## Package 'logconcens'

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Type Package

Title	Maximum Likelihood Estimation of a Log-Concave Density Based on
	Censored Data

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**Depends** R (>= 2.15.0)

Imports grDevices, graphics, stats, utils

Suggests logcondens

**Description** Based on right or interval censored data, compute the maximum likelihood estimator of a (sub)probability density under the assumption that it is log-concave. For further information see Duembgen, Rufibach and Schuhmacher (2014) <doi:10.1214/14-EJS930>.

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## **R** topics documented:

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logconcens-package

#### Description

Based on independent intervals  $X_i = [L_i, R_i]$ , where  $-\infty < L_i \le R_i \le \infty$ , compute the maximum likelihood estimator of a (sub)probability density under the assumption that it is log-concave. For further information see Duembgen, Rufibach, and Schuhmacher (2013, preprint).

#### Details

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The main function is logcon, which offers computation of the MLE for many types of censored and also exact data. Various parameters can be set that allow for fine control of the underlying EM algorithm in "difficult" situations. An object of type lcdensity is returned, for which plot, print, and summary methods are available. There is also a function loglike for computing the log-likelihood of a lcdensity object.

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#### References

Duembgen, L., Rufibach, K., and Schuhmacher, D. (2014). Maximum-likelihood estimation of a log-concave density based on censored data. Electronic Journal of Statistics, 8(1), 1405-1437. doi:10.1214/14EJS930

## Examples

- ## Simple examples with simulated data.
- ## For more detailed examples see the help for the function logcon.

## exact data
set.seed(10)

#### cure.profile

```
x <- rnorm(100)</pre>
res <- logcon(x)</pre>
## Not run:
plot(res)
xi <- seq(-3,3,0.05)
lines(xi,log(dnorm(xi)))
## End(Not run)
## interval censored data
x <- rgamma(50,3,1)
x <- cbind(x,x+rexp(50,1))</pre>
plotint(x)
res <- logcon(x)</pre>
## Not run: plot(res, type="CDF")
## right censored data with mass at infinity
set.seed(11)
x <- rgamma(50,3,1)
x <- cbind(x,ifelse(rexp(50,1/3) < x,Inf,x))</pre>
plotint(x)
res <- logcon(x,adapt.p0=TRUE)</pre>
## Not run: plot(res, type="survival")
## rounded/binned data
set.seed(12)
x <- round(rnorm(100))</pre>
x <- cbind(x-0.5,x+0.5)</pre>
plotint(x)
res <- logcon(x)</pre>
## Not run:
plot(res)
xi <- seq(-3,3,0.05)
```

```
cure.profile
```

lines(xi,log(dnorm(xi)))

## End(Not run)

```
Evaluate the Profile Log-Likelihood on a Grid of p_0-Values
```

## Description

For each of a series of values for the cure parameter  $p_0$  run the function logcon and evaluate the (normalized) log-likelihood at  $(\phi, p_0)$ , where  $\phi$  is the log subprobability density returned by logcon. This serves for (approximate) joint likelihood maximization in  $(\phi, p_0)$ .

#### Usage

#### Arguments

х	a two-column matrix of $n \ge 2$ rows containing the data intervals.
p0grid	a vector of values $p_0$ for which the profile log-likelihood is to be evaluated.
knot.prec,	reduce, control
	arguments passed to the function logcon.

## Value

A list containing the following values:

p0hat	the element in p0grid that maximizes the profile likelihood (in the very unlikely case of ties, only the smallest such element is returned).
status	the vector of (normalized) profile log-likelihood values for the elements of $p@grid$ .

## Note

For a large p0grid-vector (fine grid) computations may take a long time. Consider using the option adapt.p0 in the function logcon for a much faster method of joint likelihood maximization in  $(\phi, p_0)$ .

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## See Also

### logcon, loglike

## Examples

```
## The example from the logconcens-package help page:
set.seed(11)
x <- rgamma(50,3,1)
x <- cbind(x,ifelse(rexp(50,1/3) < x,Inf,x))</pre>
```

```
## Not run:
plotint(x)
progrid <- seq(0.1,0.6,0.025)
prores <- cure.profile(x, progrid)
plot(progrid, prores$loglike)
prores$p0hat
res <- logcon(x, p0=prores$p0hat)
plot(res, type="survival")
```

## End(Not run)

lc.control

## Description

Allows to set the control parameters for the more technical aspects of the function logcon and provides default values for any parameters that are not set.

## Usage

## Arguments

maxiter	the maximal number of iterations in the main EM algorithm. Default is 49 rather than 50, because this goes well with plotting in case of show = TRUE.
move.prec	a real number giving the threshold for the $L_1$ -distance between densities in sub- sequent steps below which the algorithm is stopped.
domind11, domin	d2r
	index numbers in the vector of sorted interval endpoints that specify the left and right boundary of the maximal domain to be considered by the algorithm; see the details section of the help for logcon. The indices are counted from the left and from the right, respectively. So the default values of domind11 = 1 and domind2r = 1 mean that the largest possible domain is used.
force.inf	logical. For experimental use only. Should the domain interval be forced to be right-infinite (if there is a right-infinite data interval)?
red.thresh	a real number indicating the threshold below which the boundary integrals are considered too small; see the details section of the help for logcon. There is a sensible default, which depends on check.red.
check.red	logical. If a boundary integral is deemed too small, should the derivative of the augmented log-likelihood be checked to confirm the domain reduction.
addpoints	logical. Should extra exact observations be added to the data at the left- and rightmost finite interval endpoints to prevent domain reduction? These observations obtain a small weight $< 1$ as compared to the weight of 1 for all the other observation intervals. The weight is specified by addeps.
addeps	a positive real number. If NULL, a default value of $1/n^2$ is computed where $n$ is the number of observation intervals. See addpoints.
preweights	a vector of weights for the observation intervals. Defaults to rep(1,n).
minw	a positive real number. This gives another way for preventing domain reduction. Instead of adding observations the weights for the internal active set algorithm are kept at or above minw at the boundary of the domain.

show	logical. Should progress of the algorithm be plotted? Warning: if TRUE, this may open many new graphics devices in case of complicated data sets.
verbose	logical. Should additional information about the progress of the algorithm be printed? This mainly prints quantities important for the decision to reduce the domain of the function and about the progress of the EM algorithm.

## Details

For further explanations about the algorithm see the help for logcon. In summary:

maxiter and move.prec provide stopping criteria for the EM algorithm.

domind11, domind2r, force.inf, red.thresh, and check.red control aspects related to domain reduction.

addpoints, addeps, preweights, winw allow for reweighing of data interval, mainly for increasing numerical stability by preventing domain reduction.

show and verbose give illustrations and background information of the run of the algorithm.

#### Value

A list with all of the above components set to their (specified or default) value.

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#### See Also

logcon

## Examples

## See the examples for logcon

## Description

Plot, print, and summary methods for objects of class lcdensity.

## lcdensity-methods

## Usage

## Arguments

x,object	objects of class lcdensity, as returned by logcon.
type	the type of plot to be produced.
sloperange	logical. In cases where the cure parameter / the right-hand side slope of the log-subdensity $\phi$ is not unique, should grey area be drawn indicating the set of possible right-hand slopes?
kinklines	logical. Should vertical lines be drawn at the kinks of the log-subdensity $\phi$ ?
kinkpoints	logical. Should fat points be plotted at the kinks of the log-subdensity $\phi$ ?
xlim, ylim	numeric vectors of length 2, giving the x and y coordinates ranges.
	further arguments passed to plot.default. Depending on the argument this may or may not work in the intended way.

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## See Also

## plotint

## Examples

## See the examples for logcon

## Description

Based on independent intervals  $X_i = [L_i, R_i]$ , where  $-\infty < L_i \le R_i \le \infty$ , compute the maximum likelihood estimator of a (sub)probability density  $\phi$  and the remaining mass  $p_0$  at infinity (also known as *cure parameter*) under the assumption that the former is log-concave. Computation is based on an EM algorithm. For further information see Duembgen, Rufibach, and Schuhmacher (2013, preprint).

## Usage

logconcure(x, p0=0, knot.prec=IQR(x[x<Inf])/75, reduce=TRUE, control=lc.control())</pre>

#### Arguments

x	a two-column matrix of $n \ge 2$ rows containing the data intervals, or a vector of length $n \ge 2$ containing the exact data points.
adapt.p0	logical. Should the algorithm be allowed to adapt $p_0$ ? In this case an alternating maximization procedure is used that is <i>believed</i> to always yield a joint maximizer $(\hat{\phi}, \hat{p_0})$ . For the much slower (but maybe safer) profile likelihood maximization method, see the function cure.profile.
9q	a number from 0 to 1 specifying the mass at infinity. If the algorithm is allowed to adapt $p_0$ , this argument only specifies the starting value. Otherwise it is assumed that the true cure parameter $p_0$ is equal to this number. In particular, for the default setting of 0, a proper probability density $\phi$ is estimated.
knot.prec	the maximal distance between two consecutive grid points, where knots (points at which the resulting log-subdensity $\phi$ may change slope) can be positioned. See details.
reduce	logical. Should the domain of the (sub)density be reduced whenever the mass at the left or the right boundary becomes too small?
control	a list of control parameters for the more technical aspects of the algorithm; usually the result of a call to lc.control.

## Details

Based on the data intervals  $X_i = [L_i, R_i]$  described above, function logcon computes a concave, piecewise linear function  $\phi$  and a probability  $p_0$  which satisfy  $\int \exp \phi(x) dx = 1 - p_0$  and jointly maximize the (normalized) log-likelihood.

$$\ell(\phi, p_0) = \frac{1}{n} \sum_{i=1}^n \left[ 1\{L_i = R_i\}\phi(X_i) + 1\{L_i < R_i\} \log\left(\int_{L_i}^{R_i} \exp\phi(x) \, dx + 1\{R_i = \infty\}p_0\right) \right],$$

If x is a two-column matrix, it is assumed to contain the left and right interval endpoints in the correct order. Intervals may have length zero (both endpoints equal) or be unbounded to the right (right endpoint is Inf). Computation is based on an EM algorithm, where the M-step uses an active set algorithm for computing the log-concave MLE for exact data with weights. The active set algorithm was described in Duembgen, Huesler, and Rufibach (2007) and Duembgen and Rufibach (2011) and is available in the R package logcondens. It has been re-implemented in C for the current package because of speed requirements. The whole algorithm for censored data has been indicated in Duembgen, Huesler, and Rufibach (2007) and was elaborated in Duembgen, Schuhmacher, and Rufibach (2013, preprint).

If x is a vector argument, it is assumed to contain the exact data points. In this case the active set algorithm is accessed directly.

In order to obtain a finite dimensional optimization problem the (supposed) domain of  $\phi$  is subdivided by a grid. Stretches between interval endpoints where for theoretical reasons no knots (points where the slope of  $\phi$  changes) can lie are left out. The argument kink.prec gives the maximal distance we allow between consecutive grid points in stretches where knots can lie. Say plotint(x) to see the grid.

The EM algorithm works only for fixed dimensionality of the problem, but the domain of the function  $\phi$  is not a priori known. Therefore there is an outer loop starting with the largest possible domain, given by the minimal and maximal endpoints of all the intervals, and decreasing the domain as soon as the EM steps let  $\phi$  become very small towards the boundary. "Very small" means that the integral of exp  $\circ \phi$  over the first or last stretch between interval endpoints within the current domain falls below a certain threshold red.thresh, which can be controlled via lc.control.

Domain reduction tends to be rather conservative. If the computed solution has a suspiciously steep slope at any of the domain boundaries, the recommended strategy is to enforce a smaller domain by increasing the parameters domind11 and/or domind2r via lc.control. The function loglike may be used to compare the (normalized) log-likelihoods of the results.

logConCens is an alias for logcon. It is introduced to provide unified naming with the main functions in the packages logcondens and logcondiscr.

logconcure is the same as logcon with adapt.p0 = TRUE fixed.

#### Value

An object of class lcdensity for which reasonable plot, print, and summary methods are available.

If the argument x is a two-column matrix (censored data case), such an object has the following components.

basedOn the string "censored" for the type of data the solution is based on.

status currently only 0 if the algorithm converged; and 1 otherwise. Note that in most cases even with status 1 the returned solution is very close to the truth. The 1 is often due to the fact that the termination criterion is not so well balanced yet.

the data entered.
the ordered vector of different interval endpoints.
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the indices of the tau-element at which the domain of the MLE $\phi$ starts/ends.
the grid vector. tau[domind1:domind2] augmented by points of subdivision.
0-1 value. For the finite elements of tplus a 1 if $\phi$ has a knot at this position, 0 otherwise.
the vector of $\phi$ -values at the finite elements of tplus.
if $\sup(\operatorname{dom}(\phi)) = \infty$ , the slope of $\phi$ after the last knot. Otherwise $-\infty$ .
a vector of length 2 specifying a range of possible values for phislr. This is for the (rather rare) situations that mass may be shifted between the interval from the rightmost tau-point to infinity and the cure parameter without changing the likelihood. Otherwise phislr.range is NA.
the cure parameter. Either the original argument $p0$ if adapt.p0 was FALSE, otheriwse the estimated cure parameter obtained by the alternating maximization procedure.
a vector of length 2 specifying a range of possible values for cure or NA. See phislr.range.
the vector of values of the distribution function $F$ of $\exp\circ\phi$ at the finite elements of tplus.
the computed value of $\lim_{t\to\infty} F(t)$ .

#### Note

If x is a vector, this function does the same as the function logConDens in the package logcondens. The latter package offers additional features such as grid-based computation with weights (for high numerical stability) and smoothing of the estimator, as well as nicer plotting. For exact data we recommend using logConDens for everyday data analysis. logcon with a vector argument is to be preferred if time is of the essence (for data sets with several thousands of points or repeated optimization in iterative algorithms) or if an additional slope functionality is required.

Two other helpful packages for log-concave density estimation based on exact data are logcondiscr for estimating a discrete distribution and LogConcDEAD for estimating a multivariate continuous distribution.

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## References

Duembgen, L., Huesler, A., and Rufibach, K. (2007). Active set and EM algorithms for log-concave densities based on complete and censored data. Technical Report 61. IMSV, University of Bern. https://arxiv.org/abs/0707.4643

Duembgen, L. and Rufibach, K., (2011). logcondens: Computations Related to Univariate Log-Concave Density Estimation. Journal of Statistical Software, 39(6), 1-28. doi:10.18637/jss.v039.i06

Duembgen, L., Rufibach, K., and Schuhmacher, D. (2014). Maximum-likelihood estimation of a log-concave density based on censored data. Electronic Journal of Statistics, 8(1), 1405-1437. doi:10.1214/14EJS930

#### See Also

lc.control,lcdensity-methods,loglike

#### Examples

```
# A function for artificially censoring exact data
censor <- function(y, timemat) {</pre>
  tm <- cbind(0,timemat,Inf)</pre>
  n <- length(y)
  res <- sapply(1:n, function(i){</pre>
            return( c( max(tm[i,][tm[i,] < y[i]]), min(tm[i,][tm[i,] >= y[i]]) ) ) } )
  return(t(res))
}
# ------
# interval censored data
# _____
set.seed(20)
n <- 100
# generate exact data:
y < - rgamma(n,3)
# generate matrix of inspection times:
itimes <- matrix(rexp(10*n),n,10)</pre>
itimes <- t(apply(itimes,1,cumsum))</pre>
# transform exact data to interval data
x <- censor(y, itimes)</pre>
# plot both
plotint(x, imarks=y)
# Compute censored log-concave MLE
# (assuming only the censored data is available to us)
res <- logcon(x)</pre>
plot(res)
# Compare it to the log-concave MLE for the exact data
# and to the true Gamma(3,1) log-density
res.ex <- logcon(y)</pre>
lines(res.ex$x, res.ex$phi, lwd=2.5, lty=2)
xi <- seq(0,14,0.05)
lines(xi,log(dgamma(xi,3,1)), col=3, lwd=2)
```

```
# ------
# censored data with cure
# ______
## Not run:
set.seed(21)
n <- 100
# generate exact data:
y <- rgamma(n,3)
cured <- as.logical(rbinom(n,1,0.3))</pre>
y[cured] <- Inf
# generate matrix of inspection times:
itimes <- matrix(rexp(6*n),n,6)</pre>
itimes <- t(apply(itimes,1,cumsum))</pre>
# transform exact data to interval data
x <- censor(y, itimes)</pre>
# plot both
plotint(x, imarks=y)
# Compute censored log-concave MLE including cure parameter
# (assuming only the censored data is available to us)
res <- logcon(x, adapt.p0=TRUE)</pre>
plot(res)
# There is a trade-off between right-hand slope and cure parameter here
# (seen by the grey area on the right), but the margin is very small:
res$cure.range
# Compare the corresponding CDF to the true CDF
plot(res, type="CDF")
xi <- seq(0,14,0.05)
lines(xi,0.7*pgamma(xi,3,1), col=3, lwd=2)
# Note that the trade-off for the right-hand slope is not visible anymore
# (in terms of the CDF the effect is too small)
## End(Not run)
# ------
# real right censored data with cure
#
 -----
# Look at data set ovarian from package survival
# Gives survival times in days for 26 patients with advanced ovarian carcinoma,
# ignoring the covariates
# Bring data to right format and plot it
## Not run:
library(survival)
data(ovarian)
sobj <- Surv(ovarian$futime, ovarian$fustat)</pre>
```

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```
x <- cbind(sobj[,1], ifelse(as.logical(sobj[,2]),sobj[,1],Inf))</pre>
plotint(x)
# Compute censored log-concave MLE including cure parameter
res <- logcon(x, adapt.p0=TRUE)</pre>
# Compare the corresponding survival function to the Kaplan-Meier estimator
plot(res, type="survival")
res.km <- survfit(sobj ~ 1)</pre>
lines(res.km, lwd=1.5)
## End(Not run)
# ------
# current status data
# ------
## Not run:
set.seed(22)
n <- 200
# generate exact data
y <- rweibull(n,2,1)</pre>
# generate vector of inspection times
itime <- matrix(rexp(n),n,1)</pre>
# transform exact data to interval data
x <- censor(y, itime)</pre>
# plot both
plotint(x, imarks=y)
# Compute censored log-concave MLE
# (assuming only the censored data is available to us)
res <- logcon(x)</pre>
plot(res, type="CDF")
# Compare it to the true Weibull(2,1) c.d.f.
xi <- seq(0,3,0.05)
lines(xi,pweibull(xi,2,1), col=3, lwd=2)
## End(Not run)
# ------
# rounded/binned data
# _____
## Not run:
set.seed(23)
n <- 100
# generate data in [0,1] rounded to one digit
y <- round(rbeta(n,2,3),1)</pre>
# bring data to right format and plot it
x <- cbind(y-0.05,y+0.05)</pre>
```

loglike

```
plotint(x)
# Compute censored log-concave MLE
res <- logcon(x)
plot(res, type="density", xlim=c(0,1))
# Compare it to the true Beta(2,3) density
xi <- seq(0,1,0.005)
lines(xi,dbeta(xi,2,3), col=3, lwd=2)
# The peaks in the estimated density are often considered unsatisfactory
# However, they are barely noticeable in the c.d.f.
plot(res, type="CDF", xlim=c(0,1))
lines(xi,pbeta(xi,2,3), col=3, lwd=2)
# To get rid of them in the density apply the smoothing
# proposed in the package logcondens (to be implemented here)
## End(Not run)</pre>
```

```
loglike
```

Compute Log-Likelihood for an Object of Class lcdensity

#### Description

Compute the (normalized) log-likelihood for an object of class lcdensity as described in the details section for the function logcon. The main use of this function is for comparing different results from logcon based on different (starting) domains.

#### Usage

loglike(lcd)

#### Arguments

lcd

an object of class lcdensity

## Value

A single numeric value, the (normalized) log-likelihood.

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## See Also

logcon

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## plotint

## Examples

```
x <- matrix(c(0,0.5,0.5,1,1,2,3,3),4,2)
res <- logcon(x)
loglike(res)</pre>
```

plotint

Plot Censored Data

## Description

Plot a graphical representation of censored data specified by a two-column matrix of left and right interval endpoints. The grid of potential knots used by logcon is also shown.

#### Usage

```
plotint(x, knot.prec = IQR(x[x<Inf])/75, imarks = NULL)</pre>
```

## Arguments

х	a two-column matrix of left and right endpoints of data intervals.
knot.prec	the maximal distance between two consecutive grid points in the depiction of the grid used by logcon.
imarks	an optional vector of "spots" to be marked by 'x' for the intervals.

## Value

Used for the side effect.

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#### See Also

plot.lcdensity

## Examples

## See the examples for logcon

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