

Package ‘tmvtnorm’

July 23, 2025

Version 1.6

Date 2023-12-05

Title Truncated Multivariate Normal and Student t Distribution

Imports stats, methods

Depends R (>= 1.9.0), mvtnorm, utils, Matrix, stats4, gmm

Encoding UTF-8

Suggests lattice

Description

Random number generation for the truncated multivariate normal and Student t distribution. Computes probabilities, quantiles and densities, including one-dimensional and bivariate marginal densities. Computes first and second moments (i.e. mean and covariance matrix) for the double-truncated multinormal case.

License GPL (>= 2)

URL <https://www.r-project.org>

NeedsCompilation yes

Author Stefan Wilhelm [aut, cre],
Manjunath B G [aut]

Maintainer Stefan Wilhelm <wilhelm@financial.com>

Repository CRAN

Date/Publication 2023-12-05 16:10:02 UTC

Contents

dtmvnorm.marginal	2
dtmvnorm.marginal2	5
dtmvt	7
gmm.tmvnorm	9
mle.tmvnorm	11
mtmvnorm	14
ptmvnorm	15
ptmvt	17

ptmvtnorm.marginal	18
qtmvtnorm-marginal	20
rtmvtnorm	21
rtmvtnorm2	27
rtmvtnorm	30
tmvtnorm	32

Index 35

dtmvnorm.marginal	<i>One-dimensional marginal density functions from a Truncated Multivariate Normal distribution</i>
-------------------	---

Description

This function computes the one-dimensional marginal density function from a Truncated Multivariate Normal density function using the algorithm given in Cartinhour (1990).

Usage

```
dtmvnorm.marginal(xn, n=1,
  mean= rep(0, nrow(sigma)),
  sigma=diag(length(mean)),
  lower=rep(-Inf, length = length(mean)),
  upper=rep( Inf, length = length(mean)),
  log=FALSE)
```

Arguments

xn	Vector of quantiles to calculate the marginal density for.
n	Index position (1..k) within the random vector x to calculate the one-dimensional marginal density for.
mean	Mean vector, default is rep(0, length = nrow(sigma)).
sigma	Covariance matrix, default is diag(length(mean)).
lower	Vector of lower truncation points,\ default is rep(-Inf, length = length(mean)).
upper	Vector of upper truncation points,\ default is rep(Inf, length = length(mean)).
log	Logical; if TRUE, densities d are given as log(d).

Details

The one-dimensional marginal density $f_i(x_i)$ of x_i is

$$f_i(x_i) = \int_{a_1}^{b_1} \dots \int_{a_{i-1}}^{b_{i-1}} \int_{a_{i+1}}^{b_{i+1}} \dots \int_{a_k}^{b_k} f(x) dx_{-i}$$

Note that the one-dimensional marginal density is not truncated normal, but only conditional densities are truncated normal.

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>

References

Cartinhour, J. (1990). One-dimensional marginal density functions of a truncated multivariate normal density function. *Communications in Statistics - Theory and Methods*, **19**, 197–203

Arnold et al. (1993). The Nontruncated Marginal of a Truncated Bivariate Normal Distribution. *Psychometrika*, **58**, 471–488

Examples

```
#####
#
# Example 1: truncated bivariate normal
#
#####

# parameters of the bivariate normal distribution
sigma = matrix(c(1, 0.95,
                 0.95, 1), 2, 2)
mu = c(0,0)

# sample from multivariate normal distribution
X = rmvnorm(5000, mu, sigma)

# tuncation in x2 with x2 <= 0
X.trunc = X[X[,2]<0,]

# plot the realisations before and after truncation
par(mfrow=c(2,2))
plot(X, col="gray", xlab=expression(x[1]), ylab=expression(x[2]),
     main="realisations from a\n truncated bivariate normal distribution")
points(X.trunc)
abline(h=0, lty=2, col="gray")
#legend("topleft", col=c("gray", "black"))

# marginal density for x1 from realisations
plot(density(X.trunc[,1]), main=expression("marginal density for "x[1]))

# one-dimensional marginal density for x1 using the formula
x <- seq(-5, 5, by=0.01)
fx <- dtmvnorm.marginal(x, n=1, mean=mu, sigma=sigma,
                        lower=c(-Inf,-Inf), upper=c(Inf,0))
lines(x, fx, lwd=2, col="red")

# marginal density for x2
plot(density(X.trunc[,2]), main=expression("marginal density for "x[2]))

# one-dimensional marginal density for x2 using the formula
x <- seq(-5, 5, by=0.01)
```

```

fx <- dtmvnorm.marginal(x, n=2, mean=mu, sigma=sigma,
  lower=c(-Inf,-Inf), upper=c(Inf,0))
lines(x, fx, lwd=2, col="blue")

#####
#
# Example 2 : truncated trivariate normal
#
#####

# parameters of the trivariate normal distribution
sigma = outer(1:3,1:3,pmin)
mu     = c(0,0,0)

# sample from multivariate normal distribution
X      = rmvnorm(2000, mu, sigma)

# truncation in x2 and x3 : x2 <= 0, x3 <= 0
X.trunc = X[X[,2]<=0 & X[,3]<=0,]

par(mfrow=c(2,3))
plot(X, col="gray", xlab=expression(x[1]), ylab=expression(x[2]),
  main="realisations from a\n truncated trivariate normal distribution")
points(X.trunc, col="black")
abline(h=0, lty=2, col="gray")

plot(X[,2:3], col="gray", xlab=expression(x[2]), ylab=expression(x[3]),
  main="realisations from a\n truncated trivariate normal distribution")
points(X.trunc[,2:3], col="black")
abline(h=0, lty=2, col="gray")
abline(v=0, lty=2, col="gray")

plot(X[,c(1,3)], col="gray", xlab=expression(x[1]), ylab=expression(x[3]),
  main="realisations from a\n truncated trivariate normal distribution")
points(X.trunc[,c(1,3)], col="black")
abline(h=0, lty=2, col="gray")

# one-dimensional marginal density for x1 from realisations and formula
plot(density(X.trunc[,1]), main=expression("marginal density for "x[1]))
x <- seq(-5, 5, by=0.01)
fx <- dtmvnorm.marginal(x, n=1, mean=mu, sigma=sigma,
  lower=c(-Inf,-Inf,-Inf), upper=c(Inf,0,0))
lines(x, fx, lwd=2, col="red")

# one-dimensional marginal density for x2 from realisations and formula
plot(density(X.trunc[,2]), main=expression("marginal density for "x[2]))
x <- seq(-5, 5, by=0.01)
fx <- dtmvnorm.marginal(x, n=2, mean=mu, sigma=sigma,
  lower=c(-Inf,-Inf,-Inf), upper=c(Inf,0,0))
lines(x, fx, lwd=2, col="red")

# one-dimensional marginal density for x3 from realisations and formula
plot(density(X.trunc[,3]), main=expression("marginal density for "x[3]))

```

```
x <- seq(-5, 5, by=0.01)
fx <- dtmvnorm.marginal(x, n=3, mean=mu, sigma=sigma,
  lower=c(-Inf,-Inf,-Inf), upper=c(Inf,0,0))
lines(x, fx, lwd=2, col="red")
```

dtmvnorm.marginal2	<i>Bivariate marginal density functions from a Truncated Multivariate Normal distribution</i>
--------------------	---

Description

This function computes the bivariate marginal density function $f(x_q, x_r)$ from a k-dimensional Truncated Multivariate Normal density function ($k \geq 2$). The bivariate marginal density is obtained by integrating out ($k-2$) dimensions as proposed by Tallis (1961). This function is basically an extraction of the Leppard and Tallis (1989) Fortran code for moments calculation, but extended to the double truncated case.

Usage

```
dtmvnorm.marginal2(xq, xr, q, r,
  mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  lower = rep(-Inf, length = length(mean)),
  upper = rep(Inf, length = length(mean)),
  log = FALSE, pmvnorm.algorithm=GenzBretz())
```

Arguments

xq	Value x_q
xr	Value x_r
q	Index position for x_q within mean vector to calculate the bivariate marginal density for.
r	Index position for x_r within mean vector to calculate the bivariate marginal density for.
mean	Mean vector, default is <code>rep(0, length = nrow(sigma))</code> .
sigma	Covariance matrix, default is <code>diag(length(mean))</code> .
lower	Vector of lower truncation points, default is <code>rep(-Inf, length = length(mean))</code> .
upper	Vector of upper truncation points, default is <code>rep(Inf, length = length(mean))</code> .
log	Logical; if TRUE, densities d are given as <code>log(d)</code> .
pmvnorm.algorithm	Algorithm used for pmvnorm

Details

The bivariate marginal density function $f(x_q, x_r)$ for $x \sim TN(\mu, \Sigma, a, b)$ and $q \neq r$ is defined as

$$F_{q,r}(x_q = c_q, x_r = c_r) = \int_{a_1}^{b_1} \dots \int_{a_{q-1}}^{b_{q-1}} \int_{a_{q+1}}^{b_{q+1}} \dots \int_{a_{r-1}}^{b_{r-1}} \int_{a_{r+1}}^{b_{r+1}} \dots \int_{a_k}^{b_k} \varphi_{\alpha\Sigma}(x_s, c_q, c_r) dx_s$$

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>, Manjunath B G <bgmanjunath@gmail.com>

References

- Tallis, G. M. (1961). The moment generating function of the truncated multinormal distribution. *Journal of the Royal Statistical Society, Series B*, **23**, 223–229
- Leppard, P. and Tallis, G. M. (1989). Evaluation of the Mean and Covariance of the Truncated Multinormal *Applied Statistics*, **38**, 543–553
- Manjunath B G and Wilhelm, S. (2009). Moments Calculation For the Double Truncated Multivariate Normal Density. Working Paper. Available at SSRN: <https://www.ssrn.com/abstract=1472153>

Examples

```
lower = c(-0.5, -1, -1)
upper = c( 2.2,  2,  2)

mean = c(0,0,0)
sigma = matrix(c(2.0, -0.6,  0.7,
                 -0.6,  1.0, -0.2,
                 0.7, -0.2,  1.0), 3, 3)

# generate random samples from untruncated and truncated distribution
Y = rmvnorm(10000, mean=mean, sigma=sigma)
X = rtmvnorm(500, mean=mean, sigma=sigma, lower=lower, upper=upper,
             algorithm="gibbs")

# compute bivariate marginal density of x1 and x2
xq <- seq(lower[1], upper[1], by=0.1)
xr <- seq(lower[2], upper[2], by=0.1)

grid <- matrix(NA, length(xq), length(xr))
for (i in 1:length(xq))
{
  for (j in 1:length(xr))
  {
    grid[i,j] = dtmvnorm.marginal2(xq=xq[i], xr=xr[j],
                                   q=1, r=2, sigma=sigma, lower=lower, upper=upper)
  }
}

plot(Y[,1], Y[,2], xlim=c(-4, 4), ylim=c(-4, 4),
```

```

    main=expression("bivariate marginal density ("*x[1]*","*x[2]*")"),
    xlab=expression(x[1]), ylab=expression(x[2]), col="gray80")
points(X[,1], X[,2], col="black")

lines(x=c(lower[1], upper[1], upper[1], lower[1], lower[1]),
      y=c(lower[2],lower[2],upper[2],upper[2],lower[2]),
      lty=2, col="red")
contour(xq, xr, grid, add=TRUE, nlevels = 8, col="red", lwd=2)

# scatterplot matrices for untruncated and truncated points
require(lattice)
splom(Y)
splom(X)

```

dtmvt

Truncated Multivariate Student t Density

Description

This function provides the joint density function for the truncated multivariate Student t distribution with mean vector equal to mean, covariance matrix sigma, degrees of freedom parameter df and lower and upper truncation points lower and upper.

Usage

```

dtmvt(x, mean = rep(0, nrow(sigma)), sigma = diag(length(mean)), df = 1,
lower = rep(-Inf, length = length(mean)),
upper = rep(Inf, length = length(mean)), log = FALSE)

```

Arguments

x	Vector or matrix of quantiles. If x is a matrix, each row is taken to be a quantile.
mean	Mean vector, default is rep(0, nrow(sigma)).
sigma	Covariance matrix, default is diag(length(mean)).
df	degrees of freedom parameter
lower	Vector of lower truncation points, default is rep(-Inf, length = length(mean)).
upper	Vector of upper truncation points, default is rep(Inf, length = length(mean)).
log	Logical; if TRUE, densities d are given as log(d).

Details

The Truncated Multivariate Student t Distribution is a conditional Multivariate Student t distribution subject to (linear) constraints $a \leq \mathbf{x} \leq b$.

The density of the p -variate Multivariate Student t distribution with ν degrees of freedom is

$$f(\mathbf{x}) = \frac{\Gamma((\nu + p)/2)}{(\pi\nu)^{p/2}\Gamma(\nu/2)\|\Sigma\|^{1/2}} \left[1 + \frac{1}{\nu}(\mathbf{x} - \mu)^T \Sigma^{-1}(\mathbf{x} - \mu)\right]^{-(\nu+p)/2}$$

The density of the truncated distribution $f_{a,b}(x)$ with constraints ($a \leq x \leq b$) is accordingly

$$f_{a,b}(x) = \frac{f(\mathbf{x})}{P(a \leq x \leq b)}$$

Value

a numeric vector with density values

Author(s)

Stefan Wilhelm <wilhelm@financial.com>

References

Geweke, J. F. (1991) Efficient simulation from the multivariate normal and Student-t distributions subject to linear constraints and the evaluation of constraint probabilities. https://www.researchgate.net/publication/2335219_Efficient_Simulation_from_the_Multivariate_Normal_and_Student-t_Distributions_Subject_to_Linear_Constraints_and_the_Evaluation_of_Constraint_Probabilities

Samuel Kotz, Saralees Nadarajah (2004). Multivariate t Distributions and Their Applications. *Cambridge University Press*

See Also

[ptmvt](#) and [rtmvt](#) for probabilities and random number generation in the truncated case, see [dmvt](#), [rmvt](#) and [pmvt](#) for the untruncated multi-t distribution.

Examples

```
# Example

x1 <- seq(-2, 3, by=0.1)
x2 <- seq(-2, 3, by=0.1)

mean <- c(0,0)
sigma <- matrix(c(1, -0.5, -0.5, 1), 2, 2)
lower <- c(-1,-1)

density <- function(x)
{
  z=dtmvt(x, mean=mean, sigma=sigma, lower=lower)
  z
}

fgrid <- function(x, y, f)
{
  z <- matrix(nrow=length(x), ncol=length(y))
  for(m in 1:length(x)){
    for(n in 1:length(y)){
```



```

z[m,n] <- f(c(x[m], y[n]))
}
}
z
}

# compute multivariate-t density d for grid
d <- fgrid(x1, x2, function(x) dtmvt(x, mean=mean, sigma=sigma, lower=lower))

# compute multivariate normal density d for grid
d2 <- fgrid(x1, x2, function(x) dtmvnorm(x, mean=mean, sigma=sigma, lower=lower))

# plot density as contourplot
contour(x1, x2, d, nlevels=5, main="Truncated Multivariate t Density",
xlab=expression(x[1]), ylab=expression(x[2]))

contour(x1, x2, d2, nlevels=5, add=TRUE, col="red")
abline(v=-1, lty=3, lwd=2)
abline(h=-1, lty=3, lwd=2)

```

gmm.tmvnorm

*GMM Estimation for the Truncated Multivariate Normal Distribution***Description**

Generalized Method of Moments (GMM) Estimation for the Truncated Multivariate Normal Distribution

Usage

```

gmm.tmvnorm(X,
  lower = rep(-Inf, length = ncol(X)),
  upper = rep(+Inf, length = ncol(X)),
  start = list(mu = rep(0, ncol(X)), sigma = diag(ncol(X))),
  fixed = list(),
  method=c("ManjunathWilhelm","Lee"),
  cholesky = FALSE,
  ...)

```

Arguments

X	Matrix of quantiles, each row is taken to be a quantile.
lower	Vector of lower truncation points, default is <code>rep(-Inf, length = ncol(X))</code> .
upper	Vector of upper truncation points, default is <code>rep(+Inf, length = ncol(X))</code> .
start	Named list with elements <code>mu</code> (mean vector) and <code>sigma</code> (covariance matrix). Initial values for optimizer.
fixed	Named list. Parameter values to keep fixed during optimization.

method	Which set of moment conditions used, possible methods are "ManjunathWilhelm" (default) and "Lee".
cholesky	if TRUE, we use the Cholesky decomposition of sigma as parametrization
...	Further arguments to pass to gmm

Details

This method performs an estimation of the parameters mean and sigma of a truncated multinormal distribution using the Generalized Method of Moments (GMM), when the truncation points lower and upper are known. `gmm.tmvnorm()` is a wrapper for the general GMM method [gmm](#), so one does not have to specify the moment conditions.

Manjunath/Wilhelm moment conditions

Because the first and second moments can be computed thanks to the [mtmvnorm](#) function, we can set up a method-of-moments estimator by equating the sample moments to their population counterparts. This way we have an exactly identified case.

Lee (1979,1983) moment conditions

The recursive moment conditions presented by Lee (1979,1983) are defined for $l = 0, 1, 2, \dots$ as

$$\sigma^{iT} E(x_i^l \mathbf{x}) = \sigma^{iT} \mu E(x_i^l) + l E(x_i^{l-1}) + \frac{a_i^l F_i(a_i)}{F} - \frac{b_i^l F_i(b_i)}{F}$$

where $E(x_i^l)$ and $E(x_i^l \mathbf{x})$ are the moments of x_i^l and $x_i^l \mathbf{x}$ respectively and $F_i(c)/F$ is the one-dimensional marginal density in variable i as calculated by [dtmvnorm.marginal](#). σ^{iT} is the i -th column of the inverse covariance matrix Σ^{-1} .

This method returns an object of class `gmm`, for which various diagnostic methods are available, like `profile()`, `confint()` etc. See examples.

Value

An object of class [gmm](#)

Author(s)

Stefan Wilhelm <wilhelm@financial.com>

References

- Tallis, G. M. (1961). The moment generating function of the truncated multinormal distribution. *Journal of the Royal Statistical Society, Series B*, **23**, 223–229
- Lee, L.-F. (1979). On the first and second moments of the truncated multi-normal distribution and a simple estimator. *Economics Letters*, **3**, 165–169
- Lee, L.-F. (1983). The determination of moments of the doubly truncated multivariate normal Tobit model. *Economics Letters*, **11**, 245–250
- Manjunath B G and Wilhelm, S. (2009). Moments Calculation For the Double Truncated Multivariate Normal Density. Working Paper. Available at SSRN: <https://www.ssrn.com/abstract=1472153>

See Also[gmm](#)**Examples**

```
## Not run:
set.seed(1.234)

# the actual parameters
lower <- c(-1, -2)
upper <- c(3, Inf)
mu <- c(0, 0)
sigma <- matrix(c(1, 0.8,
                  0.8, 2), 2, 2)

# generate random samples
X <- rtmvnorm(n=500, mu, sigma, lower, upper)

# estimate mean vector and covariance matrix sigma from random samples X
# with default start values
gmm.fit1 <- gmm.tmvnorm(X, lower=lower, upper=upper)

# diagnostic output of the estimated parameters
summary(gmm.fit1)
vcov(gmm.fit1)

# confidence intervals
confint(gmm.fit1)

# choosing a different start value
gmm.fit2 <- gmm.tmvnorm(X, lower=lower, upper=upper,
  start=list(mu=c(0.1, 0.1),
    sigma=matrix(c(1, 0.4, 0.4, 1.8), 2, 2)))
summary(gmm.fit2)

# GMM estimation with Lee (1983) moment conditions
gmm.fit3 <- gmm.tmvnorm(X, lower=lower, upper=upper, method="Lee")
summary(gmm.fit3)
confint(gmm.fit3)

# MLE estimation for comparison
mle.fit1 <- mle.tmvnorm(X, lower=lower, upper=upper)
confint(mle.fit1)

## End(Not run)
```

Description

Maximum Likelihood Estimation for the Truncated Multivariate Normal Distribution

Usage

```
mle.tmvnorm(X,
  lower = rep(-Inf, length = ncol(X)),
  upper = rep(+Inf, length = ncol(X)),
  start = list(mu = rep(0, ncol(X)), sigma = diag(ncol(X))),
  fixed = list(), method = "BFGS",
  cholesky = FALSE,
  lower.bounds = -Inf,
  upper.bounds = +Inf,
  ...)
```

Arguments

X	Matrix of quantiles, each row is taken to be a quantile.
lower	Vector of lower truncation points, default is <code>rep(-Inf, length = ncol(X))</code> .
upper	Vector of upper truncation points, default is <code>rep(+Inf, length = ncol(X))</code> .
start	Named list with elements <code>mu</code> (mean vector) and <code>sigma</code> (covariance matrix). Initial values for optimizer.
fixed	Named list. Parameter values to keep fixed during optimization.
method	Optimization method to use. See optim
cholesky	if TRUE, we use the Cholesky decomposition of <code>sigma</code> as parametrization
lower.bounds	lower bounds/box constraints for method "L-BFGS-B"
upper.bounds	upper bounds/box constraints for method "L-BFGS-B"
...	Further arguments to pass to optim

Details

This method performs a maximum likelihood estimation of the parameters mean and sigma of a truncated multinormal distribution, when the truncation points `lower` and `upper` are known. `mle.tmvnorm()` is a wrapper for the general maximum likelihood method [mle](#), so one does not have to specify the negative log-likelihood function.

The log-likelihood function for a data matrix X ($T \times n$) can be established straightforward as

$$\log L(X|\mu, \Sigma) = -T \log \alpha(\mu, \Sigma) + -T/2 \log \|\Sigma\| - \frac{1}{2} \sum_{t=1}^T (x_t - \mu)' \Sigma^{-1} (x_t - \mu)$$

As [mle](#), this method returns an object of class `mle`, for which various diagnostic methods are available, like `profile()`, `confint()` etc. See examples.

In order to adapt the estimation problem to [mle](#), the named parameters for mean vector elements are "`mu_i`" and the elements of the covariance matrix are "`sigma_ij`" for the lower triangular matrix elements, i.e. ($j \leq i$).

Value

An object of class `mle-class`

Author(s)

Stefan Wilhelm <wilhelm@financial.com>

See Also

`mle` and `mle-class`

Examples

```
## Not run:
set.seed(1.2345)

# the actual parameters
lower <- c(-1,-1)
upper <- c(1, 2)
mu    <- c(0, 0)
sigma <- matrix(c(1, 0.7,
                  0.7, 2), 2, 2)

# generate random samples
X <- rtmvnorm(n=500, mu, sigma, lower, upper)
method <- "BFGS"

# estimate mean vector and covariance matrix sigma from random samples X
# with default start values
mle.fit1 <- mle.tmvnorm(X, lower=lower, upper=upper)

# diagnostic output of the estimated parameters
summary(mle.fit1)
logLik(mle.fit1)
vcov(mle.fit1)

# profiling the log likelihood and confidence intervals
mle.profile1 <- profile(mle.fit1, X, method="BFGS", trace=TRUE)
confint(mle.profile1)

par(mfrow=c(3,2))
plot(mle.profile1)

# choosing a different start value
mle.fit2 <- mle.tmvnorm(X, lower=lower, upper=upper,
  start=list(mu=c(0.1, 0.1),
    sigma=matrix(c(1, 0.4, 0.4, 1.8),2,2)))
summary(mle.fit2)

## End(Not run)
```

mtmvnorm

Computation of Mean Vector and Covariance Matrix For Truncated Multivariate Normal Distribution

Description

Computation of the first two moments, i.e. mean vector and covariance matrix for the Truncated Multivariate Normal Distribution based on the works of Tallis (1961), Lee (1979) and Leppard and Tallis (1989), but extended to the double-truncated case with general mean and general covariance matrix.

Usage

```
mtmvnorm(mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  lower = rep(-Inf, length = length(mean)),
  upper = rep(Inf, length = length(mean)),
  doComputeVariance=TRUE,
  pmvnorm.algorithm=GenzBretz())
```

Arguments

mean	Mean vector, default is <code>rep(0, length = ncol(x))</code> .
sigma	Covariance matrix, default is <code>diag(ncol(x))</code> .
lower	Vector of lower truncation points,\ default is <code>rep(-Inf, length = length(mean))</code> .
upper	Vector of upper truncation points,\ default is <code>rep(Inf, length = length(mean))</code> .
doComputeVariance	flag whether to compute the variance for users who are interested only in the mean. Defaults to TRUE for backward compatibility.
pmvnorm.algorithm	Algorithm used for pmvnorm

Details

Details for the moment calculation under double truncation and the derivation of the formula can be found in the Manjunath/Wilhelm (2009) working paper. If only a subset of variables are truncated, we calculate the truncated moments only for these and use the Johnson/Kotz formula for the remaining untruncated variables.

Value

tmean	Mean vector of truncated variables
tvar	Covariance matrix of truncated variables

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>, Manjunath B G <bgmanjunath@gmail.com>

References

- Tallis, G. M. (1961). The moment generating function of the truncated multinormal distribution. *Journal of the Royal Statistical Society, Series B*, **23**, 223–229
- Johnson, N./Kotz, S. (1970). *Distributions in Statistics: Continuous Multivariate Distributions* Wiley & Sons, pp. 70–73
- Lee, L.-F. (1979). On the first and second moments of the truncated multi-normal distribution and a simple estimator. *Economics Letters*, **3**, 165–169
- Leppard, P. and Tallis, G. M. (1989). Evaluation of the Mean and Covariance of the Truncated Multinormal. *Applied Statistics*, **38**, 543–553
- Manjunath B G and Wilhelm, S. (2009). Moments Calculation For the Double Truncated Multivariate Normal Density. Working Paper. Available at SSRN: <https://www.ssrn.com/abstract=1472153>

Examples

```
mu    <- c(0.5, 0.5, 0.5)
sigma <- matrix(c( 1, 0.6, 0.3,
                  0.6, 1, 0.2,
                  0.3, 0.2, 2), 3, 3)

a <- c(-Inf, -Inf, -Inf)
b <- c(1, 1, 1)

# compute first and second moments
mtmvnorm(mu, sigma, lower=a, upper=b)

# compare with simulated results
X <- rtmvnorm(n=1000, mean=mu, sigma=sigma, lower=a, upper=b)
colMeans(X)
cov(X)
```

ptmvnorm

Truncated Multivariate Normal Distribution

Description

Computes the distribution function of the truncated multivariate normal distribution for arbitrary limits and correlation matrices based on the pmvnorm() implementation of the algorithms by Genz and Bretz.

Usage

```
ptmvnorm(lowerx, upperx, mean=rep(0, length(lowerx)), sigma,
  lower = rep(-Inf, length = length(mean)),
  upper = rep( Inf, length = length(mean)),
  maxpts = 25000, abseps = 0.001, releps = 0)
```

Arguments

lowerx	the vector of lower limits of length n.
upperx	the vector of upper limits of length n.
mean	the mean vector of length n.
sigma	the covariance matrix of dimension n. Either corr or sigma can be specified. If sigma is given, the problem is standardized. If neither corr nor sigma is given, the identity matrix is used for sigma.
lower	Vector of lower truncation points,\default is rep(-Inf , length = length(mean)).
upper	Vector of upper truncation points,\default is rep(Inf , length = length(mean)).
maxpts	maximum number of function values as integer.
abseps	absolute error tolerance as double.
releps	relative error tolerance as double.

Details

The computation of truncated multivariate normal probabilities and densities is done using conditional probabilities from the standard/untruncated multivariate normal distribution. So we refer to the documentation of the mvtnorm package and the methodology is described in Genz (1992, 1993) and Genz/Bretz (2009).

For properties of the truncated multivariate normal distribution see for example Johnson/Kotz (1970) and Horrace (2005).

Value

The evaluated distribution function is returned with attributes

error	estimated absolute error and
msg	status messages.

References

- Genz, A. (1992). Numerical computation of multivariate normal probabilities. *Journal of Computational and Graphical Statistics*, **1**, 141–150
- Genz, A. (1993). Comparison of methods for the computation of multivariate normal probabilities. *Computing Science and Statistics*, **25**, 400–405
- Genz, A. and Bretz, F. (2009). Computation of Multivariate Normal and t Probabilities. *Lecture Notes in Statistics*, Vol. **195**, Springer-Verlag, Heidelberg.
- Johnson, N./Kotz, S. (1970). Distributions in Statistics: Continuous Multivariate Distributions Wiley & Sons, pp. 70–73
- Horrace, W. (2005). Some Results on the Multivariate Truncated Normal Distribution. *Journal of Multivariate Analysis*, **94**, 209–221

Examples

```
sigma <- matrix(c(5, 0.8, 0.8, 1), 2, 2)
Fx <- ptmvnorm(lowerx=c(-1,-1), upperx=c(0.5,0), mean=c(0,0),
  sigma=sigma, lower=c(-1,-1), upper=c(1,1))
```

ptmvt

*Truncated Multivariate Student t Distribution***Description**

Computes the distribution function of the truncated multivariate t distribution

Usage

```
ptmvt(lowerx, upperx, mean = rep(0, length(lowerx)), sigma, df = 1,
  lower = rep(-Inf, length = length(mean)),
  upper = rep(Inf, length = length(mean)), maxpts = 25000, abseps = 0.001,
  releps = 0)
```

Arguments

lowerx	the vector of lower limits of length n.
upperx	the vector of upper limits of length n.
mean	the mean vector of length n.
sigma	the covariance matrix of dimension n. Either corr or sigma can be specified. If sigma is given, the problem is standardized. If neither corr nor sigma is given, the identity matrix is used for sigma.
df	Degrees of freedom parameter
lower	Vector of lower truncation points, default is rep(-Inf, length = length(mean)).
upper	Vector of upper truncation points, default is rep(Inf, length = length(mean)).
maxpts	maximum number of function values as integer.
abseps	absolute error tolerance as double.
releps	relative error tolerance as double.

Value

The evaluated distribution function is returned with attributes

error	estimated absolute error and
msg	status messages.

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>

References

Geweke, J. F. (1991) Efficient simulation from the multivariate normal and Student-t distributions subject to linear constraints and the evaluation of constraint probabilities. https://www.researchgate.net/publication/2335219_Efficient_Simulation_from_the_Multivariate_Normal_and_Student-t_Distributions_Subject_to_Linear_Constraints_and_the_Evaluation_of_Constraint_Probabilities

Samuel Kotz, Saralees Nadarajah (2004). Multivariate t Distributions and Their Applications. *Cambridge University Press*

Examples

```
sigma <- matrix(c(5, 0.8, 0.8, 1), 2, 2)
Fx <- ptmvt(lowerx=c(-1,-1), upperx=c(0.5,0), mean=c(0,0), sigma=sigma, df=3,
  lower=c(-1,-1), upper=c(1,1))
```

ptmvtnorm.marginal	<i>One-dimensional marginal CDF function for a Truncated Multivariate Normal and Student t distribution</i>
--------------------	---

Description

This function computes the one-dimensional marginal probability function from a Truncated Multivariate Normal and Student t density function using integration in pmvnorm() and pmvt().

Usage

```
ptmvtnorm.marginal(xn,
  n = 1,
  mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  lower = rep(-Inf, length = length(mean)),
  upper = rep(Inf, length = length(mean)))
ptmvt.marginal(xn,
  n = 1,
  mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  df = 1,
  lower = rep(-Inf, length = length(mean)),
  upper = rep(Inf, length = length(mean)))
```

Arguments

xn	Vector of quantiles to calculate the marginal probability for.
n	Index position (1..k) within the random vector xn to calculate the one-dimensional marginal probability for.
mean	the mean vector of length k.

sigma	the covariance matrix of dimension k. Either corr or sigma can be specified. If sigma is given, the problem is standardized. If neither corr nor sigma is given, the identity matrix is used for sigma.
df	degrees of freedom parameter
lower	Vector of lower truncation points, default is rep(-Inf, length = length(mean)).
upper	Vector of upper truncation points, default is rep(Inf, length = length(mean)).

Details

The one-dimensional marginal probability for index i is $F_i(x_i) = P(X_i \leq x_i)$

$$F_i(x_i) = \int_{a_1}^{b_1} \dots \int_{a_{i-1}}^{b_{i-1}} \int_{a_i}^{x_i} \int_{a_{i+1}}^{b_{i+1}} \dots \int_{a_k}^{b_k} f(x) dx = \alpha^{-1} \Phi_k(a, u, \mu, \Sigma)$$

where $u = (b_1, \dots, b_{i-1}, x_i, b_{i+1}, \dots, b_k)'$ is the upper integration bound and Φ_k is the k-dimensional normal probability (i.e. functions pmvnorm() and pmvt() in R package mvtnorm).

Value

Returns a vector of the same length as xn with probabilities.

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>

Examples

```
## Example 1: Truncated multi-normal
lower <- c(-1,-1,-1)
upper <- c(1,1,1)
mean <- c(0,0,0)
sigma <- matrix(c( 1, 0.8, 0.2,
                  0.8,  1, 0.1,
                  0.2, 0.1,  1), 3, 3)

X <- rtmvnorm(n=1000, mean=c(0,0,0), sigma=sigma, lower=lower, upper=upper)

x <- seq(-1, 1, by=0.01)
Fx <- ptmvnorm.marginal(xn=x, n=1, mean=c(0,0,0), sigma=sigma, lower=lower, upper=upper)

plot(ecdf(X[,1]), main="marginal CDF for truncated multi-normal")
lines(x, Fx, type="l", col="blue")

## Example 2: Truncated multi-t
X <- rtmtv(n=1000, mean=c(0,0,0), sigma=sigma, df=2, lower=lower, upper=upper)

x <- seq(-1, 1, by=0.01)
Fx <- ptmvt.marginal(xn=x, n=1, mean=c(0,0,0), sigma=sigma, lower=lower, upper=upper)

plot(ecdf(X[,1]), main="marginal CDF for truncated multi-t")
lines(x, Fx, type="l", col="blue")
```

qtmvnorm-marginal	<i>Quantiles of the Truncated Multivariate Normal Distribution in one dimension</i>
-------------------	---

Description

Computes the equicoordinate quantile function of the truncated multivariate normal distribution for arbitrary correlation matrices based on an inversion of the algorithms by Genz and Bretz.

Usage

```
qtmvnorm.marginal(p,
                  interval = c(-10, 10),
                  tail = c("lower.tail", "upper.tail", "both.tails"),
                  n=1,
                  mean=rep(0, nrow(sigma)),
                  sigma=diag(length(mean)),
                  lower=rep(-Inf, length = length(mean)),
                  upper=rep( Inf, length = length(mean)),
                  ...)
```

Arguments

<code>p</code>	probability.
<code>interval</code>	a vector containing the end-points of the interval to be searched by uniroot .
<code>tail</code>	specifies which quantiles should be computed. <code>lower.tail</code> gives the quantile x for which $P[X \leq x] = p$, <code>upper.tail</code> gives x with $P[X > x] = p$ and <code>both.tails</code> leads to x with $P[-x \leq X \leq x] = p$
<code>n</code>	index (1..n) to calculate marginal quantile for
<code>mean</code>	the mean vector of length <code>n</code> .
<code>sigma</code>	the covariance matrix of dimension <code>n</code> . Either <code>corr</code> or <code>sigma</code> can be specified. If <code>sigma</code> is given, the problem is standardized. If neither <code>corr</code> nor <code>sigma</code> is given, the identity matrix is used for <code>sigma</code> .
<code>lower</code>	Vector of lower truncation points,\default is <code>rep(-Inf, length = length(mean))</code> .
<code>upper</code>	Vector of upper truncation points,\default is <code>rep(Inf, length = length(mean))</code> .
<code>...</code>	additional parameters to be passed to uniroot .

Details

Only equicoordinate quantiles are computed, i.e., the quantiles in each dimension coincide. Currently, the distribution function is inverted by using the [uniroot](#) function which may result in limited accuracy of the quantiles.

Value

A list with four components: `quantile` and `f.quantile` give the location of the quantile and the value of the function evaluated at that point. `iter` and `estim.prec` give the number of iterations used and an approximate estimated precision from [uniroot](#).

See Also

[ptmvnorm](#), [pmvnorm](#)

Examples

```
# finite dimensional distribution of the Geometric Brownian Motion log-returns
# with truncation

# volatility p.a.
sigma=0.4

# risk free rate
r = 0.05

# n=3 points in time
T <- c(0.5, 0.7, 1)

# covariance matrix of Geometric Brownian Motion returns
Sigma = sigma^2*outer(T,T,pmin)

# mean vector of the Geometric Brownian Motion returns
mu     = (r - sigma^2/2) * T

# lower truncation vector a (a<=x<=b)
a = rep(-Inf, 3)

# upper truncation vector b (a<=x<=b)
b = c(0, 0, Inf)

# quantile of the t_1 returns
qtmvnorm.marginal(p=0.95, interval = c(-10, 10), tail = "lower.tail", n=1,
  mean = mu, sigma = Sigma, lower=a, upper=b)
```

rtmvnorm

Sampling Random Numbers From The Truncated Multivariate Normal Distribution

Description

This function generates random numbers from the truncated multivariate normal distribution with mean equal to `mean` and covariance matrix `sigma` (or alternatively precision matrix `H`), lower and upper truncation points `lower` and `upper` with either rejection sampling or Gibbs sampling.

Usage

```
rtmvnorm(n, mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  lower=rep(-Inf, length = length(mean)),
  upper=rep( Inf, length = length(mean)),
  D = diag(length(mean)),
  H = NULL,
  algorithm=c("rejection", "gibbs", "gibbsR"),
  ...)

rtmvnorm.sparseMatrix(n, mean = rep(0, nrow(H)),
  H = sparseMatrix(i=1:length(mean), j=1:length(mean), x=1),
  lower = rep(-Inf, length = length(mean)),
  upper = rep( Inf, length = length(mean)),
  ...)
```

Arguments

n	Number of random points to be sampled. Must be an integer ≥ 1 .
mean	Mean vector, default is <code>rep(0, length = ncol(x))</code> .
sigma	Covariance matrix, default is <code>diag(ncol(x))</code> .
lower	Vector of lower truncation points, default is <code>rep(-Inf, length = length(mean))</code> .
upper	Vector of upper truncation points, default is <code>rep(Inf, length = length(mean))</code> .
D	Matrix for linear constraints, defaults to diagonal matrix.
H	Precision matrix, default is <code>NULL</code> .
algorithm	Method used, possible methods are rejection sampling ("rejection", default), the Fortan Gibbs sampler ("gibbs") and the old Gibbs sampler implementation in R ("gibbsR").
...	additional parameters for Gibbs sampling, given to the internal method <code>rtmvnorm.gibbs()</code> , such as <code>burn.in.samples</code> , <code>start.value</code> and <code>thinning</code> , see details

Details

The generation of random numbers from a truncated multivariate normal distribution is done using either rejection sampling or Gibbs sampling.

Rejection sampling

Rejection sampling is done from the standard multivariate normal distribution. So we use the function `rmvnorm` of the **mvtnorm** package to generate proposals which are either accepted if they are inside the support region or rejected. In order to speed up the generation of N samples from the truncated distribution, we first calculate the acceptance rate α from the truncation points and then generate N/α samples iteratively until we have got N samples. This typically does not take more than 2-3 iterations. Rejection sampling may be very inefficient when the support region is small (i.e. in higher dimensions) which results in very low acceptance rates α . In this case the Gibbs sampler is preferable.

Gibbs sampling

The Gibbs sampler samples from univariate conditional distributions, so all samples can be accepted

except for a burn-in period. The number of burn-in samples to be discarded can be specified, as well as a start value of the chain. If no start value is given, we determine a start value from the support region using either lower bound or upper bound if they are finite, or 0 otherwise.

The Gibbs sampler has been reimplemented in Fortran 90 for performance reasons (`algorithm="gibbs"`). The old R implementation is still accessible through `algorithm="gibbsR"`.

The arguments to be passed along with `algorithm="gibbs"` or `algorithm="gibbsR"` are:

`burn.in.samples` number of samples in Gibbs sampling to be discarded as burn-in phase, must be non-negative.

`start.value` Start value (vector of length `length(mean)`) for the MCMC chain. If one is specified, it must lie inside the support region ($lower \leq start.value \leq upper$). If none is specified, the start value is taken componentwise as the finite lower or upper boundaries respectively, or zero if both boundaries are infinite. Defaults to `NULL`.

`thinning` Thinning factor for reducing autocorrelation of random points in Gibbs sampling. Must be an integer ≥ 1 . We create a Markov chain of length `(n*thinning)` and take only those samples `j=1:(n*thinning)` where `j %% thinning == 0` Defaults to 1 (no thinning of the chain).

Sampling with linear constraints

We extended the method to also simulate from a multivariate normal distribution subject to general linear constraints $lower \leq Dx \leq upper$. For general D , both rejection sampling or Gibbs sampling according to Geweke (1991) are available.

Gibbs sampler and the use of the precision matrix H

Why is it important to have a random sampler that works with the precision matrix? Especially in Bayesian and spatial statistics, there are a number of high-dimensional applications where the precision matrix H is readily available, but is sometimes nearly singular and cannot be easily inverted to σ . Additionally, it turns out that the Gibbs sampler formulas are much simpler in terms of the precision matrix than in terms of the covariance matrix. See the details of the Gibbs sampler implementation in the package vignette or for example Geweke (2005), pp.171-172. (Thanks to Miguel Godinho de Matos from Carnegie Mellon University for pointing me to this.) Therefore, we now provide an interface for the direct use of the precision matrix H in `rtmvnorm()`.

Gibbs sampler with sparse precision matrix H

The size of the covariance matrix σ or precision matrix H - if expressed as a dense `matrix` - grows quadratic with the number of dimensions d . For high-dimensional problems (such as $d > 5000$), it is no longer efficient and appropriate to work with dense matrix representations, as one quickly runs into memory problems.

It is interesting to note that in many applications the precision matrix, which holds the conditional dependencies, will be sparse, whereas the covariance matrix will be dense. Hence, expressing H as a sparse matrix will significantly reduce the amount of memory to store this matrix and allows much larger problems to be handled. In the current version of the package, the precision matrix (not σ since it will be dense in most cases) can be passed to `rtmvnorm.sparseMatrix()` as a `sparseMatrix` from the `Matrix` package. See the examples section below for a usage example.

Warning

A word of caution is needed for useRs that are not familiar with Markov Chain Monte Carlo methods like Gibbs sampling:

Rejection sampling is exact in the sense that we are sampling directly from the target distribution and the random samples generated are independent. So it is clearly the default method.

Markov Chain Monte Carlo methods are only approximate methods, which may suffer from several problems:

- Poor mixing
- Convergence problems
- Correlation among samples

Diagnostic checks for Markov Chain Monte Carlo include trace plots, CUSUM plots and autocorrelation plots like [acf](#). For a survey see for instance Cowles (1996).

That is, consecutive samples generated from `rtmvnorm(..., algorithm=c("gibbs", "gibbsR"))` are correlated (see also example 3 below). One way of reducing the autocorrelation among the random samples is "thinning" the Markov chain, that is recording only a subset/subsequence of the chain. For example, one could record only every 100th sample, which clearly reduces the autocorrelation and "increases the independence". But thinning comes at the cost of higher computation times, since the chain has to run much longer. We refer to autocorrelation plots in order to determine optimal thinning.

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>, Manjunath B G <bgmanjunath@gmail.com>

References

- Alan Genz, Frank Bretz, Tetsuhisa Miwa, Xuefei Mi, Friedrich Leisch, Fabian Scheipl, Torsten Hothorn (2009). *mvtnorm: Multivariate Normal and t Distributions*. R package version 0.9-7. URL <https://CRAN.R-project.org/package=mvtnorm>
- Johnson, N./Kotz, S. (1970). *Distributions in Statistics: Continuous Multivariate Distributions Wiley & Sons*, pp. 70–73
- Horrace, W. (2005). Some Results on the Multivariate Truncated Normal Distribution. *Journal of Multivariate Analysis*, **94**, 209–221
- Jayesh H. Kotecha and Petar M. Djuric (1999). Gibbs Sampling Approach For Generation of Truncated Multivariate Gaussian Random Variables *IEEE Computer Society*, 1757–1760
- Cowles, M. and Carlin, B. (1996). Markov Chain Monte Carlo Convergence Diagnostics: A Comparative Review *Journal of the American Statistical Association*, **91**, 883–904
- Geweke, J. F. (1991). Efficient Simulation from the Multivariate Normal and Student-t Distributions Subject to Linear Constraints *Computer Science and Statistics. Proceedings of the 23rd Symposium on the Interface. Seattle Washington, April 21-24, 1991*, 571–578
- Geweke, J. F. (2005). *Contemporary Bayesian Econometrics and Statistics*, Wiley & Sons, pp.171–172

See Also

[ptmvnorm](#), [pmvnorm](#), [rmvnorm](#), [dmvnorm](#)

Examples

```
#####
#
# Example 1:
# rejection sampling in 2 dimensions
#
#####

sigma <- matrix(c(4,2,2,3), ncol=2)
x <- rtmvnorm(n=500, mean=c(1,2), sigma=sigma, upper=c(1,0))
plot(x, main="samples from truncated bivariate normal distribution",
     xlim=c(-6,6), ylim=c(-6,6),
     xlab=expression(x[1]), ylab=expression(x[2]))
abline(v=1, lty=3, lwd=2, col="gray")
abline(h=0, lty=3, lwd=2, col="gray")

#####
#
# Example 2:
# Gibbs sampler for 4 dimensions
#
#####

C <- matrix(0.8, 4, 4)
diag(C) <- rep(1, 4)
lower <- rep(-4, 4)
upper <- rep(-1, 4)

# acceptance rate alpha
alpha <- pmvnorm(lower=lower, upper=upper, mean=rep(0,4), sigma=C)
alpha

# Gibbs sampler
X1 <- rtmvnorm(n=20000, mean = rep(0,4), sigma=C, lower=lower, upper=upper,
  algorithm="gibbs", burn.in.samples=100)
# Rejection sampling
X2 <- rtmvnorm(n=5000, mean = rep(0,4), sigma=C, lower=lower, upper=upper)

colMeans(X1)
colMeans(X2)

plot(density(X1[,1], from=lower[1], to=upper[1]), col="red", lwd=2,
     main="Kernel density estimates from random samples
     generated by Gibbs vs. Rejection sampling")
lines(density(X2[,1], from=lower[1], to=upper[1]), col="blue", lwd=2)
legend("topleft", legend=c("Gibbs Sampling", "Rejection Sampling"),
     col=c("red", "blue"), lwd=2, bty="n")

#####
#
# Example 3:
# Autocorrelation plot for Gibbs sampler
```

```

# with and without thinning
#
#####

sigma <- matrix(c(4,2,2,3), ncol=2)
X1 <- rtmvnorm(n=10000, mean=c(1,2), sigma=sigma, upper=c(1,0),
  algorithm="rejection")
acf(X1)
# no autocorrelation among random points

X2 <- rtmvnorm(n=10000, mean=c(1,2), sigma=sigma, upper=c(1,0),
  algorithm="gibbs")
acf(X2)
# exhibits autocorrelation among random points

X3 <- rtmvnorm(n=10000, mean=c(1,2), sigma=sigma, upper=c(1,0),
  algorithm="gibbs", thinning=2)
acf(X3)
# reduced autocorrelation among random points

plot(density(X1[,1], to=1))
lines(density(X2[,1], to=1), col="blue")
lines(density(X3[,1], to=1), col="red")

#####
#
# Example 4: Univariate case
#
#####

X <- rtmvnorm(100, mean=0, sigma=1, lower=-1, upper=1)

#####
#
# Example 5: Linear Constraints
#
#####

mean <- c(0, 0)
sigma <- matrix(c(10, 0,
                  0, 1), 2, 2)

# Linear Constraints
#
# a1 <= x1 + x2 <= b2
# a2 <= x1 - x2 <= b2
#
# [ a1 ] <= [ 1   1 ] [ x1 ] <= [b1]
# [ a2 ]   [ 1  -1 ] [ x2 ]   [b2]
a   <- c(-2, -2)
b   <- c( 2,  2)
D   <- matrix(c(1, 1,
                1, -1), 2, 2)

```

```

X <- rtmvnorm(n=10000, mean, sigma, lower=a, upper=b, D=D, algorithm="gibbsR")
plot(X, main="Gibbs sampling for multivariate normal
        with linear constraints according to Geweke (1991)")

# mark linear constraints as lines
for (i in 1:nrow(D)) {
  abline(a=a[i]/D[i, 2], b=-D[i,1]/D[i, 2], col="red")
  abline(a=b[i]/D[i, 2], b=-D[i,1]/D[i, 2], col="red")
}

#####
#
# Example 6: Using precision matrix H rather than sigma
#
#####

lower <- c(-1, -1)
upper <- c(1, 1)
mean <- c(0.5, 0.5)
sigma <- matrix(c(1, 0.8, 0.8, 1), 2, 2)
H <- solve(sigma)
D <- matrix(c(1, 1, 1, -1), 2, 2)
X <- rtmvnorm(n=1000, mean=mean, H=H, lower=lower, upper=upper, D=D, algorithm="gibbs")
plot(X, main="Gibbs sampling with precision matrix and linear constraints")

#####
#
# Example 7: Using sparse precision matrix H in high dimensions
#
#####

## Not run:
d <- 1000
I_d <- sparseMatrix(i=1:d, j=1:d, x=1)
W <- sparseMatrix(i=c(1:d, 1:(d-1)), j=c(1:d, (2:d)), x=0.5)
H <- t(I_d - 0.5 * W)
lower <- rep(0, d)
upper <- rep(2, d)

# Gibbs sampler generates n=100 draws in d=1000 dimensions
X <- rtmvnorm.sparseMatrix(n=100, mean = rep(0,d), H=H, lower=lower, upper=upper,
  burn.in.samples=100)
colMeans(X)
cov(X)

## End(Not run)

```

Description

This function generates random numbers from the truncated multivariate normal distribution with mean equal to mean and covariance matrix sigma and general linear constraints

$$lower \leq Dx \leq upper$$

with either rejection sampling or Gibbs sampling.

Usage

```
rtmvnorm2(n, mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  lower = rep(-Inf, length = length(mean)),
  upper = rep(Inf, length = length(mean)),
  D = diag(length(mean)),
  algorithm = c("gibbs", "gibbsR", "rejection"), ...)
```

Arguments

n	Number of random points to be sampled. Must be an integer ≥ 1 .
mean	Mean vector ($d \times 1$), default is <code>rep(0, length = ncol(x))</code> .
sigma	Covariance matrix ($d \times d$), default is <code>diag(ncol(x))</code> .
lower	Vector of lower truncation points ($r \times 1$), default is <code>rep(-Inf, length = length(mean))</code> .
upper	Vector of upper truncation points ($r \times 1$), default is <code>rep(Inf, length = length(mean))</code> .
D	Matrix for linear constraints ($r \times d$), defaults to diagonal matrix ($d \times d$), i.e. $r = d$.
algorithm	Method used, possible methods are the Fortan Gibbs sampler ("gibbs", default), the Gibbs sampler implementation in R ("gibbsR") and rejection sampling ("rejection")
...	additional parameters for Gibbs sampling, given to the internal method <code>rtmvnorm.gibbs()</code> , such as <code>burn.in.samples</code> , <code>start.value</code> and <code>thinning</code> , see details in rtmvnorm

Details

This method allows for $r > d$ linear constraints, whereas [rtmvnorm](#) requires a full-rank matrix D ($d \times d$) and can only handle $r \leq d$ constraints at the moment. The lower and upper bounds lower and upper are ($r \times 1$), the matrix D is ($r \times d$) and x is ($d \times 1$). The default case is $r = d$ and $D = I_d$.

Warning

This method will be merged with [rtmvnorm](#) in one of the next releases.

Author(s)

Stefan Wilhelm

See Also[rtmvnorm](#)**Examples**

```
## Not run:
#####
#
# Example 5a: Number of linear constraints r > dimension d
#
#####

# general linear restrictions  $a \leq Dx \leq b$  with  $x$  (d x 1);  $D$  (r x d);  $a, b$  (r x 1)

# Dimension d=2, r=3 linear constraints
#
#  $a_1 \leq x_1 + x_2 \leq b_1$ 
#  $a_2 \leq x_1 - x_2 \leq b_2$ 
#  $a_3 \leq 0.5x_1 - x_2 \leq b_3$ 
#
#  $\begin{bmatrix} a_1 \\ a_2 \\ a_3 \end{bmatrix} \leq \begin{bmatrix} 1 & 1 \\ 1 & -1 \\ 0.5 & -1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} \leq \begin{bmatrix} b_1 \\ b_2 \\ b_3 \end{bmatrix}$ 
#
D <- matrix(
  c( 1, 1,
    1, -1,
    0.5, -1), 3, 2, byrow=TRUE)
a <- c(0, 0, 0)
b <- c(1, 1, 1)

# mark linear constraints as lines
plot(NA, xlim=c(-0.5, 1.5), ylim=c(-1,1))
for (i in 1:3) {
  abline(a=a[i]/D[i, 2], b=-D[i,1]/D[i, 2], col="red")
  abline(a=b[i]/D[i, 2], b=-D[i,1]/D[i, 2], col="red")
}

### Gibbs sampling for general linear constraints  $a \leq Dx \leq b$ 
mean <- c(0, 0)
sigma <- matrix(c(1.0, 0.2,
  0.2, 1.0), 2, 2)
x0 <- c(0.5, 0.2) # Gibbs sampler start value
X <- rtmvnorm2(n=1000, mean, sigma, lower=a, upper=b, D, start.value=x0)

# show random points within simplex
points(X, pch=20, col="black")

## End(Not run)
```

rtmvt

*Sampling Random Numbers From The Truncated Multivariate Student
t Distribution*

Description

This function generates random numbers from the truncated multivariate Student-t distribution with mean equal to mean and covariance matrix sigma, lower and upper truncation points lower and upper with either rejection sampling or Gibbs sampling.

Usage

```
rtmvt(n, mean = rep(0, nrow(sigma)), sigma = diag(length(mean)), df = 1,
      lower = rep(-Inf, length = length(mean)),
      upper = rep(Inf, length = length(mean)),
      algorithm=c("rejection", "gibbs"), ...)
```

Arguments

n	Number of random points to be sampled. Must be an integer ≥ 1 .
mean	Mean vector, default is <code>rep(0, length = ncol(x))</code> .
sigma	Covariance matrix, default is <code>diag(ncol(x))</code> .
df	Degrees of freedom parameter (positive, may be non-integer)
lower	Vector of lower truncation points, default is <code>rep(-Inf, length = length(mean))</code> .
upper	Vector of upper truncation points, default is <code>rep(Inf, length = length(mean))</code> .
algorithm	Method used, possible methods are rejection sampling ("rejection", default) and the R Gibbs sampler ("gibbs").
...	additional parameters for Gibbs sampling, given to the internal method <code>rtmvt.gibbs()</code> , such as <code>burn.in.samples</code> , <code>start.value</code> and <code>thinning</code> , see details

Details

We sample $x \sim T(\mu, \Sigma, df)$ subject to the rectangular truncation $lower \leq x \leq upper$. Currently, two random number generation methods are implemented: rejection sampling and the Gibbs Sampler.

For rejection sampling `algorithm="rejection"`, we sample from `rmvt` and retain only samples inside the support region. The acceptance probability will be calculated with `pmvt`. `pmvt` does only accept integer degrees of freedom `df`. For non-integer `df`, `algorithm="rejection"` will throw an error, so please use `algorithm="gibbs"` instead.

The arguments to be passed along with `algorithm="gibbs"` are:

`burn.in.samples` number of samples in Gibbs sampling to be discarded as burn-in phase, must be non-negative.

`start.value` Start value (vector of length `length(mean)`) for the MCMC chain. If one is specified, it must lie inside the support region ($lower \leq start.value \leq upper$). If none is specified, the start value is taken componentwise as the finite lower or upper boundaries respectively, or zero if both boundaries are infinite. Defaults to `NULL`.

`thinning` Thinning factor for reducing autocorrelation of random points in Gibbs sampling. Must be an integer ≥ 1 . We create a Markov chain of length `(n*thinning)` and take only those samples `j=1:(n*thinning)` where `j %% thinning == 0` Defaults to 1 (no thinning of the chain).

Warning

The same warnings for the Gibbs sampler apply as for the method `rtmvnorm`.

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>, Manjunath B G <bgmanjunath@gmail.com>

References

Geweke, John F. (1991) Efficient Simulation from the Multivariate Normal and Student-t Distributions Subject to Linear Constraints. *Computer Science and Statistics. Proceedings of the 23rd Symposium on the Interface. Seattle Washington, April 21-24, 1991*, pp. 571–578 An earlier version of this paper is available at https://www.researchgate.net/publication/2335219_Efficient_Simulation_from_the_Multivariate_Normal_and_Student-t_Distributions_Subject_to_Linear_Constraints_and_the_Evaluation_of_Constraint_Probabilities

Examples

```
#####
#
# Example 1
#
#####

# Draw from multi-t distribution without truncation
X1 <- rtmvt(n=10000, mean=rep(0, 2), df=2)
X2 <- rtmvt(n=10000, mean=rep(0, 2), df=2, lower=c(-1,-1), upper=c(1,1))

#####
#
# Example 2
#
#####

df = 2
mu = c(1,1,1)
sigma = matrix(c( 1, 0.5, 0.5,
                  0.5, 1, 0.5,
                  0.5, 0.5, 1), 3, 3)
lower = c(-2,-2,-2)
upper = c(2, 2, 2)
```

```

# Rejection sampling
X1 <- rtmvt(n=10000, mu, sigma, df, lower, upper)

# Gibbs sampling without thinning
X2 <- rtmvt(n=10000, mu, sigma, df, lower, upper,
  algorithm="gibbs")

# Gibbs sampling with thinning
X3 <- rtmvt(n=10000, mu, sigma, df, lower, upper,
  algorithm="gibbs", thinning=2)

plot(density(X1[,1], from=lower[1], to=upper[1]), col="red", lwd=2,
  main="Gibbs vs. Rejection")
lines(density(X2[,1], from=lower[1], to=upper[1]), col="blue", lwd=2)
legend("topleft", legend=c("Rejection Sampling", "Gibbs Sampling"),
  col=c("red", "blue"), lwd=2)

acf(X1) # no autocorrelation in Rejection sampling
acf(X2) # strong autocorrelation of Gibbs samples
acf(X3) # reduced autocorrelation of Gibbs samples after thinning

```

tmvnorm

Truncated Multivariate Normal Density

Description

This function provides the joint density function for the truncated multivariate normal distribution with mean equal to mean and covariance matrix sigma, lower and upper truncation points lower and upper. For convenience, it furthermore serves as a wrapper function for the one-dimensional and bivariate marginal densities dtmvnorm.marginal() and dtmvnorm.marginal2() respectively when invoked with the margin argument.

Usage

```

dtmvnorm(x, mean = rep(0, nrow(sigma)),
  sigma = diag(length(mean)),
  lower=rep(-Inf, length = length(mean)),
  upper=rep( Inf, length = length(mean)),
  log=FALSE,
  margin=NULL)

```

Arguments

x	Vector or matrix of quantiles. If x is a matrix, each row is taken to be a quantile.
mean	Mean vector, default is rep(0, nrow(sigma)).
sigma	Covariance matrix, default is diag(length(mean)).
lower	Vector of lower truncation points, default is rep(-Inf, length = length(mean)).
upper	Vector of upper truncation points, default is rep(Inf, length = length(mean)).

log	Logical; if TRUE, densities d are given as log(d).
margin	if NULL then the joint density is computed (the default), if MARGIN=1 then the one-dimensional marginal density in variate q ($q = 1 \dots \text{length}(\text{mean})$) is returned, if MARGIN=c(q,r) then the bivariate marginal density in variates q and r for $q, r = 1 \dots \text{length}(\text{mean})$ and $q \neq r$ is returned.

Details

The computation of truncated multivariate normal probabilities and densities is done using conditional probabilities from the standard/untruncated multivariate normal distribution. So we refer to the documentation of the mvtnorm package and the methodology is described in Genz (1992, 1993).

Author(s)

Stefan Wilhelm <Stefan.Wilhelm@financial.com>

References

- Genz, A. (1992). Numerical computation of multivariate normal probabilities. *Journal of Computational and Graphical Statistics*, **1**, 141–150
- Genz, A. (1993). Comparison of methods for the computation of multivariate normal probabilities. *Computing Science and Statistics*, **25**, 400–405
- Johnson, N./Kotz, S. (1970). Distributions in Statistics: Continuous Multivariate Distributions Wiley & Sons, pp. 70–73
- Horrace, W. (2005). Some Results on the Multivariate Truncated Normal Distribution. *Journal of Multivariate Analysis*, **94**, 209–221

See Also

[ptmvnorm](#), [pmvnorm](#), [rmvnorm](#), [dmvnorm](#), [dtmvnorm.marginal](#) and [dtmvnorm.marginal2](#) for marginal density functions

Examples

```
dtmvnorm(x=c(0,0), mean=c(1,1), upper=c(0,0))

#####
#
# Example 1:
# truncated multivariate normal density
#
#####

x1<-seq(-2, 3, by=0.1)
x2<-seq(-2, 3, by=0.1)

density<-function(x)
{
  sigma=matrix(c(1, -0.5, -0.5, 1), 2, 2)
```

```

    z=dtmvnorm(x, mean=c(0,0), sigma=sigma, lower=c(-1,-1))
  }
  z

fgrid <- function(x, y, f)
{
  z <- matrix(nrow=length(x), ncol=length(y))
  for(m in 1:length(x)){
    for(n in 1:length(y)){
      z[m,n] <- f(c(x[m], y[n]))
    }
  }
  z
}

# compute density d for grid
d=fgrid(x1, x2, density)

# plot density as contourplot
contour(x1, x2, d, nlevels=5, main="Truncated Multivariate Normal Density",
  xlab=expression(x[1]), ylab=expression(x[2]))
abline(v=-1, lty=3, lwd=2)
abline(h=-1, lty=3, lwd=2)

#####
#
# Example 2:
# generation of random numbers
# from a truncated multivariate normal distribution
#
#####

sigma <- matrix(c(4,2,2,3), ncol=2)
x <- rtmvnorm(n=500, mean=c(1,2), sigma=sigma, upper=c(1,0))
plot(x, main="samples from truncated bivariate normal distribution",
  xlim=c(-6,6), ylim=c(-6,6),
  xlab=expression(x[1]), ylab=expression(x[2]))
abline(v=1, lty=3, lwd=2, col="gray")
abline(h=0, lty=3, lwd=2, col="gray")

```

Index

* distribution

- dtmnorm.marginal, 2
- dtmnorm.marginal2, 5
- dtmvt, 7
- mtmnorm, 14
- ptmnorm, 15
- ptmvtnorm.marginal, 18
- qtmnorm-marginal, 20
- rtmnorm, 21
- rtmnorm2, 27
- rtmvt, 30
- tmvnorm, 32

* math

- ptmvt, 17

* multivariate

- dtmnorm.marginal, 2
- dtmnorm.marginal2, 5
- dtmvt, 7
- mtmnorm, 14
- ptmnorm, 15
- ptmvt, 17
- ptmvtnorm.marginal, 18
- qtmnorm-marginal, 20
- rtmnorm, 21
- rtmnorm2, 27
- rtmvt, 30
- tmvnorm, 32

acf, 24

dmvnorm, 24, 33

dmvt, 8

dtmnorm (tmvnorm), 32

dtmnorm.marginal, 2, 10, 33

dtmnorm.marginal2, 5, 33

dtmvt, 7

gmm, 10, 11

gmm.tmvnorm, 9

matrix, 23

mle, 12, 13

mle.tmvnorm, 11

moments (mtmnorm), 14

mtmnorm, 10, 14

optim, 12

pmvnorm, 5, 14, 21, 24, 33

pmvt, 8, 30

ptmnorm, 15, 21, 24, 33

ptmnorm.marginal (ptmvtnorm.marginal), 18

ptmvt, 8, 17

ptmvt.marginal (ptmvtnorm.marginal), 18

ptmvtnorm.marginal, 18

qtmnorm-marginal, 20

qtmnorm.marginal (qtmnorm-marginal), 20

rmvnorm, 22, 24, 33

rmvt, 8, 30

rtmnorm, 21, 28, 29, 31

rtmnorm2, 27

rtmvt, 8, 30

sparseMatrix, 23

tmvnorm, 32

uniroot, 20, 21