Package ‘ashr’

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Title Methods for Adaptive Shrinkage, using Empirical Bayes
Description The R package ‘ashr’ implements an Empirical Bayes approach for large-scale hypothesis testing and false discovery rate (FDR) estimation based on the methods proposed in M. Stephens, 2016, ‘False discovery rates: a new deal’, <DOI:10.1093/biostatistics/kxw041>. These methods can be applied whenever two sets of summary statistics—estimated effects and standard errors—are available, just as ‘qvalue’ can be applied to previously computed p-values. Two main interfaces are provided: ash(), which is more user-friendly; and ash.workhorse(), which has more options and is geared toward advanced users. The ash() and ash.workhorse() also provides a flexible modeling interface that can accommodate a variety of likelihoods (e.g., normal, Poisson) and mixture priors (e.g., uniform, normal).

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Adaptive Shrinkage

Description

Implements Empirical Bayes shrinkage and false discovery rate methods based on unimodal prior distributions.

Usage

ash(betahat, sebetahat, mixcompdist = c("uniform", "halfuniform", "normal", "+uniform", "-uniform", "halfnormal"), df = NULL, ...)

ash.workhorse(betahat, sebetahat, method = c("fdr", "shrink"),
mixcompdist = c("uniform", "halfuniform", "normal", "+uniform", "-uniform", "halfnormal"), optim.method = c("mixSQP", "mixIP", "cxxMixSquarem", "mixEM", "mixVBEM", "w_mixEM"), df = NULL,
nullweight = 10, pointmass = TRUE, prior = c("nullbiased", "uniform", "unit"), mixsd = NULL, gridmult = sqrt(2),
outputlevel = 2, g = NULL, fixg = FALSE, mode = 0, alpha = 0,
grange = c(-Inf, Inf), control = list(), lik = NULL,
weights = NULL, pi_thresh = 1e-10)

Arguments

betahat a p vector of estimates
sebetahat a p vector of corresponding standard errors
distribution of components in mixture used to represent the family G. Depending on the choice of mixture component, the family G becomes more or less flexible. Options are:

- **uniform** G is (approximately) any symmetric unimodal distribution
- **normal** G is (approximately) any scale mixture of normals
- **halfuniform** G is (approximately) any unimodal distribution
- **+uniform** G is (approximately) any unimodal distribution with support constrained to be greater than the mode.
- **-uniform** G is (approximately) any unimodal distribution with support constrained to be less than the mode.
- **halfnormal** G is (approximately) any scale mixture of truncated normals where the normals are truncated at the mode.

If you are happy to assume a symmetric distribution for effects, you can use "uniform" or "normal". If you believe your effects may be asymmetric, use "halfuniform" or "halfnormal". If you want to allow only positive/negative effects use "+uniform"/"-uniform". The use of "normal" and "halfnormal" is permitted only if df=NULL.

- **df** appropriate degrees of freedom for (t) distribution of (betahat-beta)/sebetahat; default is NULL which is actually treated as infinity (Gaussian)
- **method** specifies how ash is to be run. Can be "shrinkage" (if main aim is shrinkage) or "fdr" (if main aim is to assess false discovery rate or false sign rate (fsr)). This is simply a convenient way to specify certain combinations of parameters: "shrinkage" sets pointmass=FALSE and prior="uniform"; "fdr" sets pointmass=TRUE and prior="nullbiased".
- **optmethod** specifies the function implementing an optimization method.
- **nullweight** scalar, the weight put on the prior under "nullbiased" specification, see prior
- **pointmass** Logical, indicating whether to use a point mass at zero as one of components for a mixture distribution.
- **prior** string, or numeric vector indicating Dirichlet prior on mixture proportions (defaults to "uniform", or (1,1,...,1); also can be "nullbiased" (nullweight,1,...,1) to put more weight on first component), or "unit" (1/K,...,1/K) [for optmethod=mixVBEM version only].
- **mixsd** Vector of standard deviations for underlying mixture components.
- **gridmult** the multiplier by which the default grid values for mixsd differ by one another. (Smaller values produce finer grids.)
- **outputlevel** Determines amount of output. There are several numeric options: 0 = just fitted g; 1 = also PosteriorMean and PosteriorSD; 2 = everything usually needed; 3 = also include results of mixture fitting procedure (including matrix of log-likelihoods used to fit mixture). 4 and 5 are reserved for outputting additional data required by the (in-development) flashr package. The user can also specify the output they require in detail (see Examples).
The prior distribution for beta. Usually this is unspecified (NULL) and estimated from the data. However, it can be used in conjunction with fixg=TRUE to specify the g to use (e.g. useful in simulations to do computations with the "true" g). Or, if g is specified but fixg=FALSE, the g specifies the initial value of g used before optimization, (which also implicitly specifies mixcompdist).

fixg
If TRUE, don’t estimate g but use the specified g - useful for computations under the "true" g in simulations.

mode
either numeric (indicating mode of g) or string "estimate", to indicate mode should be estimated, or a two dimension numeric vector to indicate the interval to be searched for the mode.

alpha
Numeric value of alpha parameter in the model.

grange
Two dimension numeric vector indicating the left and right limit of g. Default is c(-Inf, Inf).

control
A list of control parameters passed to optmethod.

lik
Contains details of the likelihood used; for general ash. Currently, the following choices are allowed: normal (see function lik_normal()); binomial likelihood (see function lik_binom); likelihood based on logF error distribution (see function lik_logF); mixture of normals likelihood (see function lik_normalmix); and Poisson likelihood (see function lik_pois).

weights
a vector of weights for observations; use with optmethod = "w_mixEM"; this is currently beta-functionality.

pi_thresh
a threshold below which to prune out mixture components before computing summaries (speeds up computation since empirically many components are usually assigned negligible weight). The current implementation still returns the full fitted distribution; this only affects the posterior summaries.

Details

The ash function provides a number of ways to perform Empirical Bayes shrinkage estimation and false discovery rate estimation. The main assumption is that the underlying distribution of effects is unimodal. Novice users are recommended to start with the examples provided below.

In the simplest case the inputs to ash are a vector of estimates (betahat) and their corresponding standard errors (sebetahat), and degrees of freedom (df). The method assumes that for some (unknown) "true" vector of effects beta, the statistic (betahat[j]-beta[j])/sebetahat[j] has a $t$ distribution on $df$ degrees of freedom. (The default of df=NULL assumes a normal distribution instead of a t.)

By default the method estimates the vector beta under the assumption that beta ~ g for a distribution g in G, where G is some unimodal family of distributions to be specified (see parameter mixcompdist). By default is to assume the mode is 0, and this is suitable for settings where you are interested in testing which beta[j] are non-zero. To estimate the mode see parameter mode.

As is standard in empirical Bayes methods, the fitting proceeds in two stages: i) estimate g by maximizing a (possibly penalized) likelihood; ii) compute the posterior distribution for each beta[j] | betahat[j].sebetahat[j] using the estimated g as the prior distribution.

A more general case allows that beta[j]/sebetahat[j]^alpha | sebetahat[j] ~ g.
Value

ash returns an object of class "ash", a list with some or all of the following elements (determined by outputlevel)

- **fitted_g**: fitted mixture
- **loglik**: \( \log P(D|\text{fitted}_g) \)
- **logLR**: \( \log[P(D|\text{fitted}_g)/P(D|\beta==0)] \)
- **result**: A dataframe whose columns are:
  - **NegativeProb**: A vector of posterior probability that \( \beta \) is negative.
  - **PositiveProb**: A vector of posterior probability that \( \beta \) is positive.
  - **lfsr**: A vector of estimated local false sign rate.
  - **lfdr**: A vector of estimated local false discovery rate.
  - **qvalue**: A vector of q values.
  - **svalue**: A vector of s values.
  - **PosteriorMean**: A vector consisting the posterior mean of \( \beta \) from the mixture.
  - **PosteriorSD**: A vector consisting the corresponding posterior standard deviation.

**call**

A call in which all of the specified arguments are specified by their full names

**data**

A list containing details of the data and models used (mostly for internal use)

**fit_details**

A list containing results of mixture optimization, and matrix of component log-likelihoods used in this optimization

Functions

- `ash.workhorse`: Adaptive Shrinkage with full set of options.

See Also

`ashci` for computation of credible intervals after getting the ash object return by `ash()`

Examples

```r
beta = c(rep(0,100),rnorm(100))
sebetahat = abs(rnorm(200,0,1))
betahat = rnorm(200,beta,sebetahat)
beta.ash = ash(betahat, sebetahat)
names(beta.ash)
head(beta.ash$result) # the main dataframe of results
head(get_pm(beta.ash)) # get_pm returns posterior mean
head(get_lfsr(beta.ash)) # get_lfsr returns the local false sign rate
graphics::plot(betahat,get_pm(beta.ash),xlim=c(-4,4),ylim=c(-4,4))
```

## Not run:
# Why is this example included here? -Peter
CIMatrix=ashci(beta.ash,level=0.95)
print(CIMatrix)

## End(Not run)

# Illustrating the non-zero mode feature.
betahat=betahat+5
beta.ash = ash(betahat, sebetahat)
graphics::plot(betahat,get_pm(beta.ash))
betan.ash=ash(betahat, sebetahat,mode=5)
graphics::plot(betahat,get_pm(betan.ash))
summary(betan.ash)

# Running ash with different error models
beta.ash1 = ash(betahat, sebetahat, lik = lik_normal())
beta.ash2 = ash(betahat, sebetahat, lik = lik_t(df=4))
e = rnorm(100)+log(rf(100,df1=10,df2=10)) # simulated data with log(F) error
e.ash = ash(e,1,lik=lik_logF(df1=10,df2=10))

# Specifying the output
beta.ash = ash(betahat, sebetahat, output = c("fitted_g","logLR","lfsr"))

#Running ash with a pre-specified g, rather than estimating it
beta = c(rep(0,100),rnorm(100))
sebeta = rnorm(200,0,1)
betahat = rnorm(200,beta,sebetahat)
true_g = normalmix(c(0.5,0.5),c(0,0),c(0,1)) # define true g
## Passing this g into ash causes it to i) take the sd and the means
## for each component from this g, and ii) initialize pi to the value
## from this g.
beta.ash = ash(betahat, sebetahat,g=true_g,fixg=TRUE)

# running with weights
beta.ash = ash(betahat, sebetahat, optmethod="w_mixEM",
weights = c(rep(0.5,100),rep(1,100)))

# Different algorithms can be used to compute maximum-likelihood
# estimates of the mixture weights. Here, we illustrate use of the
# EM algorithm and the (default) SQP algorithm.
set.seed(1)
betahat <- c(8.115,9.027,9.289,10.097,9.463)
sebeta <- c(0.6157,0.4129,0.3197,0.3920,0.5496)
fit.em <- ash(betahat,sebeta,mixcompdist = "normal",optmethod = "mixEM")
fit.sqp <- ash(betahat,sebeta,mixcompdist = "normal",optmethod = "mixSQP")
range(fit.em$fitted$pi - fit.sqp$fitted$pi)
**Description**

Given the ash object returned by the main function ash, this function computes a posterior credible interval (CI) for each observation. The ash object must include a data component to use this function (which it does by default).

**Usage**

```r
ashci(a, level = 0.95, betaindex, lfsr_threshold = 1, tol = 0.001, trace = FALSE)
```

**Arguments**

- `a` the fitted ash object
- `level` the level for the credible interval, (default=0.95)
- `betaindex` a vector consisting of locations of betahat where you would like to compute the credible interval
- `lfsr_threshold` a scalar, if specified then computes CIs only for observations more significant than that threshold.
- `tol` passed to uniroot; indicates desired accuracy.
- `trace` a logical variable denoting whether some of the intermediate results of iterations should be displayed to the user. Default is FALSE.

**Details**

Uses uniroot to find credible interval, one at a time for each observation. The computation cost is linear in number of observations.

**Value**

A matrix, with 2 columns, ith row giving CI for ith observation

**Examples**

```r
beta = c(rep(0,20),rnorm(20))
sebetahat = abs(rnorm(40,0,1))
betahat = rnorm(40,beta,sebetahat)
beta.ash = ash(betahat, sebetahat)
CImatrix=ashci(beta.ash,level=0.95)
CImatrix1=ashci(beta.ash,level=0.95,betaindex=c(1,2,5))
CImatrix2=ashci(beta.ash,level=0.95,lfsr_threshold=0.1)
```
Description

The main function in the ashr package is `ash`, which should be examined for more details. For simplicity only the most commonly-used options are documented under `ash`. For expert or interested users the documentation for function `ash.workhorse` provides documentation on all implemented options.

ash_pois

Performs adaptive shrinkage on Poisson data

Description

Uses Empirical Bayes to fit the model

\[ y_j | \lambda_j \sim \text{Pois}(c_j \lambda_j) \]

with

\[ h(\lambda_j) g() \]

where \( h \) is a specified link function (either "identity" or "log" are permitted).

Usage

`ash_pois(y, scale = 1, link = c("identity", "log"), ...)`

Arguments

- `y`: vector of Poisson observations.
- `scale`: vector of scale factors for Poisson observations: the model is \( y[j] \sim \text{Pois}(\text{scale}[j] \times \lambda[j]) \).
- `link`: string, either "identity" or "log", indicating the link function.
- `...`: other parameters to be passed to `ash`

Details

The model is fit in two stages: i) estimate \( g \) by maximum likelihood (over the set of symmetric unimodal distributions) to give estimate \( \hat{g} \); ii) Compute posterior distributions for \( \lambda_j \) given \( y_j, \hat{g} \). Note that the link function \( h \) affects the prior assumptions (because, e.g., assuming a unimodal prior on \( \lambda \) is different from assuming unimodal on \( \log \lambda \)), but posterior quantities are always computed for the for \( \lambda \) and *not* \( h(\lambda) \).

Examples

```r
beta = c(rep(0,50),rexp(50))
y = rpois(100,beta) # simulate Poisson observations
y.ash = ash_pois(y,scale=1)
```
calc_loglik

Compute loglikelihood for data from ash fit

Description

Return the log-likelihood of the data for a given g() prior

Usage

calc_loglik(g, data)

Arguments

g # the fitted g, or an ash object containing g
data # a data object, see set_data


calc_logLR

Compute loglikelihood ratio for data from ash fit

Description

Return the log-likelihood ratio of the data for a given g() prior

Usage

calc_logLR(g, data)

Arguments

g # the fitted g, or an ash object containing g
data # a data object, see set_data
calc_mixmean

Generic function of calculating the overall mean of the mixture

Description

Generic function of calculating the overall mean of the mixture

Usage

calc_mixmean(m)

Arguments

m a mixture of k components generated by normalmix() or unimix() or igmix()

Value

it returns scalar, the mean of the mixture distribution.

calc_mixsd

Generic function of calculating the overall standard deviation of the mixture

Description

Generic function of calculating the overall standard deviation of the mixture

Usage

calc_mixsd(m)

Arguments

m a mixture of k components generated by normalmix() or unimix() or igmix()

Value

it returns scalar
**calc_null_loglik**

*Compute loglikelihood for data under null that all beta are 0*

**Description**

Return the log-likelihood of the data betahat, with standard errors betahatsd, under the null that beta==0

**Usage**

```r
calc_null_loglik(data)
```

**Arguments**

- **data**
  - a data object; see set_data

**calc_null_vloglik**

*Compute vector of loglikelihood for data under null that all beta are 0*

**Description**

Return the vector of log-likelihoods of the data points under the null

**Usage**

```r
calc_null_vloglik(data)
```

**Arguments**

- **data**
  - a data object; see set_data

**calc_vloglik**

*Compute vector of loglikelihood for data from ash fit*

**Description**

Return the vector of log-likelihoods of the data betahat, with standard errors betahatsd, for a given g() prior on beta, or an ash object containing that

**Usage**

```r
calc_vloglik(g, data)
```

**Arguments**

- **g**
  - the fitted g, or an ash object containing g
- **data**
  - a data object, see set_data
calc_vlogLR  \hspace{1cm} \textit{Compute vector of loglikelihood ratio for data from ash fit}

\textbf{Description}

Return the vector of log-likelihood ratios of the data betahat, with standard errors betahatsd, for a given $g()$ prior on beta, or an ash object containing that, vs the null that $g()$ is point mass on 0.

\textbf{Usage}

calc_vlogLR(g, data)

\textbf{Arguments}

- \texttt{g} \hspace{1cm} the fitted $g$, or an ash object containing $g$
- \texttt{data} \hspace{1cm} a data object, see \texttt{set_data}

\textbf{cdf.ash} \hspace{1cm} \textit{cdf method for ash object}

\textbf{Description}

Computed the cdf of the underlying fitted distribution

\textbf{Usage}

cdf.ash(a, x, lower.tail = TRUE)

\textbf{Arguments}

- \texttt{a} \hspace{1cm} the fitted ash object
- \texttt{x} \hspace{1cm} the vector of locations at which cdf is to be computed
- \texttt{lower.tail} \hspace{1cm} (default=TRUE) whether to compute the lower or upper tail

\textbf{Details}

None
**cdf_conv**

**Description**
compute cdf of mixture m convoluted with error distribution either normal of sd (s) or student t with df v at locations x

**Usage**
cdf_conv(m, data)

**Arguments**
- **m**: mixture distribution with k components
- **data**: details depend on the model

---

**cdf_post**

**Description**
evaluate cdf of posterior distribution of beta at c. m is the prior on beta, a mixture; c is location of evaluation assumption is betahat | beta ~ t_v(beta,sebetahat)

**Usage**
cdf_post(m, c, data)

**Arguments**
- **m**: mixture distribution with k components
- **c**: a scalar
- **data**: details depend on model

**Value**
an n vector containing the cdf for beta_i at c

**Examples**
```r
beta = rnorm(100,0,1)
betahat = beta+rnorm(100,0,1)
sebetahat = rep(1,100)
ash.beta = ash(betahat,1,mixcompdist="normal")
cdf0 = cdf_post(ash.beta$fitted_g,0,set_data(betahat,sebetahat))
graphics::plot(cdf0,1-get_pp(ash.beta))
```
**compute_lfsr**  
*Function to compute the local false sign rate*

**Description**
Function to compute the local false sign rate

**Usage**
```
compute_lfsr(NegativeProb, ZeroProb)
```

**Arguments**
- **NegativeProb**  
  A vector of posterior probability that beta is negative.
- **ZeroProb**  
  A vector of posterior probability that beta is zero.

**Value**
The local false sign rate.

---

**comp_cdf**  
*Generic function of computing the cdf for each component*

**Description**
Generic function of computing the cdf for each component

**Usage**
```
comp_cdf(m, y, lower.tail = TRUE)
```

**Arguments**
- **m**  
  a mixture (eg of type normalmix or unimix)
- **y**  
  locations at which cdf to be computed
- **lower.tail**  
  boolean indicating whether to report lower tail

**Value**
it returns a vector of probabilities, with length equals to number of components in m
Description
compute the cdf of data for each component of mixture when convolved with error distribution

Usage
```
comp_cdf_conv(m, data)
```

Arguments

- `m` mixture distribution with k components
- `data` details depend on the model

Value

- a k by n matrix of cdfs

Description
returns cdf of convolution of each component of a normal mixture with N(0,s^2) at x. Note that convolution of two normals is normal, so it works that way

Usage
```
## S3 method for class 'normalmix'
comp_cdf_conv(m, data)
```

Arguments

- `m` mixture distribution with k components
- `data` a list with components x and s to be interpreted as a normally-distributed observation and its standard error

Value

- a k by n matrix
comp_cdf_conv

**Description**

cdf of convolution of each component of a unif mixture

**Usage**

```r
## S3 method for class 'unimix'
comp_cdf_conv(m, data)
```

**Arguments**

- `m`: a mixture of class unimix
- `data`: see set_data()

**Value**

a k by n matrix

---

comp_cdf_post

**Description**

evaluate cdf of posterior distribution of beta at c. m is the prior on beta, a mixture; c is location of evaluation assumption is betahat | beta ~ t_v(beta,sebetahat)

**Usage**

```r
comp_cdf_post(m, c, data)
```

**Arguments**

- `m`: mixture distribution with k components
- `c`: a scalar
- `data`: details depend on model

**Value**

a k by n matrix
Examples

```
  beta = rnorm(100,0,1)
  betahat= beta+rnorm(100,0,1)
  sebetahat=rep(1,100)
  ash.beta = ash(betahat,1,mixcompdist="normal")
  comp_cdf_post(get_fitted_g(ash.beta),0,data=set_data(beta,sebetahat))
```

---

**comp_dens**

*Generic function of calculating the component densities of the mixture*

**Description**

Generic function of calculating the component densities of the mixture

**Usage**

```
  comp_dens(m, y, log = FALSE)
```

**Arguments**

- **m**
  - mixture of k components generated by normalmix() or unimix() or igmix()
- **y**
  - is an n-vector of location
- **log**
  - whether to use log-scale on densities

**Value**

A k by n matrix of densities

---

**comp_dens_conv**

*comp_dens_conv*

**Description**

compute the density of data for each component of mixture when convolved with error distribution

**Usage**

```
  comp_dens_conv(m, data)
```

**Arguments**

- **m**
  - mixture distribution with k components
- **data**
  - details depend on the model

**Value**

a k by n matrix of densities
**comp_dens_conv.normalmix**

**Description**

returns density of convolution of each component of a normal mixture with $N(0,s^2)$ at x. Note that convolution of two normals is normal, so it works that way.

**Usage**

```r
## S3 method for class 'normalmix'
comp_dens_conv(m, data)
```

**Arguments**

- `m`: mixture distribution with k components
- `data`: a list with components x and s to be interpreted as a normally-distributed observation and its standard error

**Value**

a k by n matrix

**comp_dens_conv.unimix**

**Description**

density of convolution of each component of a unif mixture

**Usage**

```r
## S3 method for class 'unimix'
comp_dens_conv(m, data)
```

**Arguments**

- `m`: a mixture of class unimix
- `data`: see set_data()

**Value**

a k by n matrix
**comp_mean**

Generic function of calculating the first moment of components of the mixture

**Description**

Generic function of calculating the first moment of components of the mixture

**Usage**

```r
comp_mean(m)
```

**Arguments**

- `m` a mixture of k components generated by normalmix() or unimix() or igmix()

**Value**

it returns a vector of means.

---

**comp_mean.normalmix**

returns mean of the normal mixture

**Usage**

```r
## S3 method for class 'normalmix'
comp_mean(m)
```

**Arguments**

- `m` a normal mixture distribution with k components

**Value**

a vector of length k
comp_mean2

Generic function of calculating the second moment of components of the mixture

Description

Generic function of calculating the second moment of components of the mixture

Usage

comp_mean2(m)

Arguments

m a mixture of k components generated by normalmix() or unimix() or igmix()

Value

it returns a vector of second moments.

comp_postmean

comp_postmean

Description

output posterior mean for beta for each component of prior mixture m, given data

Usage

comp_postmean(m, data)

Arguments

m mixture distribution with k components

data details depend on the model
comp_postmean2

Description
output posterior mean-squared value given prior mixture m and data

Usage
comp_postmean2(m, data)

Arguments
m mixture distribution with k components
data details depend on the model

comp_postprob

Description
compute the posterior prob that each observation came from each component of the mixture m, output a k by n vector of probabilities computed by weighting the component densities by pi and then normalizing

Usage
comp_postprob(m, data)

Arguments
m mixture distribution with k components
data details depend on the model
comp_postsd

Description
output posterior sd for beta for each component of prior mixture m, given data

Usage
comp_postsd(m, data)

Arguments
m mixture distribution with k components
data details depend on the model

Examples
beta = rnorm(100,0,1)
betahat= beta+rnorm(100,0,1)
ash.beta = ash(betahat,1,mixcompdist="normal")
data= set_data(betahat,rep(1,100))
comp_postmean(get_fitted_g(ash.beta),data)
comp_postsd(get_fitted_g(ash.beta),data)
comp_postprob(get_fitted_g(ash.beta),data)

comp_sd

Generic function to extract the standard deviations of components of the mixture

Description
Generic function to extract the standard deviations of components of the mixture

Usage
comp_sd(m)

Arguments
m a mixture of k components generated by normalmix() or unimix() or igmix()

Value
it returns a vector of standard deviations
comp_sd.normalmix

Description
returns sds of the normal mixture

Usage
### S3 method for class 'normalmix'
comp_sd(m)

Arguments
m a normal mixture distribution with k components

Value
a vector of length k

cxxMixSquarem

Description
Brief description of function.

Usage
cxxMixSquarem(matrix_lik, prior, pi_init, control)

Arguments
matrix_lik Description of argument goes here.
prior Description of argument goes here.
pi_init Description of argument goes shere.
control Description of argument goes here.
dens

Find density at y, a generic function

Usage

dens(x, y)

Arguments

x  
A mixture of k components generated by normalmix or unimix.

y  
An n-vector of the location.

dens_conv

dens_conv

dens_conv

Description

compute density of mixture m convoluted with normal of sd (s) or student t with df v at locations x

Usage

dens_conv(m, data)

Arguments

m  
mixture distribution with k components

data  
details depend on the model
The log-F distribution

Description

Density function for the log-F distribution with \( df_1 \) and \( df_2 \) degrees of freedom (and optional non-centrality parameter \( ncp \)).

Usage

\[ dlogf(x, df_1, df_2, ncp, log = \text{FALSE}) \]

Arguments

- \( x \): vector of quantiles
- \( df_1 \): degrees of freedom
- \( df_2 \): degrees of freedom
- \( ncp \): non-centrality parameter. If omitted the central F is assumed.
- \( log \): logical; if TRUE, probabilities \( p \) are given as \( \log(p) \).

Value

The density function.

Estimate mixture proportions of a mixture \( g \) given noisy (error-prone) data from that mixture.

Description

Estimate mixture proportions of a mixture \( g \) given noisy (error-prone) data from that mixture.

Usage

\[ \text{estimate\_mixprop(data, g, prior, optmethod = c("mixSQP", "mixEM", "mixVBEM", "cxxMixSquarem", "mixIP", "w\_mixEM"), control, weights = NULL)} \]
Arguments

data list to be passed to log_comp_dens_conv; details depend on model

g an object representing a mixture distribution (eg normalmix for mixture of normals; unimix for mixture of uniforms). The component parameters of g (eg the means and variances) specify the components whose mixture proportions are to be estimated. The mixture proportions of g are the parameters to be estimated; the values passed in may be used to initialize the optimization (depending on the optmethod used)

prior numeric vector indicating parameters of "Dirichlet prior" on mixture proportions

optmethod name of function to use to do optimization

control list of control parameters to be passed to optmethod, typically affecting things like convergence tolerance

weights vector of weights (for use with w_mixEM; in beta)

Details

This is used by the ash function. Most users won’t need to call this directly, but is exported for use by some other related packages.

Value

list, including the final loglikelihood, the null loglikelihood, an n by k likelihood matrix with (j,k)th element equal to $f_k(x_j)$, the fit and results of optmethod

description Produce function to compute expectation of truncated error distribution from log cdf and log pdf (using numerical integration)

Usage

gen_etruncFUN(lcdfFUN, lpdfFUN)

Arguments

lcdfFUN the log cdfFUN of the error distribution

lpdfFUN the log pdfFUN of the error distribution
get_density

Density method for ash object

Description
Return the density of the underlying fitted distribution

Usage
get_density(a, x)

Arguments
a  the fitted ash object
x  the vector of locations at which density is to be computed

Details
None

get_lfsr
Return lfsr from an ash object

Description
These functions simply return elements of an ash object, generally without doing any calculations. (So if the value was not computed during the original call to ash, eg because of how outputlevel was set in the call, then NULL will be returned.) Accessing elements in this way rather than directly from the ash object will help ensure compatibility moving forward (e.g. if the internal structure of the ash object changes during software development.)

Usage
get_lfsr(x)
get_lfdr(a)
get_svalue(a)
get_qvalue(a)
get_pm(a)
get_psd(a)
get_lfsr(a)
get_pp(a)
get_np(a)
get_loglik(a)
get_logLR(a)
get_fitted_g(a)
get_pi0(a)

Arguments

x an ash fit (e.g. from running ash)
a an ash fit (e.g. from running ash)

Value

a vector (ash) of local false sign rates

Functions

- get_lfsr: local false sign rate
- get_lfdr: local false discovery rate
- get_svalue: svalue
- get_qvalue: qvalue
- get_pm: posterior mean
- get_psd: posterior standard deviation
- get_pp: positive probability
- get_np: negative probability
- get_loglik: log-likelihood
- get_logLR: log-likelihood ratio
- get_fitted_g: fitted g mixture
- get_pi0: pi0, the proportion of nulls
**get_post_sample**  
Sample from posterior

**Description**

Returns random samples from the posterior distribution for each observation in an ash object. A matrix is returned, with columns corresponding to observations and rows corresponding to samples.

**Usage**

\[
\text{get\_post\_sample}(a, \text{nsamp})
\]

**Arguments**

- \(a\)  
  the fitted ash object

- \(\text{nsamp}\)  
  number of samples to return (for each observation)

**Examples**

```r
beta = rnorm(100,0,1)  
betahat = beta+rnorm(100,0,1)  
ash.beta = ash(betahat,1,mixcompdist="normal")  
post.beta = get_post_sample(ash.beta,1000)
```

---

**igmix**  
Constructor for igmix class

**Description**

Creates an object of class igmix (finite mixture of univariate inverse-gammas)

**Usage**

\[
\text{igmix}(\pi, \alpha, \beta)
\]

**Arguments**

- \(\pi\)  
  vector of mixture proportions

- \(\alpha\)  
  vector of shape parameters

- \(\beta\)  
  vector of rate parameters

**Details**

None
lik_binom

**Value**

an object of class igmix

**Examples**


gmix(c(0.5,0.5),c(1,1),c(1,2))

---

**lik_binom**  
*Likelihood object for Binomial error distribution*

**Description**

Creates a likelihood object for ash for use with Binomial error distribution

**Usage**

```r
lik_binom(y, n, link = c("identity", "logit"))
```

**Arguments**

- `y`  
  Binomial observations
- `n`  
  Binomial number of trials
- `link`  
  Link function. The "identity" link directly puts unimodal prior on Binomial success probabilities \( p \), and "logit" link puts unimodal prior on \( \logit(p) \).

**Details**

Suppose we have Binomial observations \( y \) where \( y_i \sim Bin(n_i,p_i) \). We either put an unimodal prior \( g \) on the success probabilities \( p_i \sim g \) (by specifying \( link="identity" \)) or on the logit success probabilities \( \logit(p_i) \sim g \) (by specifying \( link="logit" \)). Either way, ASH with this Binomial likelihood function will compute the posterior mean of the success probabilities \( p_i \).

**Examples**

```r
p = rbeta(100,2,2)  # prior mode: 0.5
n = rpois(100,10)
y = rbinom(100,n,p)  # simulate Binomial observations
ash(rep(0,length(y)),1,lik=lik_binom(y,n))
```
lik_logF

Likelihood object for logF error distribution

Description

Creates a likelihood object for ash for use with logF error distribution

Usage

lik_logF(df1, df2)

Arguments

df1 first degree of freedom parameter of F distribution
df2 second degree of freedom parameter of F distribution

Examples

```r
e = rnorm(100) + log(rf(100, df1=10, df2=10)) # simulate some data with log(F) error
ash(e, 1, lik=lik_logF(df1=10, df2=10))
```

lik_normal

Likelihood object for normal error distribution

Description

Creates a likelihood object for ash for use with normal error distribution

Usage

lik_normal()

Examples

```r
z = rnorm(100) + rnorm(100) # simulate some data with normal error
ash(z, 1, lik=lik_normal())
```
lik_normalmix

*Likelihood object for normal mixture error distribution*

**Description**

Creates a likelihood object for ash for use with normal mixture error distribution

**Usage**

```r
lik_normalmix(pilik, sdlik)
```

**Arguments**

- `pilik`: a k vector of mixture proportions (k is the number of mixture components), or an n*k matrix that the j’th row the is mixture proportions for betahat_j
- `sdlik`: a k vector of component-wise standard deviations, or an n*k matrix that the j’th row the is component-wise standard deviations for betahat_j

**Examples**

```r
e = rnorm(100,0,0.8)
e[seq(1,100,by=2)] = rnorm(50,0,1.5)  # generate e~0.5*N(0,0.8^2)+0.5*N(0,1.5^2)
betahat = rnorm(100)+e
ash(betahat, 1, lik=lik_normalmix(c(0.5,0.5),c(0.8,1.5)))
```

lik_poiss

*Likelihood object for Poisson error distribution*

**Description**

Creates a likelihood object for ash for use with Poisson error distribution

**Usage**

```r
lik_poiss(y, scale = 1, link = c("identity", "log"))
```

**Arguments**

- `y`: Poisson observations.
- `scale`: Scale factor for Poisson observations: y~Pois(scale*lambda).
- `link`: Link function. The "identity" link directly puts unimodal prior on Poisson intensities lambda, and "log" link puts unimodal prior on log(lambda).
Details

Suppose we have Poisson observations \( y_i \sim \text{Poisson}(c_i \lambda_i) \). We either put an unimodal prior \( g \) on the (scaled) intensities \( \lambda_i \sim g \) (by specifying \text{link}="identity") or on the log intensities \( \logit(\lambda_i) \sim g \) (by specifying \text{link}="log"). Either way, ASH with this Poisson likelihood function will compute the posterior mean of the intensities \( \lambda_i \).

Examples

```r
beta = c(rnorm(100,50,5)) # prior mode: 50
y = rpois(100,beta) # simulate Poisson observations
ash(rep(0,length(y)),1,lik=lik_pois(y))
```

```r
lik_t
Likelihood object for t error distribution
```

Description

Creates a likelihood object for ash for use with t error distribution

Usage

```r
lik_t(df)
```

Arguments

- \( df \) degree of freedom parameter of t distribution

Examples

```r
z = rnorm(100) + rt(100,df=4) # simulate some data with t error
ash(z,1,lik=lik_t(df=4))
```

```r
loglik_conv
loglik_conv
```

Description

find log likelihood of data using convolution of mixture with error distribution

Usage

```r
loglik_conv(m, data)
```

Arguments

- \( m \) mixture distribution with k components
- \( data \) details depend on the model
**Description**

The default version of loglik_conv.

**Usage**

```r
## Default S3 method:
loglik_conv(m, data)
```

**Arguments**

- `m` mixture distribution with k components
- `data` data whose details depend on model

---

**Description**

compute the log density of the components of the mixture m when convoluted with a normal with standard deviation s or a scaled (se) student.t with df v, the density is evaluated at x

**Usage**

```r
log_comp_dens_conv(m, data)
```

**Arguments**

- `m` mixture distribution with k components
- `data` details depend on the model

**Value**

a k by n matrix of log densities
**Description**

returns log-density of convolution of each component of a normal mixture with N(0,s^2) or s*t(v) at x. Note that convolution of two normals is normal, so it works that way

**Usage**

```r
## S3 method for class 'normalmix'
log_comp_dens_conv(m, data)
```

**Arguments**

- `m` mixture distribution with k components
- `data` a list with components x and s to be interpreted as a normally-distributed observation and its standard error

**Value**

a k by n matrix

---

**Description**

log density of convolution of each component of a unif mixture

**Usage**

```r
## S3 method for class 'unimix'
log_comp_dens_conv(m, data)
```

**Arguments**

- `m` a mixture of class unimix
- `data` see set_data()

**Value**

a k by n matrix of densities
Description

Returns cdf for a mixture (generic function)

Usage

mixcdf(x, y, lower.tail = TRUE)

Arguments

x
a mixture (eg of type normalmix or unimix)
y
locations at which cdf to be computed
lower.tail
boolean indicating whether to report lower tail

Details

None

Value

an object of class normalmix

Examples

mixcdf(normalmix(c(0.5,0.5),c(0,0),c(1,2)),seq(-4,4,length=100))

Description.default

The default version of mixcdf.

Usage

## Default S3 method:
mixcdf(x, y, lower.tail = TRUE)

Arguments

x
a mixture (eg of type normalmix or unimix)
y
locations at which cdf to be computed
lower.tail
boolean indicating whether to report lower tail
mixEM

Estimate mixture proportions of a mixture model by EM algorithm

Description

Given the individual component likelihoods for a mixture model, estimates the mixture proportions by an EM algorithm.

Usage

mixEM(matrix_lik, prior, pi_init = NULL, control = list())

Arguments

matrix_lik, a n by k matrix with (j,k)th element equal to \( f_k(x_j) \).

prior, a k vector of the parameters of the Dirichlet prior on \( \pi \). Recommended to be rep(1,k).

pi_init, the initial value of \( \pi \) to use. If not specified defaults to (1/k,...,1/k).

control A list of control parameters for the SQUAREM algorithm, default value is set to be control.default=list(K = 1, method=3, square=TRUE, step.min0=1, step.max0=1, mstep=4, kr=1, objfn.inc=1, tol=1.e-07, maxiter=5000, trace=FALSE).

Details

Fits a k component mixture model

\[
f(x|\pi) = \sum_k \pi_k f_k(x)
\]

to independent and identically distributed data \( x_1, \ldots, x_n \). Estimates mixture proportions \( \pi \) by maximum likelihood, or by maximum a posteriori (MAP) estimation for a Dirichlet prior on \( \pi \) (if a prior is specified). Uses the SQUAREM package to accelerate convergence of EM. Used by the ash main function; there is no need for a user to call this function separately, but it is exported for convenience.

Value

A list, including the estimates (pihat), the log likelihood for each iteration (B) and a flag to indicate convergence.
mixIP

Estimate mixture proportions of a mixture model by Interior Point method

Description

Given the individual component likelihoods for a mixture model, estimates the mixture proportions.

Usage

mixIP(matrix_lik, prior, pi_init = NULL, control = list(), weights = NULL)

Arguments

- **matrix_lik**, a n by k matrix with (j,k)th element equal to $f_k(x_j)$.
- **prior**, a k vector of the parameters of the Dirichlet prior on $\pi$. Recommended to be `rep(1,k)`.
- **pi_init**, the initial value of $\pi$ to use. If not specified defaults to (1/k,...,1/k).
- **control** A list of control parameters to be passed to REBayes::KWDual
- **weights** weights to be assigned to the observations (an n vector)

Details

Optimizes

$$L(\pi) = \sum_j w_j \log(\sum_k \pi_k f_{jk}) + h(\pi)$$

subject to $\pi_k$ non-negative and $\sum_k \pi_k = 1$. Here

$$h(\pi)$$

is a penalty function $h(\pi) = \sum_k (\text{prior}_k-1) \log \pi_k$. Calls REBayes::KWDual in the REBayes package, which is in turn a wrapper to the mosek convex optimization software. So REBayes must be installed to use this. Used by the ash main function; there is no need for a user to call this function separately, but it is exported for convenience.

Value

A list, including the estimates (pihat), the log likelihood for each iteration (B) and a flag to indicate convergence
mixmean2

Generic function of calculating the overall second moment of the mixture

Description

Generic function of calculating the overall second moment of the mixture

Usage

mixmean2(m)

Arguments

m a mixture of k components generated by normalmix() or unimix() or igmix()

Value

it returns scalar

mixprop

Generic function of extracting the mixture proportions

Description

Generic function of extracting the mixture proportions

Usage

mixprop(m)

Arguments

m a mixture of k components generated by normalmix() or unimix() or igmix()

Value

it returns a vector of component probabilities, summing up to 1.
mixSQP

*Estimate mixture proportions of a mixture model using mix-SQP algorithm.*

**Description**

Estimate mixture proportions of a mixture model using mix-SQP algorithm.

**Usage**

```r
mixSQP(matrix_lik, prior, pi_init = NULL, control = list(), weights = NULL)
```

**Arguments**

- `matrix_lik`: A matrix containing the conditional likelihood values, possibly normalized.
- `prior`: A vector of the parameters of the Dirichlet prior on the mixture weights.
- `pi_init`: The initial estimate of the mixture weights.
- `control`: A list of settings for the mix-SQP optimization algorithm; see `mixsqp` for details.
- `weights`: The weights to be assigned to the observations. Must be a vector of length equal the number of rows of `matrix_lik`. If `weights = NULL`, all observations are assigned the same weight.

**Value**

A list object including the estimates (`pihat`) and a flag (`control`) indicating convergence success or failure.

mixVBEM

*Estimate posterior distribution on mixture proportions of a mixture model by a Variational Bayes EM algorithm*

**Description**

Given the individual component likelihoods for a mixture model, estimates the posterior on the mixture proportions by an VBEM algorithm. Used by the ash main function; there is no need for a user to call this function separately, but it is exported for convenience.

**Usage**

```r
mixVBEM(matrix_lik, prior, pi_init = NULL, control = list())
```
Arguments

- **matrix_lik**: a n by k matrix with (j,k)th element equal to \( f_k(x_j) \).
- **prior**: a k vector of the parameters of the Dirichlet prior on \( \pi \). Recommended to be \text{rep}(1,k)
- **pi_init**: the initial value of the posterior parameters. If not specified defaults to the prior parameters.
- **control**: A list of control parameters for the SQUAREM algorithm, default value is set to be \text{control.default=list(K = 1, method=3, square=TRUE, step.min0=1, step.max0=1, mstep=4, kr=1, objfn.inc=1, tol=1.e-07, maxiter=5000, trace=FALSE)}.

Details

Fits a k component mixture model

\[
    f(x|\pi) = \sum_k \pi_k f_k(x)
\]

to independent and identically distributed data \( x_1, \ldots, x_n \). Estimates posterior on mixture proportions \( \pi \) by Variational Bayes, with a Dirichlet prior on \( \pi \). Algorithm adapted from Bishop (2009), Pattern Recognition and Machine Learning, Chapter 10.

Value

A list, whose components include point estimates (pihat), the parameters of the fitted posterior on \( \pi \) (pipost), the bound on the log likelihood for each iteration (B) and a flag to indicate convergence (converged).

---

**my_e2truncbeta**  
*second moment of truncated Beta distribution*

Description

Compute second moment of the truncated Beta.

Usage

```r
my_e2truncbeta(a, b, alpha, beta)
```

Arguments

- **a**: left limit of distribution
- **b**: right limit of distribution
- **alpha, beta**: shape parameters of Beta distribution
my_e2truncgamma  second moment of truncated gamma distribution

Description

Compute second moment of the truncated gamma.

Usage

my_e2truncgamma(a, b, shape, rate)

Arguments

a  left limit of distribution
b  right limit of distribution
shape  shape of gamma distribution
rate  rate of gamma distribution

my_e2truncnorm  Expected Squared Value of Truncated Normal

Description

Computes the expected squared values of truncated normal distributions with parameters a, b, mean, and sd. Arguments can be scalars, vectors, or matrices. Arguments of shorter length will be recycled according to the usual recycling rules, but a and b must have the same length. Missing values are accepted for all arguments.

Usage

my_e2truncnorm(a, b, mean = 0, sd = 1)

Arguments

a  The lower limit for the support of the truncated normal. Can be -Inf.
b  The upper limit for the support. Can be Inf. a and b must have the same length, and each element of a should be less than or equal to the corresponding element of b.
mean  The mean of the untruncated normal.
sd  The standard deviation of the untruncated normal. Standard deviations of zero are interpreted as numerically (rather than exactly) zero, so that the square of the untruncated mean is returned if it lies within [a, b] and the square of the nearer of a and b is returned otherwise.
**Value**

The expected squared values of truncated normal distributions with parameters \( a, b, \text{mean}, \text{sd} \). If any of the arguments is a matrix, then a matrix will be returned.

**See Also**

`my_etruncnorm, my_vtruncnorm`

---

**Description**

Compute second moment of the truncated \( t \). Uses results from O'Hagan, Biometrika, 1973

**Usage**

```
my_e2trunct(a, b, df)
```

**Arguments**

- **a**: left limit of distribution
- **b**: right limit of distribution
- **df**: degree of freedom of error distribution

---

**Description**

Compute mean of the truncated Beta.

**Usage**

```
my_etruncbeta(a, b, alpha, beta)
```

**Arguments**

- **a**: left limit of distribution
- **b**: right limit of distribution
- **alpha, beta**: shape parameters of Beta distribution
Description
Compute mean of the truncated gamma.

Usage
my_etruncgamma(a, b, shape, rate)

Arguments
a        left limit of distribution
b        right limit of distribution
shape    shape of gamma distribution
rate     rate of gamma distribution

Description
Compute expectation of truncated log-F distribution.

Usage
my_etrunclogf(a, b, df1, df2)

Arguments
a        Left limit of distribution.
b        Right limit of distribution.
df1, df2  degrees of freedom
my_etruncnorm

Description
Computes the means of truncated normal distributions with parameters \(a\), \(b\), \(\text{mean}\), and \(\text{sd}\). Arguments can be scalars, vectors, or matrices. Arguments of shorter length will be recycled according to the usual recycling rules, but \(a\) and \(b\) must have the same length. Missing values are accepted for all arguments.

Usage
my_etruncnorm(a, b, mean = 0, sd = 1)

Arguments
- \(a\): The lower limit for the support of the truncated normal. Can be \(-\infty\).
- \(b\): The upper limit for the support. Can be \(\infty\). \(a\) and \(b\) must have the same length, and each element of \(a\) should be less than or equal to the corresponding element of \(b\).
- \(\text{mean}\): The mean of the untruncated normal.
- \(\text{sd}\): The standard deviation of the untruncated normal. Standard deviations of zero are interpreted as numerically (rather than exactly) zero, so that the untruncated mean is returned if it lies within \([a, b]\) and the nearer of \(a\) and \(b\) is returned otherwise.

Value
The expected values of truncated normal distributions with parameters \(a\), \(b\), \(\text{mean}\), and \(\text{sd}\). If any of the arguments is a matrix, then a matrix will be returned.

See Also
my_e2truncnorm, my_vtruncnorm

my_etrunct

Description
Compute second moment of the truncated t. Uses results from O’Hagan, Biometrika, 1973

Usage
my_etrunct(a, b, df)
Arguments

- **a**: left limit of distribution
- **b**: right limit of distribution
- **df**: degree of freedom of error distribution

### my_vtruncnorm

Variance of Truncated Normal

**Description**

Computes the variance of truncated normal distributions with parameters \(a, b, \text{mean}, \text{sd}\). Arguments can be scalars, vectors, or matrices. Arguments of shorter length will be recycled according to the usual recycling rules, but \(a\) and \(b\) must have the same length. Missing values are accepted for all arguments.

**Usage**

```r
my_vtruncnorm(a, b, mean = 0, sd = 1)
```

**Arguments**

- **a**: The lower limit for the support of the truncated normal. Can be \(-\infty\).
- **b**: The upper limit for the support. Can be \(\infty\). \(a\) and \(b\) must have the same length, and each element of \(a\) should be less than or equal to the corresponding element of \(b\).
- **mean**: The mean of the untruncated normal.
- **sd**: The standard deviation of the untruncated normal.

**Value**

The variance of truncated normal distributions with parameters \(a, b, \text{mean}, \text{sd}\). If any of the arguments is a matrix, then a matrix will be returned.

**See Also**

- `my_etruncnorm`
- `my_e2truncnorm`
**ncomp**

---

<table>
<thead>
<tr>
<th>ncomp</th>
<th>ncomp</th>
</tr>
</thead>
</table>

**Description**

ncomp

**Usage**

ncomp(m)

**Arguments**

m a mixture of k components generated by normalmix() or unimix() or igmix()

---

<table>
<thead>
<tr>
<th>ncomp.default</th>
<th>ncomp.default</th>
</tr>
</thead>
</table>

**Description**

The default version of ncomp.

**Usage**

## Default S3 method:

ncomp(m)

**Arguments**

m a mixture of k components generated by normalmix() or unimix() or igmix()

---

<table>
<thead>
<tr>
<th>normalmix</th>
<th>Constructor for normalmix class</th>
</tr>
</thead>
</table>

**Description**

Creates an object of class normalmix (finite mixture of univariate normals)

**Usage**

normalmix(pi, mean, sd)
Arguments

- \(\pi\): vector of mixture proportions
- mean: vector of means
- sd: vector of standard deviations

Details

None

Value

an object of class normalmix

Examples

normalmix(c(0.5, 0.5), c(0, 0), c(1, 2))

Description

"parallel" vector version of \texttt{cdf_post} where \(c\) is a vector, of same length as \(\text{betahat}\) and \(\text{sebetahat}\)

Usage

\texttt{pcdf_post}(m, c, \text{data})

Arguments

- \(m\): mixture distribution with \(k\) components
- \(c\): a numeric vector with \(n\) elements
- data: depends on context

Value

an \(n\) vector, whose \(\text{ith element}\) is the \(\text{cdf}\) for \(\text{beta\_i}\) at \(c\_i\)

Examples

\texttt{beta = rnorm(100, 0, 1)}
\texttt{betahat = beta + rnorm(100, 0, 1)}
\texttt{sebetahat = rep(1, 100)}
\texttt{ash.beta = ash(betahat, 1, mixcompdist="normal")}
\texttt{c = pcdf_post(get_fitted_g(ash.beta), beta, set_data(betahat, sebetahat))}
plogf

The log-F distribution

Description

Distribution function for the log-F distribution with df1 and df2 degrees of freedom (and optional non-centrality parameter ncp).

Usage

plogf(q, df1, df2, ncp, lower.tail = TRUE, log.p = FALSE)

Arguments

q vector of quantiles
df1, df2 degrees of freedom
ncp non-centrality parameter. If omitted the central F is assumed.
lower.tail logical; if TRUE (default), probabilities are P[X <= x], otherwise, P[X > x].
log.p logical; if TRUE, probabilities p are given as log(p).

Value

The distribution function.

plot.ash

Plot method for ash object

Description

Plot the cdf of the underlying fitted distribution

Usage

## S3 method for class 'ash'
plot(x, ..., xmin, xmax)

Arguments

x the fitted ash object
... Arguments to be passed to methods, such as graphical parameters (see plot)
xmin xlim lower range, default is the lowest value of betahat
xmax xlim upper range, default is the highest value of betahat

Details

None
plot_diagnostic  Diagnostic plots for ash object

Description
Generate several plots to diagnose the fitness of ASH on the data

Usage
plot_diagnostic(x, plot.it = TRUE, sebetahat.tol = 0.001, plot.hist, 
    xmin, xmax, breaks = "Sturges", alpha = 0.01, pch = 19, 
    cex = 0.25)

Arguments
- x: the fitted ash object
- plot.it: logical. whether to plot the diagnostic result
- sebetahat.tol: tolerance to test the equality of betahat
- plot.hist: logical. whether to plot the histogram of betahat when sebetahat is not constant
- xmin, xmax: range of the histogram of betahat to be plotted
- breaks: histograms parameter (see hist)
- alpha: error level for the de-trended diagnostic plot
- pch, cex: plot parameters for dots

Details
None.

pm_on_zero  Generic function to extract which components of mixture are point mass on 0

Description
Generic function to extract which components of mixture are point mass on 0

Usage
pm_on_zero(m)

Arguments
- m: a mixture of k components generated by normalmix() or unimix() or igmix()

Value
a boolean vector indicating which components are point mass on 0
posterior_dist

Description
Return the posterior on beta given a prior (g) that is a mixture of normals (class normalmix) and observation \( \text{betahat} \sim N(\text{beta}, \text{sebetahat}) \)

Usage
posterior_dist(g, betahat, sebetahat)

Arguments
- \( g \): a normalmix with components indicating the prior; works only if \( g \) has means 0
- \( \text{betahat} \): (n vector of observations)
- \( \text{sebetahat} \): (n vector of standard errors/deviations of observations)

Details
This can be used to obt

Value
A list, \((\pi_1, \mu_1, \sigma_1)\) whose components are each \( k \) by \( n \) matrices where \( k \) is number of mixture components in \( g \), \( n \) is number of observations in \( \text{betahat} \)

postmean

Description
postmean

Usage
postmean(m, data)

Arguments
- \( m \): mixture distribution with \( k \) components
- \( \text{data} \): details depend on the model
**Description**

output posterior mean-squared value given prior mixture `m` and data

**Usage**

```r
postmean2(m, data)
```

**Arguments**

- `m`  
  mixture distribution with k components
- `data`  
  details depend on the model

---

**Description**

output posterior sd given prior mixture `m` and data

**Usage**

```r
postsd(m, data)
```

**Arguments**

- `m`  
  mixture distribution with k components
- `data`  
  details depend on the model
Description

returns random samples from the posterior, given a prior distribution m and n observed datapoints.

Usage

post_sample(m, data, nsamp)

Arguments

m prior distribution (eg of type normalmix)
data a list with components x and s, each vectors of length n, to be interpreted as a normally-distributed observations and corresponding standard errors
nsamp number of random samples to return for each observation

Details

exported, but mostly users will want to use ‘get_post_sample’

Value

an nsamp by n matrix

Description

returns random samples from the posterior, given a prior distribution m and n observed datapoints.

Usage

## S3 method for class 'normalmix'
post_sample(m, data, nsamp)

Arguments

m mixture distribution with k components
data a list with components x and s to be interpreted as a normally-distributed observation and its standard error
nsamp number of samples to return for each observation
### post_sample.unimix

**Description**
returns random samples from the posterior, given a prior distribution m and n observed datapoints.

**Usage**
```r
## S3 method for class 'unimix'
post_sample(m, data, nsamp)
```

**Arguments**
- `m`: mixture distribution with k components
- `data`: a list with components x and s to be interpreted as a normally-distributed observation and its standard error
- `nsamp`: number of samples to return for each observation

**Value**
a nsamp by n matrix

---

### print.ash

**Description**
Print the fitted distribution of beta values in the EB hierarchical model

**Usage**
```r
## S3 method for class 'ash'
print(x, ...)
```

**Arguments**
- `x`: the fitted ash object
- `...`: not used, included for consistency as an S3 generic/method.

**Details**
None
prune

Description
prunes out mixture components with low weight

Usage
prune(m, thresh = 1e-10)

Arguments
m
What is this argument?
thresh
the threshold below which components are removed

qval.from.lfdr
Function to compute q values from local false discovery rates

Description
Computes q values from a vector of local fdr estimates

Usage
qval.from.lfdr(lfdr)

Arguments
lfdr,
a vector of local fdr estimates

Details
The q value for a given lfdr is an estimate of the (tail) False Discovery Rate for all findings with a smaller lfdr, and is found by the average of the lfdr for all more significant findings. See Storey (2003), Annals of Statistics, for definition of q value.

Value
vector of q values
set_data  
*Takes raw data and sets up data object for use by ash*

**Description**
Takes raw data and sets up data object for use by ash

**Usage**
```r
define set_data(betahat, sebetahat, lik = NULL, alpha = 0)
```

**Arguments**
- `betahat`: vector of betahats
- `sebetahat`: vector of standard errors
- `lik`: a likelihood (see e.g., `lik_normal()`)  
- `alpha`: specifies value of alpha to use (model is for betahat/sebetahat^alpha | sebetahat)

**Details**
The data object stores both the data, and details of the model to be used for the data. For example, in the generalized version of ash the cdf and pdf of the likelihood are stored here.

**Value**
data object (list)

---

summary.ash  
*Summary method for ash object*

**Description**
Print summary of fitted ash object

**Usage**
```r
## S3 method for class `ash`
summary(object, ...)
```

**Arguments**
- `object`: the fitted ash object
- `...`: not used, included for consistency as an S3 generic/method.
unimix

Details
summary prints the fitted mixture, the fitted log likelihood with 10 digits and a flag to indicate convergence

unimix Constructor for unimix class

Description
Creates an object of class unimix (finite mixture of univariate uniforms)

Usage
unimix(pi, a, b)

Arguments
pi vector of mixture proportions
a vector of left hand ends of uniforms
b vector of right hand ends of uniforms

Details
None

Value
an object of class unimix

Examples
unimix(c(0.5,0.5),c(0,0),c(1,2))

vcdf_post vcdf_post

Description
vectorized version of cdf_post

Usage
vcdf_post(m, c, data)
w_mixEM

Estimate mixture proportions of a mixture model by EM algorithm (weighted version)

Arguments

- **m**: mixture distribution with k components
- **c**: a numeric vector
- **data**: depends on context

Value

- an n vector containing the cdf for beta_i at c

Examples

```r
beta = rnorm(100,0,1)
betahat = beta+rnorm(100,0,1)
sebetahat = rep(1,100)
ash.beta = ash(betahat,1,mixcompdist="normal")
c = vcdf_post(get_fitted_g(ash.beta),seq(-5,5,length=1000),data = set_data(betahat,sebetahat))
```

Description

Given the individual component likelihoods for a mixture model, and a set of weights, estimates the mixture proportions by an EM algorithm.

Usage

```r
w_mixEM(matrix_lik, prior, pi_init = NULL, weights = NULL, control = list())
```

Arguments

- **matrix_lik**, a n by k matrix with (j,k)th element equal to \( f_k(x_j) \).
- **prior**, a k vector of the parameters of the Dirichlet prior on \( \pi \). Recommended to be \( \text{rep}(1,k) \).
- **pi_init**, the initial value of \( \pi \) to use. If not specified defaults to \( (1/k,...,1/k) \).
- **weights**, an n vector of weights
- **control** A list of control parameters for the SQUAREM algorithm, default value is set to be control.default=list(K = 1, method=3, square=TRUE, step.min0=1, step.max0=1, mstep=4, kr=1, objfn.inc=1,tol=1.e-07, maxiter=5000, trace=FALSE).
Details

Fits a k component mixture model

\[ f(x|\pi) = \sum_k \pi_k f_k(x) \]

to independent and identically distributed data \( x_1, \ldots, x_n \) with weights \( w_1, \ldots, w_n \). Estimates mixture proportions \( \pi \) by maximum likelihood, or by maximum a posteriori (MAP) estimation for a Dirichlet prior on \( \pi \) (if a prior is specified). Here the log-likelihood for the weighted data is defined as \( l(\pi) = \sum_j w_j \log f(x_j|\pi) \). Uses the SQUAREM package to accelerate convergence of EM. Used by the ash main function; there is no need for a user to call this function separately, but it is exported for convenience.

Value

A list, including the estimates (pihat), the log likelihood for each iteration (B) and a flag to indicate convergence.
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