

- M. Nagode and M. Fajdiga. The rebmix algorithm for the univariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(5):876-892, 2011a. doi:[10.1080/03610920903480890](https://doi.org/10.1080/03610920903480890).
- M. Nagode and M. Fajdiga. The rebmix algorithm for the multivariate finite mixture estimation. Communications in Statistics - Theory and Methods, 40(11):2022-2034, 2011b. doi:[10.1080/03610921003725788](https://doi.org/10.1080/03610921003725788).
- M. Nagode. Finite mixture modeling via REBMIX. Journal of Algorithms and Optimization, 3(2):14-28, 2015. <https://repozitorij.uni-lj.si/Dokument.php?id=127674&lang=eng>.

RCLRMIX-class

Class "RCLRMIX"

## Description

Object of class RCLRMIX.

## Objects from the Class

Objects can be created by calls of the form `new("RCLRMIX", ...)`. Accessor methods for the slots are `a.Dataset(x = NULL)`, `a.pos(x = NULL)`, `a.Zt(x = NULL)`, `a.Zp(x = NULL, s = expression(c))`, `a.c(x = NULL)`, `a.p(x = NULL, s = expression(c))`, `a.pi(x = NULL, s = expression(c))`, `a.P(x = NULL, s = expression(c))`, `a.tau(x = NULL, s = expression(c))`, `a.prob(x = NULL)`, `a.Rule(x = NULL)`, `a.from(x = NULL)`, `a.to(x = NULL)`, `a.EN(x = NULL)` and `a.ED(x = NULL)`, where `x` stands for an object of class RCLRMIX and `s` a desired number of clusters for which the slot is calculated.

## Slots

**x:** an object of class REBMIX.

**Dataset:** a data frame or an object of class Histogram to be clustered.

**pos:** a desired row number in `x@summary` for which the clustering is performed. The default value is 1.

**Zt:** a factor of true cluster membership.

**Zp:** a factor of predictive cluster membership.

**c:** number of nonempty clusters.

**p:** a vector of length  $c$  containing prior probabilities of cluster memberships  $p_l$  summing to 1. The value is returned only if all variables in slot `x` follow either binomial or Dirac parametric families. The default value is `numeric()`.

**pi:** a list of length  $d$  of matrices of size  $c \times K_i$  containing cluster conditional probabilities  $\pi_{ilk}$ . Let  $\pi_{ilk}$  denote the cluster conditional probability that an observation in cluster  $l = 1, \dots, c$  produces the  $k$ th outcome on the  $i$ th variable. Suppose we observe  $i = 1, \dots, d$  polytomous categorical variables (the manifest variables), each of which contains  $K_i$  possible outcomes for observations  $j = 1, \dots, n$ . A manifest variable is a variable that can be measured or

observed directly. It must be coded as whole number starting at zero for the first outcome and increasing to the possible number of outcomes minus one. It is presumed here that all variables are statistically independent and within clusters and that  $\mathbf{y}_1, \dots, \mathbf{y}_n$  stands for an observed  $d$  dimensional dataset of size  $n$  of vector observations  $\mathbf{y}_j = (y_{1j}, \dots, y_{ij}, \dots, y_{dj})^\top$ . The value is returned only if all variables in slot  $x$  follow either binomial or Dirac parametric families. The default value is `list()`.

- P:** a data frame containing true  $N_t(\mathbf{y}_j)$  and predictive  $N_p(\mathbf{y}_j)$  frequencies calculated for unique  $\mathbf{y}_{\tilde{j}} \in \{\mathbf{y}_1, \dots, \mathbf{y}_n\}$ , where  $\tilde{j} = 1, \dots, \tilde{n}$  and  $\tilde{n} \leq n$ .
- tau:** a matrix of size  $n \times c$  containing conditional probabilities  $\tau_{jl}$  that observations  $\mathbf{y}_1, \dots, \mathbf{y}_n$  arise from clusters  $1, \dots, c$ .
- prob:** a vector of length  $c$  containing probabilities of correct clustering for  $s = 1, \dots, c$ .
- Rule:** a character containing the merging rule. One of "Entropy" and "Demp". The default value is "Entropy".
- from:** a vector of length  $c - 1$  containing clusters merged to to clusters.
- to:** a vector of length  $c - 1$  containing clusters originating from from clusters.
- EN:** a vector of length  $c - 1$  containing entropies for combined clusters.
- ED:** a vector of length  $c - 1$  containing decrease of entropies for combined clusters.
- A:** an adjacency matrix of size  $c_{\max} \times c_{\max}$ , where  $c_{\max} \geq c$ .

### Author(s)

Marko Nagode, Branislav Panic

### References

J. P. Baudry, A. E. Raftery, G. Celeux, K. Lo and R. Gottardo. Combining mixture components for clustering. Journal of Computational and Graphical Statistics, 19(2):332-353, 2010. [doi:10.1198/jcgs.2010.08111](https://doi.org/10.1198/jcgs.2010.08111)

S. Kyoya and K. Yamanishi. Summarizing finite mixture model with overlapping quantification. Entropy, 23(11):1503, 2021. [doi:10.3390/e23111503](https://doi.org/10.3390/e23111503)

### Examples

```
devAskNewPage(ask = TRUE)

# Generate normal dataset.

n <- c(500, 200, 400)

Theta <- new("RNGMVNORM.Theta", c = 3, d = 2)

a.theta1(Theta, 1) <- c(3, 10)
a.theta1(Theta, 2) <- c(8, 6)
a.theta1(Theta, 3) <- c(12, 11)
a.theta2(Theta, 1) <- c(3, 0.3, 0.3, 2)
a.theta2(Theta, 2) <- c(5.7, -2.3, -2.3, 3.5)
a.theta2(Theta, 3) <- c(2, 1, 1, 2)
```

```

normal <- RNGMIX(model = "RNGMVNORM", Dataset.name = "normal_1", n = n, Theta = a.Theta(Theta))

# Estimate number of components, component weights and component parameters.

normalest <- REBMIX(model = "REBMVNORM",
Dataset = a.Dataset(normal),
Preprocessing = "histogram",
cmax = 6,
Criterion = "BIC")

summary(normalest)

# Plot finite mixture.

plot(normalest)

# Cluster dataset.

normalclu <- RCLRMIX(model = "RCLRMVNORM", x = normalest, Zt = a.Zt(normal))

# Plot clusters.

plot(normalclu)

summary(normalclu)

```

RCLRMIX-methods

*Predicts Cluster Membership Based Upon a Model Trained by REBMIX*

## Description

Returns as default the RCLRMIX algorithm output for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities, following the methodology proposed in the article cited in the references. If `model` equals "RCLRMVNORM" output for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices is returned.

## Usage

```

## S4 method for signature 'RCLRMIX'
RCLRMIX(model = "RCLRMIX", x = NULL, Dataset = NULL,
         pos = 1, Zt = factor(), Rule = character(), ...)
## ... and for other signatures
## S4 method for signature 'RCLRMIX'
summary(object, ...)
## ... and for other signatures

```

### Arguments

model	see Methods section below.
x	an object of class REBMIX.
Dataset	a data frame or an object of class Histogram to be clustered.
pos	a desired row number in x@summary for which the clustering is performed. The default value is 1.
Zt	a factor of true cluster membership. The default value is factor().
Rule	a character containing the merging rule. One of "Entropy" or "Demp". The default value is "Entropy".
object	see Methods section below.
...	currently not used.

### Value

Returns an object of class RCLRMIX or RCLRMVNORM.

### Methods

```
signature(model = "RCLRMIX") a character giving the default class name "RCLRMIX" for mix-
tures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial,
Poisson, Dirac, uniform or von Mises component densities.

signature(model = "RCLRMVNORM") a character giving the class name "RCLRMVNORM" for mix-
tures of multivariate normal component densities with unrestricted variance-covariance matri-
ces.

signature(object = "RCLRMIX") an object of class RCLRMIX.

signature(object = "RCLRMVNORM") an object of class RCLRMVNORM.
```

### Author(s)

Marko Nagode

### References

J. P. Baudry, A. E. Raftery, G. Celeux, K. Lo and R. Gottardo. Combining mixture components for clustering. Journal of Computational and Graphical Statistics, 19(2):332-353, 2010. [doi:10.1198/jcgs.2010.08111](https://doi.org/10.1198/jcgs.2010.08111)

### Examples

```
devAskNewPage(ask = TRUE)

# Generate Poisson dataset.

n <- c(500, 200, 400)

Theta <- new("RNGMIX.Theta", c = 3, pdf = "Poisson")
```

```

a.theta1(Theta) <- c(3, 12, 36)

poisson <- RNGMIX(Dataset.name = "Poisson_1", n = n, Theta = a.Theta(Theta))

# Estimate number of components, component weights and component parameters.

EM <- new("EM.Control", strategy = "exhaustive")

poissonest <- REBMIX(Dataset = a.Dataset(poisson),
  Preprocessing = "histogram",
  cmax = 6,
  Criterion = "BIC",
  pdf = rep("Poisson", 1),
  EMcontrol = EM)

summary(poissonest)

# Plot finite mixture.

plot(poissonest)

# Cluster dataset.

poissonclu <- RCLRMIX(x = poissonest, Zt = a.Zt(poisson))

summary(poissonclu)

# Plot clusters.

plot(poissonclu)

# Create new dataset.

Dataset <- sample.int(n = 50, size = 10, replace = TRUE)

Dataset <- as.data.frame(Dataset)

# Cluster the dataset.

poissonclu <- RCLRMIX(x = poissonest, Dataset = Dataset, Rule = "Demp")

a.Dataset(poissonclu)

```

RCLS.chunk-class      *Class "RCLS.chunk"*

### Description

Object of class RCLS.chunk.

### Objects from the Class

Objects can be created by calls of the form `new("RCLS.chunk", ...)`. Accessor methods for the slots are `a.s(x = NULL)`, `a.levels(x = NULL)`, `a.ntrain(x = NULL)`, `a.train(x = NULL)`, `a.Zr(x = NULL)`, `a.ntest(x = NULL)`, `a.test(x = NULL)` and `a.Zt(x = NULL)`, where `x` stands for an object of class `RCLS.chunk`.

### Slots

- s:** finite set of size  $s$  of classes  $\Omega = \{\Omega_g; g = 1, \dots, s\}$ .
- levels:** a character vector of length  $s$  containing class names  $\Omega_g$ .
- ntrain:** a vector of length  $s$  containing numbers of observations in train datasets  $Y_{\text{traing}}$ .
- train:** a list of length  $n_D$  of data frames containing train datasets  $Y_{\text{traing}}$  of length  $n_{\text{traing}}$ .
- Zr:** a list of factors of true class membership  $\Omega_g$  for the train datasets.
- ntest:** number of observations in test dataset  $Y_{\text{test}}$ .
- test:** a data frame containing test dataset  $Y_{\text{test}}$  of length  $n_{\text{test}}$ .
- Zt:** a factor of true class membership  $\Omega_g$  for the test dataset.

### Author(s)

Marko Nagode

### References

D. M. Dziuda. Data Mining for Genomics and Proteomics: Analysis of Gene and Protein Expression Data. John Wiley & Sons, New York, 2010.

### Description

Object of class `RCLSMIX`.

### Objects from the Class

Objects can be created by calls of the form `new("RCLSMIX", ...)`. Accessor methods for the slots are `a.o(x = NULL)`, `a.Dataset(x = NULL)`, `a.s(x = NULL)`, `a.ntrain(x = NULL)`, `a.P(x = NULL)`, `a.ntest(x = NULL)`, `a.Zt(x = NULL)`, `a.Zp(x = NULL)`, `a.CM(x = NULL)`, `a.Accuracy(x = NULL)`, `a.Error(x = NULL)`, `a.Precision(x = NULL)`, `a.Sensitivity(x = NULL)`, `a.Specificity(x = NULL)` and `a.Chunks(x = NULL)`, where `x` stands for an object of class `RCLSMIX`.

## Slots

**x:** a list of objects of class REBMIX of length  $o$  obtained by running REBMIX on  $g = 1, \dots, s$  train datasets  $Y_{\text{traing}}$  all of length  $n_{\text{traing}}$ . For the train datasets the corresponding class membership  $\Omega_g$  is known. This yields  $n_{\text{train}} = \sum_{g=1}^s n_{\text{traing}}$ , while  $Y_{\text{traing}} \cap Y_{\text{traing}} = \emptyset$  for all  $q \neq g$ . Each object in the list corresponds to one chunk, e.g.,  $(y_{1j}, y_{3j})^\top$ .

**o:** number of chunks  $o$ .  $Y = \{\mathbf{y}_j; j = 1, \dots, n\}$  is an observed  $d$ -dimensional dataset of size  $n$  of vector observations  $\mathbf{y}_j = (y_{1j}, \dots, y_{dj})^\top$  and is partitioned into train and test datasets. Vector observations  $\mathbf{y}_j$  may further be split into  $o$  chunks when running REBMIX, e.g., for  $d = 6$  and  $o = 3$  the set of chunks substituting  $\mathbf{y}_j$  may be as follows  $(y_{1j}, y_{3j})^\top$ ,  $(y_{2j}, y_{4j}, y_{6j})^\top$  and  $y_{5j}$ .

**Dataset:** a data frame containing test dataset  $Y_{\text{test}}$  of length  $n_{\text{test}}$ . For the test dataset the corresponding class membership  $\Omega_g$  is not known.

**s:** finite set of size  $s$  of classes  $\Omega = \{\Omega_g; g = 1, \dots, s\}$ .

**ntrain:** a vector of length  $s$  containing numbers of observations in train datasets  $Y_{\text{traing}}$ .

**P:** a vector of length  $s$  containing prior probabilities  $P(\Omega_g) = \frac{n_{\text{train}}}{n_{\text{train}}}$ .

**ntest:** number of observations in test dataset  $Y_{\text{test}}$ .

**Zt:** a factor of true class membership  $\Omega_g$  for the test dataset.

**Zp:** a factor of predictive class membership  $\Omega_g$  for the test dataset.

**CM:** a table containing confusion matrix for multiclass classifier. It contains number  $x_{qg}$  of test observations with the true class  $q$  that are classified into the class  $g$ , where  $q, g = 1, \dots, s$ .

**Accuracy:** proportion of all test observations that are classified correctly. Accuracy =  $\frac{\sum_{g=1}^s x_{gg}}{n_{\text{test}}}$ .

**Error:** proportion of all test observations that are classified wrongly. Error =  $1 - \text{Accuracy}$ .

**Precision:** a vector containing proportions of predictive observations in class  $g$  that are classified correctly into class  $g$ . Precision( $g$ ) =  $\frac{x_{gg}}{\sum_{q=1}^s x_{qg}}$ .

**Sensitivity:** a vector containing proportions of test observations in class  $g$  that are classified correctly into class  $g$ . Sensitivity( $g$ ) =  $\frac{x_{gg}}{\sum_{q=1}^s x_{qg}}$ .

**Specificity:** a vector containing proportions of test observations that are not in class  $g$  and are classified into the non  $g$  class. Specificity( $g$ ) =  $\frac{n_{\text{test}} - \sum_{q=1}^s x_{qg}}{n_{\text{test}} - \sum_{q=1}^s x_{qg}}$ .

**Chunks:** a vector containing selected chunks.

## Author(s)

Marko Nagode

## References

D. M. Dziuda. Data Mining for Genomics and Proteomics: Analysis of Gene and Protein Expression Data. John Wiley & Sons, New York, 2010.

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<b>RCLSMIX-methods</b>	<i>Predicts Class Membership Based Upon a Model Trained by REBMIX</i>
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## Description

Returns as default the RCLSMIX algorithm output for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities. If model equals "RCLSMVNORM" output for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices is returned.

## Usage

```
## S4 method for signature 'RCLSMIX'
RCLSMIX(model = "RCLSMIX", x = list(), Dataset = data.frame(),
         Zt = factor(), ...)
## ... and for other signatures
## S4 method for signature 'RCLSMIX'
summary(object, ...)
## ... and for other signatures
```

## Arguments

model	see Methods section below.
x	a list of objects of class REBMIX of length $o$ obtained by running <a href="#">REBMIX</a> on $g = 1, \dots, s$ train datasets $Y_{\text{train}g}$ all of length $n_{\text{train}g}$ . For the train datasets the corresponding class membership $\Omega_g$ is known. This yields $n_{\text{train}} = \sum_{g=1}^s n_{\text{train}g}$ , while $Y_{\text{train}q} \cap Y_{\text{train}g} = \emptyset$ for all $q \neq g$ . Each object in the list corresponds to one chunk, e.g., $(y_{1j}, y_{3j})^\top$ . The default value is <code>list()</code> .
Dataset	a data frame containing test dataset $Y_{\text{test}}$ of length $n_{\text{test}}$ . For the test dataset the corresponding class membership $\Omega_g$ is not known. The default value is <code>data.frame()</code> .
Zt	a factor of true class membership $\Omega_g$ for the test dataset. The default value is <code>factor()</code> .
object	see Methods section below.
...	currently not used.

## Value

Returns an object of class RCLSMIX or RCLSMVNORM.

## Methods

`signature(model = "RCLSMIX")` a character giving the default class name "RCLSMIX" for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities.

```

signature(model = "RCLSMVNORM") a character giving the class name "RCLSMVNORM" for mix-
tures of multivariate normal component densities with unrestricted variance-covariance matri-
ces.

signature(object = "RCLSMIX") an object of class RCLSMIX.

signature(object = "RCLSMVNORM") an object of class RCLSMVNORM.

```

### **Author(s)**

Marko Nagode

### **References**

R. O. Duda and P. E. Hart. Pattern Classification and Scene Analysis. John Wiley & Sons, New York, 1973.

### **Examples**

```

## Not run:
devAskNewPage(ask = TRUE)

data(adult)

# Find complete cases.

adult <- adult[complete.cases(adult),]

# Replace levels with numbers.

adult <- as.data.frame(data.matrix(adult))

# Find numbers of levels.

cmax <- unlist(lapply(apply(adult[, c(-1, -16)], 2, unique), length))

cmax

# Split adult dataset into train and test subsets for two Incomes
# and remove Type and Income columns.

Adult <- split(p = list(type = 1, train = 2, test = 1),
Dataset = adult, class = 16)

# Estimate number of components, component weights and component parameters
# for the set of chunks 1:14.

adulttest <- list()

for (i in 1:14) {
  adulttest[[i]] <- REBMIX(Dataset = a.train(chunk(Adult, i)),
  Preprocessing = "histogram",
  cmax = min(120, cmax[i]),
}

```

```

Criterion = "BIC",
pdf = "Dirac",
K = 1)
}

# Class membership prediction based upon the best first search algorithm.

adultcla <- BFSMIX(x = adultest,
Dataset = a.test(Adult),
Zt = a.Zt(Adult))

adultcla

summary(adultcla)

# Plot selected chunks.

plot(adultcla, nrow = 5, ncol = 2)

## End(Not run)

```

REBMIX-class

Class "REBMIX"

## Description

Object of class REBMIX.

## Objects from the Class

Objects can be created by calls of the form `new("REBMIX", ...)`. Accessor methods for the slots are `a.Dataset(x = NULL, pos = 0)`, `a.Preprocessing(x = NULL)`, `a.cmax(x = NULL)`, `a.cmin(x = NULL)`, `a.Criterion(x = NULL)`, `a.Variables(x = NULL)`, `a.pdf(x = NULL)`, `a.theta1(x = NULL)`, `a.theta2(x = NULL)`, `a.theta3(x = NULL)`, `a.K(x = NULL)`, `a.ymin(x = NULL)`, `a.ymax(x = NULL)`, `a.ar(x = NULL)`, `a.Restraints(x = NULL)`, `a.Mode(x = NULL)`, `a.w(x = NULL, pos = 0)`, `a.Theta(x = NULL, pos = 0)`, `a.summary(x = NULL, col.name = character(), pos = 0)`, `a.summary.EM(x = NULL, col.name = character(), pos = 0)`, `a.pos(x = NULL)`, `a.opt.c(x = NULL)`, `a.opt.IC(x = NULL)`, `a.opt.logL(x = NULL)`, `a.opt.Dmin(x = NULL)`, `a.opt.D(x = NULL)`, `a.all.K(x = NULL)`, `a.all.IC(x = NULL)`, `a.theta1.all(x = NULL, pos = 1)`, `a.theta2.all(x = NULL, pos = 1)` and `a.theta3.all(x = NULL, pos = 1)`, where `x`, `pos` and `col.name` stand for an object of class REBMIX, a desired slot item and a desired column name, respectively.

## Slots

**Dataset:** a list of length  $n_D$  of data frames or objects of class `Histogram`. Data frames should have size  $n \times d$  containing  $d$ -dimensional datasets. Each of the  $d$  columns represents one random variable. Numbers of observations  $n$  equal the number of rows in the datasets.

**Preprocessing:** a character vector giving the preprocessing types. One of "histogram", "kernel density estimation" or "k-nearest neighbour".

**cmax:** maximum number of components  $c_{\max} > 0$ . The default value is 15.

**cmin:** minimum number of components  $c_{\min} > 0$ . The default value is 1.

**Criterion:** a character giving the information criterion type. One of default Akaike "AIC", "AIC3", "AIC4" or "AICc", Bayesian "BIC", consistent Akaike "CAIC", Hannan-Quinn "HQC", minimum description length "MDL2" or "MDL5", approximate weight of evidence "AWE", classification likelihood "CLC", integrated classification likelihood "ICL" or "ICL-BIC", partition coefficient "PC", total of positive relative deviations "D" or sum of squares error "SSE".

**Variables:** a character vector of length  $d$  containing types of variables. One of "continuous" or "discrete".

**pdf:** a character vector of length  $d$  containing continuous or discrete parametric family types. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or "vonMises".

**theta1:** a vector of length  $d$  containing initial component parameters. One of  $n_{il} = \text{number of categories} - 1$  for "binomial" distribution.

**theta2:** a vector of length  $d$  containing initial component parameters. Currently not used.

**theta3:** a vector of length  $d$  containing initial component parameters. One of  $\xi_{il} \in \{-1, \text{NA}, 1\}$  for "Gumbel" distribution.

**K:** a character or a vector or a list of vectors containing numbers of bins  $v$  for the histogram and the kernel density estimation or numbers of nearest neighbours  $k$  for the  $k$ -nearest neighbour. There is no genuine rule to identify  $v$  or  $k$ . Consequently, the REBMIX algorithm identifies them from the set K of input values by minimizing the information criterion. The Sturges rule  $v = 1 + \log_2(n)$ , Log<sub>10</sub> rule  $v = 10\log_{10}(n)$  or RootN rule  $v = 2\sqrt{n}$  can be applied to estimate the limiting numbers of bins or the rule of thumb  $k = \sqrt{n}$  to guess the intermediate number of nearest neighbours. If, e.g., K = c(10, 20, 40, 60) and minimum IC coincides, e.g., 40, brackets are set to 20 and 60 and the golden section is applied to refine the minimum search. See also `kseq` for sequence of bins or nearest neighbours generation. The default value is "auto".

**ymin:** a vector of length  $d$  containing minimum observations. The default value is `numeric()`.

**ymax:** a vector of length  $d$  containing maximum observations. The default value is `numeric()`.

**ar:** acceleration rate  $0 < a_r \leq 1$ . The default value is 0.1 and in most cases does not have to be altered.

**Restraints:** a character giving the restraints type. One of "rigid" or default "loose". The rigid restraints are obsolete and applicable for well separated components only.

**Mode:** a character giving the mode type. One of "all", "outliers" or default "outliersplus". The modes are determined in decreasing order of magnitude from all observations if Mode = "all". If Mode = "outliers", the modes are determined in decreasing order of magnitude from outliers only. In the meantime, some outliers are reclassified as inliers. Finally, when all observations are inliers, the procedure is completed. If Mode = "outliersplus", the modes are determined in decreasing magnitude from the outliers only. In the meantime, some outliers are reclassified as inliers. Finally, if all observations are inliers, they are converted to outliers and the mode determination procedure is continued.

**w:** a list of vectors of length  $c$  containing component weights  $w_l$  summing to 1.

**Theta:** a list of lists each containing  $c$  parametric family types pdf1. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or circular

"vonMises" defined for  $0 \leq y_i \leq 2\pi$ . Component parameters  $\theta_{1l}$  follow the parametric family types. One of  $\mu_{il}$  for normal, lognormal, Gumbel and von Mises distributions,  $\theta_{il}$  for Weibull, gamma, binomial, Poisson and Dirac distributions and  $a$  for uniform distribution. Component parameters  $\theta_{2l}$  follow  $\theta_{1l}$ . One of  $\sigma_{il}$  for normal, lognormal and Gumbel distributions,  $\beta_{il}$  for Weibull and gamma distributions,  $p_{il}$  for binomial distribution,  $\kappa_{il}$  for von Mises distribution and  $b$  for uniform distribution. Component parameters  $\theta_{3l}$  follow  $\theta_{2l}$ . One of  $\xi_{il}$  for Gumbel distribution.

**summary:** a data frame with additional information about dataset, preprocessing,  $c_{\max}$ ,  $c_{\min}$ , information criterion type,  $a_r$ , restraints type, mode type, optimal  $c$ , optimal  $v$  or  $k$ ,  $K$ ,  $y_{i0}$ ,  $y_{\min}$ ,  $y_{\max}$ , optimal  $h_i$ , information criterion IC, log likelihood  $\log L$  and degrees of freedom  $M$ .

**summary.EM:** a data frame with additional information about dataset, strategy for the EM algorithm strategy, variant of the EM algorithm variant, acceleration type acceleration, tolerance tolerance, acceleration multiplier acceleration.multiplier, maximum allowed number of iterations maximum.iterations, number of iterations used for obtaining optimal solution opt.iterations.nbr and total number of iterations of the EM algorithm total.iterations.nbr.

**pos:** position in the summary data frame at which log likelihood  $\log L$  attains its maximum.

**opt.c:** a list of vectors containing numbers of components for optimal  $v$  for the histogram and the kernel density estimation or for optimal number of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**opt.IC:** a list of vectors containing information criteria for optimal  $v$  for the histogram and the kernel density estimation or for optimal number of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**opt.logL:** a list of vectors containing log likelihoods for optimal  $v$  for the histogram and the kernel density estimation or for optimal number of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**opt.Dmin:** a list of vectors containing  $D_{\min}$  values for optimal  $v$  for the histogram and the kernel density estimation or for optimal number of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**opt.D:** a list of vectors containing totals of positive relative deviations for optimal  $v$  for the histogram and the kernel density estimation or for optimal number of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**all.K:** a list of vectors containing all processed numbers of bins  $v$  for the histogram and the kernel density estimation or all processed numbers of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

**all.IC:** a list of vectors containing information criteria for all processed numbers of bins  $v$  for the histogram and the kernel density estimation or for all processed numbers of nearest neighbours  $k$  for the  $k$ -nearest neighbour.

## Author(s)

Marko Nagode

## Description

Returns as default the REBMIX algorithm output for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities. If model equals "REBMVNORM" output for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices is returned.

## Usage

```
## S4 method for signature 'REBMIX'
REBMIX(model = "REBMIX", Dataset = list(), Preprocessing = character(),
        cmax = 15, cmin = 1, Criterion = "AIC", pdf = character(),
        theta1 = numeric(), theta2 = numeric(), theta3 = numeric(), K = "auto",
        ymin = numeric(), ymax = numeric(), ar = 0.1,
        Restraints = "loose", Mode = "outliersplus", EMcontrol = NULL, ...)
## ... and for other signatures
## S4 method for signature 'REBMIX'
summary(object, ...)
## ... and for other signatures
```

## Arguments

<b>model</b>	see Methods section below.
<b>Dataset</b>	a list of length $n_D$ of data frames or objects of class <code>Histogram</code> . Data frames should have size $n \times d$ containing $d$ -dimensional datasets. Each of the $d$ columns represents one random variable. Numbers of observations $n$ equal the number of rows in the datasets.
<b>Preprocessing</b>	a character giving the preprocessing type. One of "histogram", "kernel density estimation" or "k-nearest neighbour".
<b>cmax</b>	maximum number of components $c_{\max} > 0$ . The default value is 15.
<b>cmin</b>	minimum number of components $c_{\min} > 0$ . The default value is 1. If $c_{\min} > 1$ , it may happen that no solution is found, and an error is returned by the method.
<b>Criterion</b>	a character giving the information criterion type. One of default Akaike "AIC", "AIC3", "AIC4" or "AICc", Bayesian "BIC", consistent Akaike "CAIC", Hannan-Quinn "HQC", minimum description length "MDL2" or "MDL5", approximate weight of evidence "AWE", classification likelihood "CLC", integrated classification likelihood "ICL" or "ICL-BIC", partition coefficient "PC", total of positive relative deviations "D" or sum of squares error "SSE".
<b>pdf</b>	a character vector of length $d$ containing continuous or discrete parametric family types. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or "vonMises".

theta1	a vector of length $d$ containing initial component parameters. One of $n_{il} =$ number of categories – 1 for "binomial" distribution.
theta2	a vector of length $d$ containing initial component parameters. Currently not used.
theta3	a vector of length $d$ containing initial component parameters. One of $\xi_{il} \in \{-1, \text{NA}, 1\}$ for "Gumbel" distribution.
K	a character or a vector or a matrix of size $n_D \times d$ containing numbers of bins $v$ or $v_1, \dots, v_d$ for the histogram and the kernel density estimation or numbers of nearest neighbours $k$ for the $k$ -nearest neighbour. There is no genuine rule to identify $v$ or $k$ . Consequently, the REBMIX algorithm identifies them from the set K of input values by minimizing the information criterion. The Sturges rule $v = 1 + \log_2(n)$ , Log <sub>10</sub> rule $v = 10\log_{10}(n)$ or RootN rule $v = 2\sqrt{n}$ can be applied to estimate the limiting numbers of bins or the rule of thumb $k = \sqrt{n}$ to guess the intermediate number of nearest neighbours. If, e.g., K = c(10, 20, 40, 60) and minimum IC coincides, e.g., 40, brackets are set to 20 and 60 and the golden section is applied to refine the minimum search. If, e.g., K = matrix(c(10, 15, 18, 5, 7, 9), byrow = TRUE, ncol = 3) than $d = 3$ and the list Dataset contains $n_D = 2$ frames. Hence, different numbers of bins can be assigned to $y_1, \dots, y_d$ . See also kseq for sequence of bins or nearest neighbours generation. The default value is "auto".
ymin	a vector of length $d$ containing minimum observations. The default value is numeric().
ymax	a vector of length $d$ containing maximum observations. The default value is numeric().
ar	acceleration rate $0 < a_r \leq 1$ . The default value is 0.1 and in most cases does not have to be altered.
Restraints	a character giving the restraints type. One of "rigid" or default "loose". The rigid restraints are obsolete and applicable for well separated components only.
Mode	a character giving the mode type. One of "all", "outliers" or default "outliersplus". The modes are determined in decreasing order of magnitude from all observations if Mode = "all". If Mode = "outliers", the modes are determined in decreasing order of magnitude from outliers only. In the meantime, some outliers are reclassified as inliers. Finally, when all observations are inliers, the procedure is completed. If Mode = "outliersplus", the modes are determined in decreasing magnitude from the outliers only. In the meantime, some outliers are reclassified as inliers. Finally, if all observations are inliers, they are converted to outliers and the mode determination procedure is continued.
EMcontrol	an object of class EM.Control.
object	see Methods section below.
...	currently not used.

**Value**

Returns an object of class REBMIX or REBMVNORM.

## Methods

```
signature(model = "REBMIX") a character giving the default class name "REBMIX" for mixtures
of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Pois-
son, Dirac, uniform or von Mises component densities.

signature(model = "REBMVNORM") a character giving the class name "REBMVNORM" for mixtures
of multivariate normal component densities with unrestricted variance-covariance matrices.

signature(object = "REBMIX") an object of class REBMIX.

signature(object = "REBMVNORM") an object of class REBMVNORM.
```

## Author(s)

Marko Nagode

## References

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## Examples

```
# Generate and plot univariate normal dataset.

n <- c(998, 263, 1086, 487)

Theta <- new("RNGMIX.Theta", c = 4, pdf = "normal")

a.theta1(Theta) <- c(688, 265, 30, 934)
a.theta2(Theta) <- c(72, 54, 34, 28)

normal <- RNGMIX(Dataset.name = "complex1",
rseed = -1,
n = n,
Theta = a.Theta(Theta))

normal

a.Dataset(normal, 1)[1:20,]

# Estimate number of components, component weights and component parameters.

normalest <- REBMIX(Dataset = a.Dataset(normal),
Preprocessing = "h",
cmax = 8,
Criterion = "BIC",
pdf = "n")

normalest

BIC(normalest)

logL(normalest)

# Plot finite mixture.

plot(normalest, nrow = 2, what = c("pdf", "marginal cdf"), npts = 1000)

# EM algorithm utilization

# Load iris data.

data(iris)

Dataset <- list(data.frame(iris[, c(1:4)]))

# Create EM.Control object.

EM <- new("EM.Control",
```

```

strategy = "exhaustive",
variant = "EM",
acceleration = "fixed",
tolerance = 1e-4,
acceleration.multiplier = 1.0,
maximum.iterations = 1000)

# Mixture parameter estimation using REBMIX and EM algorithm.

irisest <- REBMIX(model = "REBMVNORM",
Dataset = Dataset,
Preprocessing = "histogram",
cmax = 10,
Criterion = "BIC",
EMcontrol = EM)

irisest

# Print total number of EM iterations used in Ehxaustive strategy from summary.EM slot.

a.summary.EM(irisest, col.name = "total.iterations.nbr", pos = 1)

```

REBMIX.boot-class      *Class "REBMIX.boot"*

## Description

Object of class REBMIX.boot.

## Objects from the Class

Objects can be created by calls of the form `new("REBMIX.boot", ...)`. Accessor methods for the slots are `a.rseed(x = NULL)`, `a.pos(x = NULL)`, `a.Bootstrap(x = NULL)`, `a.B(x = NULL)`, `a.n(x = NULL)`, `a.replace(x = NULL)`, `a.prob(x = NULL)`, `a.c(x = NULL)`, `a.c.se(x = NULL)`, `a.c.cv(x = NULL)`, `a.c.mode(x = NULL)`, `a.c.prob(x = NULL)`, `a.w(x = NULL)`, `a.w.se(x = NULL)`, `a.w.cv(x = NULL)`, `a.Theta(x = NULL)`, `a.Theta.se(x = NULL)` and `a.Theta.cv(x = NULL)`, where `x` stands for an object of class REBMIX.boot.

## Slots

**x:** an object of class REBMIX.

**rseed:** set the random seed to any negative integer value to initialize the sequence. The first bootstrap dataset corresponds to it. For each next bootstrap dataset the random seed is decremented  $r_{\text{seed}} = r_{\text{seed}} - 1$ . The default value is -1.

**pos:** a desired row number in `x@summary` to be bootstrapped. The default value is 1.

**Bootstrap:** a character giving the bootstrap type. One of default "parametric" or "nonparametric".

**B:** number of bootstrap datasets. The default value is 100.

**n:** number of observations. The default value is `numeric()`.

**replace:** logical. The sampling is with replacement if TRUE, see also [sample](#). The default value is TRUE.

**prob:** a vector of length  $n$  containing probability weights, see also [sample](#). The default value is [numeric\(\)](#).

**c:** a vector containing numbers of components for  $B$  bootstrap datasets.

**c.se:** standard error of numbers of components c.

**c.cv:** coefficient of variation of numbers of components c.

**c.mode:** mode of numbers of components c.

**c.prob:** probability of mode c.mode.

**w:** a matrix containing component weights for  $\leq B$  bootstrap datasets.

**w.se:** a vector containing standard errors of component weights w.

**w.cv:** a vector containing coefficients of variation of component weights w.

**Theta:** a list of matrices containing component parameters theta1.1, theta2.1 and theta3.1 for  $\leq B$  bootstrap datasets.

**Theta.se:** a list of vectors containing standard errors of component parameters theta1.1, theta2.1 and theta3.1.

**Theta.cv:** a list of vectors containing coefficients of variation of component parameters theta1.1, theta2.1 and theta3.1.

## Author(s)

Marko Nagode

RNGMIX-class

Class "RNGMIX"

## Description

Object of class RNGMIX.

## Objects from the Class

Objects can be created by calls of the form `new("RNGMIX", ...)`. Accessor methods for the slots are `a.Dataset.name(x = NULL)`, `a.rseed(x = NULL)`, `a.n(x = NULL)`, `a.Theta(x = NULL)`, `a.Dataset(x = NULL, pos = 0)`, `a.Zt(x = NULL)`, `a.w(x = NULL)`, `a.Variables(x = NULL)`, `a.ymin(x = NULL)` and `a.ymax(x = NULL)`, where x and pos stand for an object of class RNGMIX and a desired slot item, respectively.

## Slots

- Dataset.name:** a character vector containing list names of data frames of size  $n \times d$  that  $d$ -dimensional datasets are written in.
- rseed:** set the random seed to any negative integer value to initialize the sequence. The first file in `Dataset.name` corresponds to it. For each next file the random seed is decremented  $r_{\text{seed}} = r_{\text{seed}} - 1$ . The default value is  $-1$ .
- n:** a vector containing numbers of observations in classes  $n_l$ , where number of observations  $n = \sum_{l=1}^c n_l$ .
- Theta:** a list containing  $c$  parametric family types `pdf1`. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or circular "vonMises" defined for  $0 \leq y_i \leq 2\pi$ . Component parameters  $\theta_{il}$  follow the parametric family types. One of  $\mu_{il}$  for normal, lognormal, Gumbel and von Mises distributions,  $\theta_{il}$  for Weibull, gamma, binomial, Poisson and Dirac distributions and  $a$  for uniform distribution. Component parameters  $\theta_{2.1}$  follow  $\theta_{1.1}$ . One of  $\sigma_{il}$  for normal, lognormal and Gumbel distributions,  $\beta_{il}$  for Weibull and gamma distributions,  $p_{il}$  for binomial distribution,  $\kappa_{il}$  for von Mises distribution and  $b$  for uniform distribution. Component parameters  $\theta_{3.1}$  follow  $\theta_{2.1}$ . One of  $\xi_{il} \in \{-1, 1\}$  for Gumbel distribution.
- Dataset:** a list of length  $n_D$  of data frames of size  $n \times d$  containing  $d$ -dimensional datasets. Each of the  $d$  columns represents one random variable. Numbers of observations  $n$  equal the number of rows in the datasets.
- Zt:** a factor of true cluster membership.
- w:** a vector of length  $c$  containing component weights  $w_l$  summing to 1.
- Variables:** a character vector containing types of variables. One of "continuous" or "discrete".
- ymin:** a vector of length  $d$  containing minimum observations.
- ymax:** a vector of length  $d$  containing maximum observations.

## Author(s)

Marko Nagode

## Description

Returns as default the RNGMIX univariate or multivariate random datasets for mixtures of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Poisson, Dirac, uniform or von Mises component densities. If `model` equals "RNGMVNORM" multivariate random datasets for mixtures of multivariate normal component densities with unrestricted variance-covariance matrices are returned.

## Usage

```
## S4 method for signature 'RNGMIX'
RNGMIX(model = "RNGMIX", Dataset.name = character(),
        rseed = -1, n = numeric(), Theta = list(), ...)
## ... and for other signatures
```

## Arguments

model	see Methods section below.
Dataset.name	a character vector containing list names of data frames of size $n \times d$ that $d$ -dimensional datasets are written in.
rseed	set the random seed to any negative integer value to initialize the sequence. The first file in Dataset.name corresponds to it. For each next file the random seed is decremented $r_{\text{seed}} = r_{\text{seed}} - 1$ . The default value is -1.
n	a vector containing numbers of observations in classes $n_l$ , where number of observations $n = \sum_{l=1}^c n_l$ .
Theta	a list containing $c$ parametric family types pdfl. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or circular "vonMises" defined for $0 \leq y_i \leq 2\pi$ . Component parameters theta1.1 follow the parametric family types. One of $\mu_{il}$ for normal, lognormal, Gumbel and von Mises distributions, $\theta_{il}$ for Weibull, gamma, binomial, Poisson and Dirac distributions and $a$ for uniform distribution. Component parameters theta2.1 follow theta1.1. One of $\sigma_{il}$ for normal, lognormal and Gumbel distributions, $\beta_{il}$ for Weibull and gamma distributions, $p_{il}$ for binomial distribution, $\kappa_{il}$ for von Mises distribution and $b$ for uniform distribution. Component parameters theta3.1 follow theta2.1. One of $\xi_{il} \in \{-1, 1\}$ for Gumbel distribution.
...	currently not used.

## Details

RNGMIX is based on the "Minimal" random number generator ran1 of Park and Miller with the Bays-Durham shuffle and added safeguards that returns a uniform random deviate between 0.0 and 1.0 (exclusive of the endpoint values).

## Value

Returns an object of class RNGMIX or RNGMVNORM.

## Methods

```
signature(model = "RNGMIX") a character giving the default class name "RNGMIX" for mixtures
of conditionally independent normal, lognormal, Weibull, gamma, Gumbel, binomial, Pois-
son, Dirac, uniform or von Mises component densities.

signature(model = "RNGMVNORM") a character giving the class name "RNGMVNORM" for mixtures
of multivariate normal component densities with unrestricted variance-covariance matrices.
```

## Author(s)

Marko Nagode

## References

W. H. Press, S. A. Teukolsky, W. T. Vetterling and B. P. Flannery. Numerical Recipes in C: The Art of Scientific Computing. Cambridge University Press, Cambridge, 1992.

## Examples

```
devAskNewPage(ask = TRUE)

# Generate and print multivariate normal datasets with diagonal
# variance-covariance matrices.

n <- c(75, 100, 125, 150, 175)

Theta <- new("RNGMIX.Theta", c = 5, pdf = rep("normal", 4))

a.theta1(Theta, 1) <- c(10, 12, 10, 12)
a.theta1(Theta, 2) <- c(8.5, 10.5, 8.5, 10.5)
a.theta1(Theta, 3) <- c(12, 14, 12, 14)
a.theta1(Theta, 4) <- c(13, 15, 7, 9)
a.theta1(Theta, 5) <- c(7, 9, 13, 15)
a.theta2(Theta, 1) <- c(1, 1, 1, 1)
a.theta2(Theta, 2) <- c(1, 1, 1, 1)
a.theta2(Theta, 3) <- c(1, 1, 1, 1)
a.theta2(Theta, 4) <- c(2, 2, 2, 2)
a.theta2(Theta, 5) <- c(3, 3, 3, 3)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1:25, sep = ""),
  rseed = -1,
  n = n,
  Theta = a.Theta(Theta))

simulated

plot(simulated, pos = 22, nrow = 2, ncol = 3)

# Generate and print multivariate normal datasets with unrestricted
# variance-covariance matrices.

n <- c(200, 50, 50)

Theta <- new("RNGMVNORM.Theta", c = 3, d = 3)

a.theta1(Theta, 1) <- c(0, 0, 0)
a.theta1(Theta, 2) <- c(-6, 3, 6)
a.theta1(Theta, 3) <- c(6, 6, 4)
a.theta2(Theta, 1) <- c(9, 0, 0, 0, 4, 0, 0, 0, 1)
a.theta2(Theta, 2) <- c(4, -3.2, -0.2, -3.2, 4, 0, -0.2, 0, 1)
a.theta2(Theta, 3) <- c(4, 3.2, 2.8, 3.2, 4, 2.4, 2.8, 2.4, 2)
```

```

simulated <- RNGMIX(model = "RNGMVNORM",
Dataset.name = paste("simulated_", 1:2, sep = ""),
rseed = -1,
n = n,
Theta = a.Theta(Theta))

simulated

plot(simulated, pos = 2, nrow = 3, ncol = 1)

# Generate and print multivariate mixed continuous-discrete datasets.

n <- c(400, 100, 500)

Theta <- new("RNGMIX.Theta", c = 3, pdf = c("lognormal", "Poisson", "binomial", "Weibull"))

a.theta1(Theta, 1) <- c(1, 2, 10, 2)
a.theta1(Theta, 2) <- c(3.5, 10, 10, 10)
a.theta1(Theta, 3) <- c(2.5, 15, 10, 25)
a.theta2(Theta, 1) <- c(0.3, NA, 0.9, 3)
a.theta2(Theta, 2) <- c(0.2, NA, 0.1, 7)
a.theta2(Theta, 3) <- c(0.4, NA, 0.7, 20)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1:5, sep = ""),
rseed = -1,
n = n,
Theta = a.Theta(Theta))

simulated

plot(simulated, pos = 4, nrow = 2, ncol = 3)

# Generate and print univariate mixed Weibull dataset.

n <- c(75, 100, 125, 150, 175)

Theta <- new("RNGMIX.Theta", c = 5, pdf = "Weibull")

a.theta1(Theta) <- c(12, 10, 14, 15, 9)
a.theta2(Theta) <- c(2, 4.1, 3.2, 7.1, 5.3)

simulated <- RNGMIX(Dataset.name = "simulated",
rseed = -1,
n = n,
Theta = a.Theta(Theta))

simulated

plot(simulated, pos = 1)

# Generate and print multivariate normal datasets with unrestricted
# variance-covariance matrices.

```

```

# Set dimension, dataset size, number of components and seed.

d <- 2; n <- 1000; c <- 10; set.seed(123)

# Component weights are generated.

w <- runif(c, 0.1, 0.9); w <- w / sum(w)

# Set range of means and rang of eigenvalues.

mu <- c(-100, 100); lambda <- c(1, 100)

# Component means and variance-covariance matrices are calculated.

Mu <- list(); Sigma <- list()

for (l in 1:c) {
  Mu[[l]] <- runif(d, mu[1], mu[2])
  Lambda <- diag(runif(d, lambda[1], lambda[2])), nrow = d, ncol = d)
  P <- svd(matrix(runif(d * d, -1, 1), nc = d))$u
  Sigma[[l]] <- P
}

# Numbers of observations are calculated and component means and
# variance-covariance matrices are stored.

n <- round(w * n); Theta <- list()

for (l in 1:c) {
  Theta[[paste0("pdf", l)]] <- rep("normal", d)
  Theta[[paste0("theta1.", l)]] <- Mu[[l]]
  Theta[[paste0("theta2.", l)]] <- as.vector(Sigma[[l]])
}

# Dataset is generated.

simulated <- RNGMIX(model = "RNGMVNORM", Dataset.name = "mvnorm_1",
rseed = -1, n = n, Theta = Theta)

plot(simulated)

# Generate and print bivariate mixed uniform-Gumbel dataset.

n <- c(100, 150)

Theta <- new("RNGMIX.Theta", c = 2, pdf = c("uniform", "Gumbel"))

a.theta1(Theta, l = 1) <- c(2, 10)
a.theta2(Theta, l = 1) <- c(10, 2.3)
a.theta3(Theta, l = 1) <- c(NA, 1.0)
a.theta1(Theta, l = 2) <- c(10, 50)
a.theta2(Theta, l = 2) <- c(30, 4.2)

```

```
a.theta3(Theta, l = 2) <- c(NA, -1.0)

simulated <- RNGMIX(Dataset.name = paste("simulated_", 1, sep = ""),
rseed = -1,
n = n,
Theta = a.Theta(Theta))

plot(simulated)
```

RNGMIX.Theta-class      *Class "RNGMIX.Theta"*

## Description

Object of class RNGMIX.Theta.

## Objects from the Class

Objects can be created by calls of the form `new("RNGMIX.Theta", ...)`. Accessor methods for the slots are `a.c(x = NULL)`, `a.d(x = NULL)`, `a.pdf(x = NULL)` and `a.Theta(x = NULL)`, where `x` stands for an object of class RNGMIX.Theta. Setter methods `a.theta1(x = NULL, l = numeric())`, `a.theta2(x = NULL, l = numeric())` and `a.theta3(x = NULL, l = numeric())`, `a.theta1.all(x = NULL)`, `a.theta2.all(x = NULL)` and `a.theta3.all(x = NULL)` are provided to write to Theta slot, where  $l = 1, \dots, c$ .

## Slots

**c:** number of components  $c > 0$ . The default value is 1.

**d:** number of dimensions.

**pdf:** a character vector of length  $d$  containing continuous or discrete parametric family types. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or "vonMises".

**Theta:** a list containing  $c$  parametric family types `pdfl`. One of "normal", "lognormal", "Weibull", "gamma", "Gumbel", "binomial", "Poisson", "Dirac", "uniform" or circular "vonMises" defined for  $0 \leq y_i \leq 2\pi$ . Component parameters `theta1.l` follow the parametric family types. One of  $\mu_{il}$  for normal, lognormal, Gumbel and von Mises distributions,  $\theta_{il}$  for Weibull, gamma, binomial, Poisson and Dirac distributions and  $a$  for uniform distribution. Component parameters `theta2.l` follow `theta1.l`. One of  $\sigma_{il}$  for normal, lognormal and Gumbel distributions,  $\beta_{il}$  for Weibull and gamma distributions,  $p_{il}$  for binomial distribution,  $\kappa_{il}$  for von Mises distribution and  $b$  for uniform distribution. Component parameters `theta3.l` follow `theta2.l`. One of  $\xi_{il} \in \{-1, 1\}$  for Gumbel distribution.

## Author(s)

Marko Nagode

## Examples

```

Theta <- new("RNGMIX.Theta", c = 2, pdf = c("normal", "Gumbel"))

a.theta1(Theta, l = 1) <- c(2, 10)
a.theta2(Theta, l = 1) <- c(0.5, 2.3)
a.theta3(Theta, l = 1) <- c(NA, 1.0)
a.theta1(Theta, l = 2) <- c(20, 50)
a.theta2(Theta, l = 2) <- c(3, 4.2)
a.theta3(Theta, l = 2) <- c(NA, -1.0)

Theta

Theta <- new("RNGMIX.Theta", c = 2, pdf = c("normal", "Gumbel"))

a.theta1.all(Theta) <- c(2, 10, 20, 50)
a.theta2.all(Theta) <- c(0.5, 2.3, 3, 4.2)
a.theta3.all(Theta) <- c(NA, 1.0, NA, -1.0)

Theta

Theta <- new("RNGMVNORM.Theta", c = 2, d = 3)

a.theta1(Theta, l = 1) <- c(2, 10, -20)
a.theta1(Theta, l = 2) <- c(-2.4, -15.1, 30)

Theta

```

## Description

These data are the results of a sensorless drive diagnosis procedure. Features are extracted from the electric current drive signals. The drive has intact and defective components. This results in 11 different classes with different conditions. Each condition has been measured several times by 12 different operating conditions, this means by different speeds, load moments and load forces. The current signals are measured with a current probe and an oscilloscope on two phases. The original dataset contains 49 features, however, here only 3 are used, that is, features 5, 7 and 11. First class (1) are the healthy drives and the rest are the drives with fault components.

## Usage

```
data(sensorlessdrive)
```

## Format

`sensorlessdrive` is a data frame with 58509 cases (rows) and 4 variables (columns) named:

1. V5 continuous.

2. V7 continuous.
3. V11 continuous.
4. Class discrete 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or 11.

### Source

A. Asuncion and D. J. Newman. Uci machine learning repository, 2007. <http://archive.ics.uci.edu/ml/>.

### References

F. Paschke1, C. Bayer, M. Bator, U. Moenks, A. Dicks, O. Enge-Rosenblatt and V. Lohweg. Sensorlose Zustandsueberwachung an Synchronmotoren. 23. Workshop Computational Intelligence VDI/VDE-Gesellschaft Mess- und Automatisierungstechnik (GMA), 2013.

M. Bator, A. Dicks, U. Moenks and V. Lohweg. Feature extraction and reduction applied to sensorless drive diagnosis. 22. Workshop Computational Intelligence VDI/VDE-Gesellschaft Mess- und Automatisierungstechnik (GMA), 2012. doi:10.13140/2.1.2421.5689.

### Examples

```
## Not run:
data(sensorlessdrive)

# Split dataset into train (75

set.seed(3)

Drive <- split(p = 0.75, Dataset = sensorlessdrive, class = 4)

# Estimate number of components, component weights and component
# parameters for train subsets.

driveest <- REBMIX(model = "REBMVNORM",
Dataset = a.train(Drive),
Preprocessing = "histogram",
cmax = 15,
Criterion = "BIC")

# Classification.

drivecla <- RCLSMIX(model = "RCLSMVNORM",
x = list(driveest),
Dataset = a.test(Drive),
Zt = a.Zt(Drive))

drivecla

summary(drivecla)

## End(Not run)
```

**split-methods***Splits Dataset into Train and Test Datasets***Description**

Returns (invisibly) the object containing train and test observations  $y_1, \dots, y_n$  as well as true class membership  $\Omega_g$  for the test dataset.

**Usage**

```
## S4 method for signature 'numeric'
split(p = 0.75, Dataset = data.frame(), class = numeric(), ...)
## S4 method for signature 'list'
split(p = list(), Dataset = data.frame(), class = numeric(), ...)
## ... and for other signatures
```

**Arguments**

- |                      |  |
|----------------------|--|
| <code>p</code>       | see Methods section below.   |
| <code>Dataset</code> | a data frame containing dataset $Y$ of length $n$ . For the dataset the corresponding class membership $\Omega_g$ is known. The default value is <code>data.frame()</code> . |
| <code>class</code>   | a column number in <code>Dataset</code> containing the class membership information. The default value is <code>numeric()</code> .   |
| <code>...</code>     | further arguments to <code>sample</code> .   |

**Value**

Returns an object of class `RCLS.chunk`.

**Methods**

`signature(p = "numeric")` a number specifying the fraction of observations for training  $0.0 \leq p \leq 1.0$ . The default value is  $0.75$ .

`signature(p = "list")` a list composed of column number `p$type` in `Dataset` containing the type membership information followed by the corresponding train `p$train` and test `p$test` values. The default value is `list()`.

**Author(s)**

Marko Nagode

**Examples**

```
## Not run:  
data(iris)  
  
# Split dataset into train (75  
  
set.seed(5)  
  
Iris <- split(p = 0.75, Dataset = iris, class = 5)  
  
Iris  
  
# Generate simulated dataset.  
  
N <- 1000  
  
class <- c(rep("A", 0.4 * N), rep("B", 0.2 * N),  
rep("C", 0.1 * N), rep("D", 0.05 * N), rep("E", 0.25 * N))  
  
type <- c(rep("train", 0.75 * N), rep("test", 0.25 * N))  
  
n <- 300  
  
Dataset <- data.frame(1:n, sample(class, n))  
  
colnames(Dataset) <- c("y", "class")  
  
# Split dataset into train (60  
  
simulated <- split(p = 0.6, Dataset = Dataset, class = 2)  
  
simulated  
  
# Generate simulated dataset.  
  
Dataset <- data.frame(1:n, sample(class, n), sample(type, n))  
  
colnames(Dataset) <- c("y", "class", "type")  
  
# Split dataset into train and test subsets.  
  
simulated <- split(p = list(type = 3, train = "train",  
test = "test"), Dataset = Dataset, class = 2)  
  
simulated  
  
## End(Not run)
```

## Description

Returns the sum of squares error at pos.

## Usage

```
## S4 method for signature 'REBMIX'
SSE(x = NULL, pos = 1, ...)
## ... and for other signatures
```

## Arguments

- x see Methods section below.
- pos a desired row number in x@summary for which the information criterion is calculated. The default value is 1.
- ... currently not used.

## Methods

- `signature(x = "REBMIX")` an object of class REBMIX.
- `signature(x = "REBMVNORM")` an object of class REBMVNORM.

## Author(s)

Marko Nagode

## References

- C. M. Bishop. Neural Networks for Pattern Recognition. Clarendon Press, Oxford, 1995.

## Description

These data are the results of an extraction process from images of faults of steel plates. There are seven different faults: Pastry (1), Z\_Scratch (2), K\_Scratch (3), Stains (4), Dirtiness (5), Bumps (6), Other faults (7).

## Usage

```
data(steelplates)
```

## Format

steelplates is a data frame with 1941 cases (rows) and 28 variables (columns) named:

1. X\_Minimum integer.
2. X\_Maximum integer.
3. Y\_Minimum integer.
4. Y\_Maximum integer.
5. Pixels\_Areas integer.
6. X\_Perimeter integer.
7. Y\_Perimeter integer.
8. Sum\_of\_Luminosity integer.
9. Minimum\_of\_Luminosity integer.
10. Maximum\_of\_Luminosity integer.
11. Length\_of\_Conveyer integer.
12. TypeOfSteel\_A300 binary.
13. TypeOfSteel\_A400 binary.
14. Steel\_Plate\_Thickness integer.
15. Edges\_Index continuous.
16. Empty\_Index continuous.
17. Square\_Index continuous.
18. Outside\_X\_Index continuous.
19. Edges\_X\_Index continuous.
20. Edges\_Y\_Index continuous.
21. Outside\_Global\_Index continuous.
22. LogOfAreas continuous.
23. Log\_X\_Index continuous.
24. Log\_Y\_Index continuous.
25. Orientation\_Index continuous.
26. Luminosity\_Index continuous.
27. SigmoidOfAreas continuous.
28. Class discrete 1, 2, 3, 4, 5, 6 or 7.

## Source

A. Asuncion and D. J. Newman. Uci machine learning repository, 2007. <http://archive.ics.uci.edu/ml/>.

## References

M. Buscema, S. Terzi, W. Tastle. A new meta-classifier. Annual Conference of the North American Fuzzy Information Processing Society - NAFIPS, 2010. doi:[10.1109/NAFIPS.2010.5548298](https://doi.org/10.1109/NAFIPS.2010.5548298).

M. Buscema. MetaNet\*: The theory of independent judges. Substance Use & Misuse. 33(2):439-461, 1998. doi:[10.3109/10826089809115875](https://doi.org/10.3109/10826089809115875).

## Examples

```
## Not run:
data(steelplates)

# Split dataset into train (75
set.seed(3)

Steelplates <- split(p = 0.75, Dataset = steelplates, class = 28)

# Estimate number of components, component weights and component
# parameters for train subsets.

steelplatesest <- REBMIX(model = "REBMVNORM",
  Dataset = a.train(Steelplates),
  Preprocessing = "histogram",
  cmax = 15,
  Criterion = "BIC")

# Classification.

steelplatescla <- RCLSMIX(model = "RCLSMVNORM",
  x = list(steelplatesest),
  Dataset = a.test(Steelplates),
  Zt = a.Zt(Steelplates))

steelplatescla

summary(steelplatescla)

## End(Not run)
```

**truck**

*Truck Dataset*

## Description

The dataset contains amplitudes and means measured on a truck wheels.

## Usage

```
data(truck)
```

**Format**

truck is a data frame with 31665 rows and 2 variables (columns) named:

1. Amplitude continuous.
2. Mean continuous.

**Author(s)**

Mitja Franko

**Examples**

```
data(truck)
```

---

weibull

*Weibull Dataset 8.I*

---

**Description**

The complete data are the failure times in weeks.

**Usage**

```
data(weibull)
```

**Format**

weibull is a data frame with 50 cases (rows) and 1 variables (columns) named:

1. Failure.Time continuous.

**References**

D. N. P. Murthy, M. Xie and R. Jiang. Weibull Models. John Wiley & Sons, New York, 2003.

**Examples**

```
data(weibull)
```

**weibullnormal***Weibull-normal Simulated Dataset***Description**

The dataset contains amplitudes and means simulated from a three component Weibull-normal mixture.

**Usage**

```
data(weibullnormal)
```

**Format**

`weibullnormal` is a data frame with 10000 rows and 2 variables (columns) named:

1. Amplitude continuous.
2. Mean continuous.

**Author(s)**

Mitja Franko

**Examples**

```
data(weibullnormal)
```

**wine***Wine Recognition Data***Description**

These data are the results of a chemical analysis of wines grown in the same region in Italy but derived from three different cultivars (1-3). The analysis determined the quantities of 13 constituents: alcohol, malic acid, ash, alcalinity of ash, magnesium, total phenols, flavanoids, nonflavanoid phenols, proanthocyanins, colour intensity, hue, OD280/OD315 of diluted wines, and proline found in each of the three types of the wines. The number of instances in classes 1 to 3 is 59, 71 and 48, respectively.

**Usage**

```
data(wine)
```

## Format

wine is a data frame with 178 cases (rows) and 14 variables (columns) named:

1. Alcohol continuous.
2. Malic.Acid continuous.
3. Ash continuous.
4. Alcalinity.of.Ash continuous.
5. Magnesium continuous.
6. Total.Phenols continuous.
7. Flavanoids continuous.
8. Nonflavanoid.Phenols continuous.
9. Proanthocyanins continuous.
10. Color.Intensity continuous.
11. Hue continuous.
12. OD280.OD315.of.Diluted.Wines continuous.
13. Proline continuous.
14. Cultivar discrete 1, 2 or 3.

## Source

A. Asuncion and D. J. Newman. Uci machine learning repository, 2007. <http://archive.ics.uci.edu/ml/>.

## References

S. J. Roberts, R. Everson and I. Rezek. Maximum certainty data partitioning. Pattern Recognition, 33(5):833-839, 2000. doi:10.1016/S00313203(99)000862.

## Examples

```
## Not run:  
devAskNewPage(ask = TRUE)  
  
data(wine)  
  
# Show level attributes.  
  
levels(factor(wine[["Cultivar"]]))  
  
# Split dataset into train (75  
  
set.seed(3)  
  
Wine <- split(p = 0.75, Dataset = wine, class = 14)  
  
# Estimate number of components, component weights and component
```

```

# parameters for train subsets.

n <- range(a.ntrain(Wine))

K <- c(as.integer(1 + log2(n[1])), # Minimum v follows Sturges rule.
       as.integer(10 * log10(n[2]))) # Maximum v follows log10 rule.

K <- c(floor(K[1]^(1/13)), ceiling(K[2]^(1/13)))

wineest <- REBMIX(model = "REBMVNORM",
                    Dataset = a.train(Wine),
                    Preprocessing = "kernel density estimation",
                    cmax = 10,
                    Criterion = "ICL-BIC",
                    pdf = rep("normal", 13),
                    K = K[1]:K[2],
                    Restraints = "loose",
                    Mode = "outliersplus")

plot(wineest, pos = 1, nrow = 7, ncol = 6, what = c("pdf"))
plot(wineest, pos = 2, nrow = 7, ncol = 6, what = c("pdf"))
plot(wineest, pos = 3, nrow = 7, ncol = 6, what = c("pdf"))

# Selected chunks.

winecla <- RCLSMIX(model = "RCLSMVNORM",
                     x = list(wineest),
                     Dataset = a.test(Wine),
                     Zt = a.Zt(Wine))

winecla

summary(winecla)

# Plot selected chunks.

plot(winecla, nrow = 7, ncol = 6)

## End(Not run)

```

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