Package ‘nor1mix’

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Title Normal aka Gaussian (1-d) Mixture Models (S3 Classes and Methods)

Version 1.3-0

Date 2019-06-13

Description Onedimensional Normal (i.e. Gaussian) Mixture Models Classes, for, e.g., density estimation or clustering algorithms research and teaching; providing the widely used Marron-Wand densities. Efficient random number generation and graphics. Fitting to data by efficient ML (Maximum Likelihood) or traditional EM estimation.

Imports stats, graphics

Suggests cluster

License GPL (>= 2)

Encoding UTF-8

NeedsCompilation no

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Details

The DESCRIPTION file:

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Title: Normal aka Gaussian (1-d) Mixture Models (S3 Classes and Methods)
Version: 1.3-0
Date: 2019-06-13
Authors@R: c(person("Martin", "Maechler", role = c("aut","cre"), email = "maechler@stat.math.ethz.ch", comment = c(ORCID = "0000-0001-7278-1983")) , person("Erik", "Jørgensen", role = "ctb", comment = "pnorMix(), qnorMix()")
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Maintainer: Martin Maechler <maechler@stat.math.ethz.ch>

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Note that direct Maximum Likelihood ML (via optim()) is typically much faster converging (and more reliably detecting convergence correctly), notably thanks to a smart re-parametrization: use norMixMLE().

Author(s)
NA
Maintainer: NA

See Also
The Marron-Wand examples of normal (gaussian) mixtures MarronWand.
Multivariate distributions from copulas Mvdc from the copula package can use norMix marginals.

Examples
example(dnorMix)

clus2norMix  Transform Clustering / Grouping to Normal Mixture

Description
Simple transformation of a clustering or grouping to a normal mixture object (class "norMix", see, norMix).

Usage
clus2norMix(gr, x, name = deparse(sys.call()))

Arguments
  gr  a grouping/clustering vector with values in \{1, ..., \(K\}\}; possibly a factor.
  x  numeric vector of (original) data (of the same length as gr).
  name  name for norMix() object; constructed from the call by default.
dnorMix

Value

A call to `norMix()` with (mu, sig2, w) set to the empirical values of the groups (as defined by `split(x, gr)`).

Note

Via this function, any simple clustering algorithm (such as `pam`) can be used as simple mixture model fitting procedure.

Author(s)

Martin Maechler, Dec. 2007

See Also

`norMix`; further `pam()` (or `clara()`) from package `cluster` for sensible clusterings.

Examples

```r
x9 <- rnorMix(500, MW.nm9)
require("cluster")
pxc <- pam(x9, k=3)
plot(pxc, which = 2)# silhouette

(nm.p9 <- clus2norMix(pxc$clustering, x9))
plot(nm.p9, p.norm=FALSE)
lines(MW.nm9, col="thistle")
```

---

dnorMix

**Normal Mixture Density**

Description

Evaluate the density function of the normal mixture specified as `norMix` object.

Usage

```r
dnorMix (x, obj, log = FALSE)
dnorMixL(obj, x = NULL, log = FALSE, xlim = NULL, n = 511)
dpnorMix(x, obj, lower.tail = TRUE)
```
Arguments

obj    an object of class norMix.

x    numeric vector with abscissa values where to evaluate the density (and probability, for dpnorMix()). For dnorMixL() by default, when NULL, it is constructed from n (and xlim if that is specified).

log    logical indicating log-density values should be returned.

xlim    range of abscissa values, used if x == NULL. By default, xlim is taken as mean plus/minus 3 standard deviations of the normal mixture.

n    number of abscissa values to generate if x is not specified.

lower.tail    logical; if TRUE (default), probabilities are $P[X \leq x]$, otherwise, $P[X > x]$.

Value

dnorMix(x) returns the numeric vector of density values $f(x)$, logged if log is TRUE.
dnorMixL() returns a list with components

x    the abscissa values.

y    the density values $f(x)$ as for dnorMix().

dpnorMix() returns a list with components

d    the density values $f(x)$ as for dnorMix().

p    the probability values $F(x)$ as for pnorMix().

See Also

rnorMix for random number generation, and norMix for the construction and further methods, particularly plot.norMix which makes use dnorMix.

Examples

ff <- dnorMixL(MW.nm7)
str(ff)
plot(ff, type = "h", ylim = c(0,1)) # rather use plot(ff, ...)

x <- seq(-4,5, length=501)
dp <- dpnorMix(x, MW.mm7)
lines(x, dp$d, col = "tomato", lwd=3)
lines(x, dp$p, col = 3, lwd=2)# does not fit y-wise
stopifnot(all.equal(dp$d, dnorMix(x, MW.mm7), tolerance=1e-12),
all.equal(dp$p, pnorMix(x, MW.mm7), tolerance=1e-12))
**llnorMix**

*Likelihood, Parametrization and EM-Steps For 1D Normal Mixtures*

**Description**

These functions work with an almost unconstrained parametrization of univariate normal mixtures.

- `llnorMix(p, *)` computes the log likelihood.
- `obj <- par2norMix(p)` maps parameter vector `p` to a `norMix` object `obj`.
- `p <- nM2par(obj)` maps from a `norMix` object `obj` to parameter vector `p`.

Where `p` is always a parameter vector in our parametrization.

Partly for didactical reasons, the following functions provide the basic ingredients for the EM algorithm (see also `norMixEM`) to parameter estimation:

- `estep.nm(x, obj, p)` computes 1 E-step for the data `x`, given either a "norMix" object `obj` or parameter vector `p`.
- `mstep.nm(x, z)` computes 1 M-step for the data `x` and the probability matrix `z`.
- `emstep.nm(x, obj)` computes 1 E- and 1 M-step for the data `x` and the "norMix" object `obj`.

Where again, `p` is a parameter vector in our parametrization, `x` is the (univariate) data, and `z` is a `$n \times m$ matrix` of (posterior) conditional probabilities, and `θ` is the full parameter vector of the mixture model.

**Usage**

```r
llnorMix(p, x, m = (length(p) + 1)/3, trafo = c("clr1", "logit"))
par2norMix(p, trafo = c("clr1", "logit"), name = )
nM2par(obj, trafo = c("clr1", "logit"))
estep.nm(x, obj, par)
mstep.nm(x, z)
emstep.nm(x, obj)
```

**Arguments**

- `p, par` numeric vector: our parametrization of a univariate normal mixture, see details.
- `x` numeric: the data for which the likelihood is to be computed.
- `m` integer number of mixture components; this is not to be changed for a given `p`.
- `trafo` character string specifying the transformation of the component weight `$w \cdot m$-vector` (mathematical notation in `norMix`: `$π_j, j = 1, \ldots, m$`) to an `$m - 1$-dimensional unconstrained parameter vector in our parametrization. "logit" has been hard-wired up to `norlimix` version 1.2-3, and has been replaced as default in 2019 for `norlimix` version 1.2-4 by "clr1" which is more symmetric and basically Aitchinson’s centered log ratio, see also CRAN package `compositions’s clr()`.
name (for par2norMix()): a name for the "norMix" object that is returned, uses a smart default.

obj a "norMix" object, see norMix.

z a \( n \times m \) matrix of (posterior) conditional probabilities, \( z_{ij} = P(x_i \in C_j|\theta) \), where \( C_j \) denotes the \( j \)-th group ("cluster").

Details

We use a parametrization of a (finite) \( m \)-component univariate normal mixture which is particularly apt for likelihood maximization, namely, one whose parameter space is almost a full \( \mathbb{R}^M \), \( M = 3m - 1 \).

For an \( m \)-component mixture, we map to and from a parameter vector \( \theta (== p \text{ as } \mathbb{R}\text{-vector}) \) of length \( 3m - 1 \). For mixture density

\[
\sum_{j=1}^{m} \pi_j \phi((t - \mu_j)/\sigma_j)/\sigma_j,
\]

we transform the \( \pi_j \) (for \( j \in 1, \ldots, m \)) via the transform specified by trafo (see below), and log-transform the \( \sigma_j \). Consequently, \( \theta \) is partitioned into

\[
p[1:(m-1)]: \quad \text{For}
\]

\[
\text{trafo = "logit": } p[j] = \log(\pi_{j+1}) \quad \text{and } \pi_1 \text{ is given implicitly as } \pi_1 = 1 - \sum_{j=2}^{m} \pi_j.
\]

\[
\text{trafo = "clr1": } (\text{centered log ratio, omitting 1st element): Set } \ell_j := \ln(\pi_j) \text{ for } j = 1, \ldots, m, \text{ and } p[j] = \ell_{j+1} - 1/m \sum_{j=1}^{m-1} \ell_j \text{ for } j = 1, \ldots, m-1.
\]

\[
p[m:(2m-1)]: \quad p[m-1+j] = \mu_j, \text{ for } j=1:m.
\]

\[
p[2m:(3m-1)]: \quad p[2*m-1+j] = \log(\sigma_j), \text{ i.e., } \sigma_j^2 = \exp(2 * p[. + j]).
\]

Value

llnorMix() returns a number, namely the log-likelihood.

par2norMix() returns "norMix" object, see norMix.

nM2par() returns the parameter vector \( \theta \) of length \( 3m - 1 \).

estep.nm() returns \( z \), the matrix of (conditional) probabilities.

mstep.nm() returns the model parameters as a list with components \( w, \mu, \) and \( \sigma \), corresponding to the arguments of norMix(). (and see the 'Examples' on using do.call(norMix, *) with it.)

emstep.nm() returns an updated "norMix" object.

Author(s)

Martin Maechler
See Also

`norMix`, `logLik`. Note that the log likelihood of a "norMix" object is directly given by `sum(dnorMix(x, obj, log=TRUE)).

To fit, using the EM algorithm, rather use `norMixEM()` than the `e.step`, `m.step`, or `em.step` functions.

Note that direct likelihood maximization, i.e., MLE, is typically considerably more efficient than
the EM, and typically converges well with our parametrization, see `norMixMLE`.

Examples

```r
(obj <- MW.nm10) # "the Claw" -- m = 6 components
length(pp <- nM2par(obj)) # 17 == (3*6) - 1
par2norMix(pp)
## really the same as the initial 'obj' above

## Log likelihood (of very artificial data):
llnorMix(pp, x = seq(-2, 2, length=1000))
set.seed(47)## of more realistic data:
x <- rnorMix(1000, obj)
llnorMix(pp, x)

## Consistency check : nM2par() and par2norMix() are inverses
all.EQ <- function(x,y, tol = 1e-15, ...) all.equal(x,y, tolerance=tol, ...)
stopifnot(all.EQ(pp, nM2par(par2norMix(pp))),
  all.EQ(obj, par2norMix(nM2par(obj))),
  check.attributes=FALSE),
## Direct computation of log-likelihood:
all.EQ(sum(dnorMix(x, obj, log=TRUE)),
      llnorMix(pp, x))

## E- and M- steps : ------------------------------
rE1 <- estep.nm(x, obj)
rE2 <- estep.nm(x, par=pp) # the same as rE1
z <- rE1
str( rM <- mstep.nm(x, z))
(rEM <- emstep.nm(x, obj))

stopifnot(all.EQ(rE1, rE2),
      all.EQ(rEM, do.call(norMix, c(rM, name="")))
```

Description

The fifteen density examples used in Marron and Wand (1992)’s simulation study have been used in quite a few subsequent studies, can all be written as normal mixtures and are provided here for convenience and didactical examples of normal mixtures. Number 16 has been added by Jansen et al.
Usage

MW.nm1 # Gaussian
MW.nm2 # Skewed
MW.nm2.old # Skewed(old)
MW.nm3 # Str Skew
MW.nm4 # Kurtotic
MW.nm5 # Outlier
MW.nm6 # Bimodal
MW.nm7 # Separated (bimodal)
MW.nm8 # Asymmetric Bimodal
MW.nm9 # Trimodal
MW.nm10 # Claw
MW.nm11 # Double Claw
MW.nm12 # Asymmetric Claw
MW.nm13 # Asymm. Double Claw
MW.nm14 # Smooth Comb
MW.nm15 # Discrete Comb
MW.nm16 # Distant Bimodal

Author(s)

Martin Maechler

Source

They have been translated from Steve Marron’s Matlab code, now at
https://marronwebfiles.sites.oasis.unc.edu/oldResearch/parameters/nmpar.m, however
for number 2, the Matlab code had MW.nm2.old; and I’ve defined MW.nm2 as from the Annals paper;
see also the last example below.

References

Marron, S. and Wand, M. (1992) Exact Mean Integrated Squared Error; *Annals of Statistics* 20,
712–736.

For number 16,

Examples

```r
MW.nm10
plot(MW.nm14)
```

## These are defined as norMix() calls in ../R/zMarrWand-dens.R

```r
nms <- ls(pat="^MW.nm", "package:nor1mix")
nms <- nms[order(as.numeric(substring(nms,6)))]
for(n in nms) {
```
norMix

Mixtures of Univariate Normal Distributions

Description

Objects of class `norMix` represent finite mixtures of (univariate) normal (aka Gaussian) distributions. Methods for construction, printing, plotting, and basic computations are provided.

Usage

```r
norMix(mu, sig2 = rep(1, m), sigma = rep(1, m),
```

```r
```

## Plot all of them:
op <- par(mfrow=c(4,4), mgp = c(1.2, 0.5, 0), tcl = -0.2,
       mar = .1 + c(2,2,2,1), oma = c(0,0,3,0))
for(n in nms[-17]) plot(get(n, "package:nor1mix"))

## and their Q-Q-plots (not really fast):
prob <- ppoints(N <- 100)
for(n in nms[-17])
qqnorm(qnorMix(prob, get(n, "package:nor1mix")), main = n)

## "object" overview:
cbind(sapply(nms, function(n) { o <- get(n)
       sprintf("%-18s: K =%2d; rng = [%3.1f, %2.1f],
                attr(o, "name"), nrow(o),
                min(o[,"mu"] - 3*sqrt(o[,"sig2"])),
                max(o[,"mu"] + 3*sqrt(o[,"sig2"])) )
       }))

## Note that Marron-Wand (1992), p.720 give #2 as
## the parameters of which at first look quite different from
## which has been the definition in the above "Source" Matlab code.
## It's easy to see that mu_{nm2} = -.3 + 1.2 * mu_{paper},
## and correspondingly, s2_{nm2} = 1.2^2 * s2_{paper}
## such that they are "identical" apart from scale and location:
op. <- par(mfrow=2:1, mgp= c(1.2, 0.5,0), tcl= -0.2, mar=.1+c(2,2,2,1))
plot(MW.nm2)
plot(MW.nm2.old)
par(op.)
is.norMix(obj)
m.norMix(obj)
var.norMix(x, ...)
## S3 method for class 'norMix'
mean(x, ...)
## S3 method for class 'norMix'
print(x, ...)
## S3 method for class 'norMix'

Arguments

mu
numeric vector of length \( K \), say, specifying the means \( \mu \) of the \( K \) normal components.

sig2
**deprecated!** numeric vector of length \( K \), specifying the variances \( \sigma^2 \) of the \( K \) normal components. Do specify sigma instead!

sigma
numeric vector of length \( K \), specifying the standard deviations \( \sigma \) of the \( K \) normal components.

w
numeric vector of length \( K \), specifying the mixture proportions \( \pi_j \) of the normal components, \( j = 1, \ldots, K \). Defaults to equal proportions

name
optional name tag of the result (used for printing).

long.name
logical indicating if the name attribute should use punctuation and hence be slightly larger than by default.

obj, x
an object of class norMix.

i, j, drop
for indexing, see the generic [ extractor function.

... further arguments passed to methods.

Details

The (one dimensional) normal mixtures, \( \mathbb{R} \) objects of class "norMix", are constructed by norMix and tested for by is.norMix. m.norMix() returns the number of mixture components; the mean() method for class "norMix" returns the (theoretical / true) mean \( E[X] \) and var.norMix() the true variance \( E[(X - E[X])^2] \) where \( X \sim \langle \text{norm.mxt} \rangle \).

The subsetting aka “extract” method (x[i,j]; for generic [])—when called as x[i,]—will typically return a "norMix" object unless matrix indexing selects only one row in which case x[i, , drop=FALSE] will return the normal mixture (of one component only).

For further methods (density, random number generation, fitting, ...), see below.

Value

norMix returns objects of class "norMix" which are currently implemented as 3-column matrix with column names mu, sigma, and w, and further attributes. The user should rarely need to access the underlying structure directly.
Note

For *estimation* of the parameters of such a normal mixture, we provide a smart parametrization and an efficient implementation of the direct MLE or also the EM algorithm, see `norMixMLE()` which includes `norMixEM()`.

Author(s)

Martin Maechler

See Also

dnorMix for the density, pnorMix for the cumulative distribution and the quantile function (qnorMix), and rnorMix for random numbers and plot.norMix, the plot method.

MarronWand has the Marron-Wand densities as normal mixtures.

`norMixMLE()` and `norMixEM()` provide *fitting* of univariate normal mixtures to data.

Examples

```r
ex <- norMix(mu = c(1,2,5)) # defaults: sigma = 1, equal proportions ('w')
ex
plot(ex, p.comp = TRUE)# looks like a mixture of only 2; 'p.comp' plots components

## The 2nd Marron-Wand example, see also ?MW.nm2
ex2 <- norMix(name = "#2 Skewed",
             mu = c(0,.5,13/12),
             sigma = c(1,2/3,5/9),
             w = c(.2,.2,.6))

m.norMix(ex2)
mean(ex2)
var.norMix(ex2)
(e23 <- ex2[2:3,]) # (with re-normalized weights)
stopifnot(is.norMix(e23),
          all.equal(var.norMix(ex2), 719/1080, tol=1e-14),
          all.equal(var.norMix(ex), 35/9, tol=1e-14),
          all.equal(var.norMix(ex[2:3,]), 13/4, tol=1e-14),
          all.equal(var.norMix(e23), 53^2/(12^3*4), tol=1e-14))

plot(ex2, log = "y")# maybe "revealing"
```

---

**norMix2call**

*Transform ”norMix” object into Call, Expression or Function*

Description

E.g., for taking symbolic derivatives, it may be useful to get an R **call**, **expression**, or **function** in / of x from a specific ”norMix” object.
norMix2call

Usage

norMix2call(obj, oneArg = TRUE)
## S3 method for class 'norMix'
as.expression(x, oneArg = TRUE, ...)
## S3 method for class 'norMix'
as.function(x, oneArg = TRUE, envir = parent.frame(), ...)

Arguments

obj, x  an R object of class "norMix".
oneArg  logical specifying if expressions of the form dnorm((x - mu)/sig) should be used, i.e. one Argument only, instead of dnorm(x, mu, sig).
envir   an environment; often the default is perfect.
...    potentially further arguments (not used in any examples yet).

Value

according to the function used, an R 'language' object, i.e., a call, expression, or function, respectively.

Author(s)

Martin Maechler

See Also

norMix. Note that deriv() currently only works correctly in case of the default oneArg = TRUE.

Examples

(cMW2 <- norMix2call(MW.nm2))
deriv(cMW2, "x")

(fMW1 <- as.function (MW.nm1))
(eMW3 <- as.expression(MW.nm3))
stopifnot(is.call (cMW2), is.call(norMix2call(MW.nm2, FALSE)),
is.function (fMW1), is.function (as.function (MW.nm4)),
is.expression(eMW3), is.expression(as.expression(MW.nm5)))
EM and MLE Estimation of Univariate Normal Mixtures

Description

These functions estimate the parameters of a univariate (finite) normal mixture using the EM algorithm or Likelihood Maximization via optim(., method = "BFGS").

Usage

norMixEM(x, m, name = NULL, sd.min = 1e-07 * diff(range(x))/m, trafo = c("clr1", "logit"), maxiter = 100, tol = sqrt(.Machine$double.eps), trace = 1)
norMixMLE(x, m, name = NULL, trafo = c("clr1", "logit"), maxiter = 100, tol = sqrt(.Machine$double.eps), trace = 2)

Arguments

x numeric: the data for which the parameters are to be estimated.
m integer or factor: If m has length 1 it specifies the number of mixture components, otherwise it is taken to be a vector of initial cluster assignments, see details below.
name character, passed to norMix. The default, NULL, uses match.call().
sd.min number: the minimal value that the normal components’ standard deviations (sd) are allowed to take. A warning is printed if some of the final sd’s are this boundary.
trafo character string specifying the transformation of the component weight $w m$-vector (mathematical notation in norMix: $\pi_j, j = 1, \ldots, m$) to an $(m - 1)$-dimensional unconstrained parameter vector in our parametrization. See nM2par for details.
maxiter integer: maximum number of EM iterations.
tol numeric: EM iterations stop if relative changes of the log-likelihood are smaller than tol.
trace integer (or logical) specifying if the iterations should be traced and how much output should be produced. The default, 1 prints a final one line summary, where trace = 2 produces one line of output per iteration.

Details

Estimation of univariate mixtures can be very sensitive to initialization. By default, norMixEM and norMixMLE cut the data into $m$ groups of approximately equal size. See examples below for other initialization possibilities.
The EM algorithm consists in repeated application of E- and M- steps until convergence. Mainly for didactical reasons, we also provide the functions `estep.nm`, `mstep.nm`, and `emstep.nm`.

The MLE, Maximum Likelihood Estimator, maximizes the likelihood using `optim`, using the same advantageous parametrization as `llnorMix`.

**Value**

An object of class `norMix`.

**Author(s)**

EM: Friedrich Leisch, originally; Martin Maechler vectorized it in `m`, added `trace` etc.

MLE: M.Maechler

**Examples**

```r
ex <- norMix(mu = c(-1,2,5), sig2 = c(1, 0.5, 3))
plot(ex, col="gray", p.norm=FALSE)

x <- rnorMix(100, ex)
lines(density(x))
rug(x)

## EM estimation may fail depending on random sample
ex1 <- norMixEM(x, 3, trace=2) #-> warning (sometimes)
ex1
plot(ex1)

## initialization by cut() into intervals of equal length:
ex2 <- norMixEM(x, cut(x, 3))
ex2

## initialization by kmeans():
k3 <- kmeans(x, 3)$cluster
ex3 <- norMixEM(x, k3)
ex3

## Now, MLE instead of EM:
exM <- norMixMLE(x, k3, tol = 1e-12, trace=4)
exM

## real data
data(faithful)
plot(density(faithful$waiting, bw = "SJ"), ylim=c(0,0.044))
rug(faithful$waiting)

(nmF <- norMixEM(faithful$waiting, 2))
lines(nmF, col=2)
## are three components better?
nmF3 <- norMixEM(faithful$waiting, 3, maxiter = 200)
lines(nmF3, col="forestgreen")
```
Description

The `plot` and `lines` methods for `norMix` objects draw the normal mixture density, optionally additionally with a fitted normal density.

Usage

```r
## S3 method for class 'norMix'
plot(x, type = "l", n = 511, xout = NULL, xlim = NULL, ylim,
     xlab = "x", ylab = "f(x)", main = attr(x, "name"), lwd = 1.4,
     p.norm = !p.comp, p.h0 = TRUE, p.comp = FALSE,
     parNorm = list(col = 2, lty = 2, lwd = 0.4),
     parH0 = list(col = 3, lty = 3, lwd = 0.4),
     parComp = list(col = "blue3", lty = 3, lwd = 0.4), ...)

## S3 method for class 'norMix'
lines(x, type = "l", n = 511, xout = NULL,
      lwd = 1.4, p.norm = FALSE, parNorm = list(col = 2, lty = 2, lwd = 0.4),
      ...)```

Arguments

- `x` object of class `norMix`.
- `type` character denoting type of plot, see, e.g. `lines`.
- `n` number of points to generate if `xout` is unspecified.
- `xout` numeric or `NULL` giving the abscissae at which to draw the density.
- `xlim` range of x values to use; particularly important if `xout` is not specified where `xlim` is passed to `dnorMix` and gets a smart default if unspecified.
- `ylim` range of y values to use; by default, if not specified (or containing NA), a smart default is used.
- `xlab`, `ylab` labels for the x and y axis with defaults.
- `main` main title of plot, defaulting to the `norMix` name.
- `lwd` line width for plotting with a non-standard default.
- `p.norm` logical indicating if the normal density with the same mean and variance should be drawn as well.
- `p.h0` logical indicating if the line `y = 0` should be drawn.
- `p.comp` logical indicating if the Gaussian components should also be drawn individually.
- `parNorm` graphical parameters for drawing the normal density if `p.norm` is true.
- `parH0` graphical parameters for drawing the line `y = 0` if `p.h0` is true.
- `parComp` graphical parameters for drawing the single components if `p.comp` is true.
- `...` further arguments passed to and from methods.
pnorMix

Author(s)
Martin Maechler

See Also
norMix for the construction and further methods, particularly dnorMix which is used here.

Examples
plot(norMix(m=c(0,3), sigma = c(2,1))) # -> var = c(2^2, 1) = c(4, 1)
plot(MW.nm4, p.norm=FALSE, p.comp = TRUE)
plot(MW.nm4, p.norm=FALSE, p.comp = TRUE, ylim = c(0, 2))# now works
stopifnot(all.equal(c(0,2), par("yaxp")[1:2], tol= 1e-15))

## Further examples in ?norMix and ?rnorMix

---

### pnorMix

Normal Mixture Cumulative Distribution and Quantiles

#### Description
Compute cumulative probabilities or quantiles (the inverse) for a normal mixture specified as norMix object.

#### Usage

```r
pnorMix(q, obj, lower.tail = TRUE, log.p = FALSE)
qnorMix(p, obj, lower.tail = TRUE, log.p = FALSE,
       tol = .Machine$double.eps^0.25, maxiter = 1000, traceRootsearch = 0,
       method = c("interpQspline", "interpspline", "eachRoot", "root2"),
       l.interp = pmax(1, pmin(20, 1000 / m)), n.mu.interp = 100)
```

#### Arguments

- `obj` an object of class norMix.
- `q` numeric vector of quantiles.
- `p` numeric vector of probabilities. Note that for all methods but "eachRoot", qnorMix(p, *) works with the full vector `p`, typically using (inverse) interpolation approaches; consequently the result is very slightly dependent on `p` as a whole.
- `lower.tail` logical; if TRUE (default), probabilities are \( P[X \leq x] \), otherwise, \( P[X > x] \).
- `log.p` logical; if TRUE, probabilities `p` are given as log(p).
- `tol, maxiter` tolerance and maximal number of iterations for the root search algorithm, see method below and uniroot.
traceRootsearch

logical or integer in \{0, 1, 2, 3\}, determining the amount of information printed during root search.

method

a string specifying which algorithm is used for the “root search”. Originally, the only method was a variation of “eachRoot”, which is the default now when only very few quantiles are sought. For large \texttt{m.normMix()}, the default is set to “root2”, currently.

l.interp

positive integer for method = “interQpspline” or “interpspline”, determining the number of values in each “mu-interval”.

n.mu.interp

positive integer for method = “interQpspline” or “interpspline”, determining the (maximal) number of mu-values to be used as knots for inverse interpolation.

Details

Whereas the distribution function \texttt{pnormMix} is the trivial sum of weighted normal probabilities (\texttt{pnorm}), its inverse, \texttt{qnormMix} is computed numerically: For each \(p\) we search for \(q\) such that \(\texttt{pnormMix(obj, q) == p}\), i.e., \(f(q) = 0\) for \(f(q) := \texttt{pnormMix(obj, q) - p}\). This is a root finding problem which can be solved by \texttt{uniroot(f, lower, upper, *)}. If \(\text{length}(p) <= 2\) or method = “eachRoot”, this happens one for one for the sorted \(p\)’s. Otherwise, we start by doing this for the outermost non-trivial (0 < \(p < 1\)) values of \(p\).

For method = “interQpspline” or “interpspline”, we now compute \(p. <- \texttt{pnormMix(q., obj)}\) for values \(q.\) which are a grid of length \(l.interp\) in each interval \([q_j, q_{j+1}]\), where \(q_j\) are the “X-extremes” plus (a sub sequence of length \(n.mu.interp\) of) the ordered mu\([j]\)’s. Then, we use \texttt{montone} inverse interpolation (\texttt{splinefun(q., p., method=“monoH.FC”)}) plus a few (maximally maxiter, typically one!) Newton steps. The default, “interQpspline”, additionally logit-transforms the \(p.\) values to make the interpolation more linear. This method is faster, particularly for large \(\text{length}(p)\).

Value

a numeric vector of the same length as \(p\) or \(q\), respectively.

Author(s)

Very first version (for length-1 \(p, q\)) by Erik Jørgensen <Erik.Jorgensen@agrsci.dk>.

See Also

\texttt{dnorMix} for the density function.

Examples

\texttt{mW.mm3} # the "strange skew" one
plot(mW.mm3)
## now the cumulative :
x <- seq(-4,4, length=1001)
plot(x, pnormMix(x, mW.mm3), type="l", col=2)
## and some of its inverse :

\texttt{pnorMix}
```r
pp <- seq(.1, .9, by=.1)
plot(qnorMix(pp, MW.nm3), pp)

## The "true" median of a normal mixture:
median.norMix <- function(x) qnorMix(1/2, x)
median.norMix(MW.nm3) ## -2.32
```

---

### r.norMix

**Ratio of Normal Mixture to Corresponding Normal**

**Description**

Compute \( r(x) = f(x)/f_0(x) \) where \( f() \) is a normal mixture density and \( f_0 \) the normal density with the same mean and variance as \( f \).

**Usage**

```r
r.norMix(obj, x = NULL, xlim = NULL, n = 511, xy.return = TRUE)
```

**Arguments**

- **obj**: an object of class `norMix`.
- **x**: numeric vector with abscissa values where to evaluate the density. Default is constructed from \( n \) (and \( xlim \) if specified).
- **xlim**: range of abscissa values, used if \( x == NULL \). By default, \( xlim \) taken as mean plus/minus 3 standard deviations of the normal mixture.
- **n**: number of abscissa values to generate if \( x \) is not specified.
- **xy.return**: logical indicating if the result should be a list or just a numeric vector, see below.

**Value**

It depends on `xy.return`. If it’s false, a numeric vector of the same length as \( x \), if true (as per default), a list that can be plotted, with components

- **x**: abscissa values corresponding to argument \( x \).
- **y**: corresponding values \( r(x) \).
- **f0**: values of the moment matching normal density \( f_0(x) \).

**Note**

The ratio function is used in certain semi-parametric density estimation methods (and theory).
Examples

d3 <- norMix(m = 5*(0:2), w = c(0.6, 0.3, 0.1))
plot(d3)
rd3 <- r.norMix(d3)
str(rd3)
stopifnot(rd3 $ y == r.norMix(d3, xy.ret = FALSE))
par(new = TRUE)
plot(rd3, type = "l", col = 3, axes = FALSE, xlab = "", ylab="")
axis(4, col.axis=3)

rnorMix

Generate 'Normal Mixture' Distributed Random Numbers

Description

Generate n random numbers, distributed according to a normal mixture.

Usage

rnorMix(n, obj)

Arguments

n the number of random numbers desired.
obj an object of class norMix.

Details

For a mixture of m, i.e., m.norMix(obj), components, generate the number in each component as multinomial, and then use rnorm for each.

Note that the these integer (multinomial) numbers are generated via sample(), which is by .Random.seed, notably from RNGkind(sample.kind = ..) which changed with R version 3.6.0.

Value

numeric vector of length n.

See Also

dnorMix for the density, and norMix for the construction and further methods.

Examples

x <- rnorMix(5000, MW.nm10)
hist(x)# you don't see the claw
plot(density(x), ylim = c(0,0.6),
     main = "Estim. and true 'MW.nm10' density")
lines(MW.nm10, col = "orange")
Description

Sorting a "norMix" object (see `norMix`), sorts along the mu values; i.e., for the default `decreasing = FALSE` the resulting `x[, "mu"]` are sorted from left to right.

Usage

```r
## S3 method for class 'norMix'
sort(x, decreasing = FALSE, ...)
```

Arguments

- `x` an object of class "norMix".
- `decreasing` logical indicating if sorting should be up or down.
- `...` further arguments passed to `sort(x[, "mu"], *)`.

Value

a "norMix" object like `x`.

Examples

```r
sort(MW.nm9)
stopifnot(identical(MW.nm2, sort(MW.nm2)))
```
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