

# Package ‘mederrRank’

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**Title** Bayesian Methods for Identifying the Most Harmful Medication Errors

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**Description** Two distinct but related statistical approaches to the problem of identifying the combinations of medication error characteristics that are more likely to result in harm are implemented in this package: 1) a Bayesian hierarchical model with optimal Bayesian ranking on the log odds of harm, and 2) an empirical Bayes model that estimates the ratio of the observed count of harm to the count that would be expected if error characteristics and harm were independent. In addition, for the Bayesian hierarchical model, the package provides functions to assess the sensitivity of results to different specifications of the random effects distributions.

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**mederrRank-package**      *Bayesian Methods for Identifying the Most Harmful Medication Errors*

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**Description**

Two distinct but related statistical approaches to the problem of identifying the combinations of medication error characteristics that are more likely to result in harm are implemented in this package: 1) a Bayesian hierarchical model with optimal Bayesian ranking on the log odds of harm, and 2) an empirical Bayes model that estimates the ratio of the observed count of harm to the count that would be expected if error characteristics and harm were independent. In addition, for the Bayesian hierarchical model, the package provides functions to assess the sensitivity of results to different specifications of the random effects distributions.

**Details**

The package is loaded with the usual `library(mederrRank)` command. The most important functions are `bhm.mcmc`, `bhm.resample` and `mixnegbinom.em`.

**Author(s)**

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## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[bayes.rank](#), [bhm.mcmc](#), [bhm.resample](#), [mixnegbinom.em](#).

`bayes.rank`

*Optimal Bayesian Ranking*

## Description

This function estimates the ranks of the log odds of harm of the various medication error profiles as described in Myers et al. (2011).

## Usage

```
bayes.rank(model)
```

## Arguments

`model` a [mederrFit](#) object.

## Details

Using the posterior samples of the  $\theta_i$ , the function estimates the ranks of the log odds of harm of the various error profiles. Optimal Bayesian ranking gives estimates of rank for profile  $i$  as

$$\hat{R}_i = \sum_{k=1}^n \hat{P}(\theta_k \leq \theta_i | \mathbf{y}, \mathbf{N}),$$

where  $\hat{P}(\theta_k \leq \theta_i | \mathbf{y}, \mathbf{N})$  is the posterior probability that  $\theta_k \leq \theta_i$ .

## Value

`bayes.rank` returns the numerical vector of Optimal Bayesian ranks for the chosen [mederrFit](#) model (see the references for the details).

## Author(s)

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## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[bhm.constr.resamp](#), [bhm.mcmc](#).

## Examples

```
## Not run:
data("simdata", package = "mederrRank")
summary(simdata)

fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
ranks <- bayes.rank(fit)

summary(ranks)

## End(Not run)
```

**bhm.constr.resamp**

*Markov Chain Monte Carlo Estimation (Step 2) of the Bayesian Hierarchical Model for Identifying the Most Harmful Medication Errors*

## Description

This function represents the "constructor" function for the resampling procedure used in this package. [bhm.resample](#) calculates the importance ratios, and performs the sampling, and then this function constructs the resampled model based on that information.

## Usage

`bhm.constr.resamp(model, resample, k, eta)`

## Arguments

<code>model</code>	an object of <a href="#">class "mederrFit"</a> .
<code>resample</code>	an object of <a href="#">class "mederrResample"</a> .
<code>k</code>	$k$ (number of degrees of freedom) value to use in the resampling procedure.
<code>eta</code>	$\eta$ (skewing parameter) value to use in the resampling procedure.

## Details

Deviations from the normal, i.e. ( $k = \infty, \eta = 1$ ), random effects distribution using a different pair of  $k$  and  $\eta$  values are considered. The methodology implemented here is the importance link function resampling approach introduced by MacEachern and Peruggia (2000): based on the ( $k = \infty, \eta = 1$ ) chain, new posterior samples under a new set of ( $k, \eta$ ) values is obtained.

**Value**

`bhm.constr.resamp` returns an object of the class "[mederrFit](#)".

**Author(s)**

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Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

MacEachern, S. and Peruggia, M. (2000), "Importance Link Function Estimation for Markov Chain Monte Carlo Methods", *Journal of Computational and Graphical Statistics*, 9, 99-121.

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bhm.mcmc](#), [bhm.resample](#), [mederrData](#), [mederrFit](#).

**Examples**

```
## Not run:
data("simdata", package = "mederrRank")
summary(simdata)

fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
resamp <- bhm.resample(fit, simdata, p.resample = .1,
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)
plot(fit, fit2, simdata)

theta0 <- c(10, 6, 100, 100, .1)
ans <- mixnegbinom.em(simdata, theta0, 50000, 0.01, se = TRUE,
stratified = TRUE)

summary(fit2, ans, simdata)

## End(Not run)
```

**Description**

This function implements the Markov Chain Monte Carlo estimation methodology for the Bayesian hierarchical model described in Myers et al. (2011).

## Usage

```
bhm.mcmc(dat, nsim = 2000, burnin = 500, scale.factor = 1,
adaptive.int = 100, adaptive.max = 1000, prior = NULL,
init = NULL, tuneD = NULL, tuneT = NULL)
```

## Arguments

<code>dat</code>	an object of <a href="#">class "mederrData"</a> .
<code>nsim</code>	number of iterations.
<code>burnin</code>	number of burn-in iterations.
<code>scale.factor</code>	scale factor of the random effects proposal distribution.
<code>adaptive.int</code>	iteration interval at which the standard error of the random effects proposal distribution is updated.
<code>adaptive.max</code>	last iteration at which the standard error of the random effects proposal distribution is updated.
<code>prior</code>	an optional list of the hyperparameters values; see the Details section below.
<code>init</code>	an optional list of initial values for the model parameters; see the Details section below.
<code>tuneD</code>	an optional vector of the $\delta_j$ proposal distribution variances.
<code>tuneT</code>	an optional vector of the $\theta_i$ proposal distribution variances.

## Details

The Bayesian hierarchical model (with crossed random effects) implemented here for identifying the medication error profiles with the largest log odds of harm is

$$\begin{aligned}
y_{ij} | N_{ij}, p_{ij} &\sim \text{Bin}(N_{ij}, p_{ij}) \\
\text{logit}(p_{ij}) &= \gamma + \theta_i + \delta_j \\
\theta_i | \sigma, \eta, k &\sim \text{St}(0, \sigma, k, \eta), \quad i = 1, \dots, n \\
\delta_j | \tau^2 &\sim N(0, \tau^2), \quad j = 1, \dots, J \\
\gamma &\sim N(g, G) \\
\sigma^2 &\sim \text{IG}(a_1, b_1) \\
\tau^2 &\sim \text{IG}(a_2, b_2) \\
k &\sim \text{Unif}(0, \infty) \\
\eta &\sim \text{Unif}(0, \infty),
\end{aligned}$$

where  $N_{ij}$  denotes the number of times that the error profile  $i$  is cited on a report from hospital  $j$  and  $y_{ij}$  is the corresponding number of times that profile  $i$  in hospital  $j$  was reported with harm. This function implements the first model estimation step in which the values  $k = \infty$  and  $k = 1$ , i.e. a symmetric normal distribution, is forced for the error profiles' random effects. A sample from the joint posterior distribution of all other parameters via Markov Chain Monte Carlo with adaptive Metropolis steps for each set of random effects is obtained. For more details see Myers et al. (2011).

**Value**

`bhm.mcmc` returns an object of the class "[mederrFit](#)".

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**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bhm.resample](#), [mederrData](#), [mederrFit](#).

**Examples**

```
## Not run:  
data("simdata", package = "mederrRank")  
summary(simdata)  
  
fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)  
resamp <- bhm.resample(fit, simdata, p.resample = .1,  
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))  
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)  
plot(fit, fit2, simdata)  
  
theta0 <- c(10, 6, 100, 100, .1)  
ans <- mixnegbinom.em(simdata, theta0, 50000, 0.01, se = TRUE,  
stratified = TRUE)  
  
summary(fit2, ans, simdata)  
## End(Not run)
```

---

**bhm.resample**

*Resampling Transformation for the Markov Chain Monte Carlo Estimation Simulation of the Bayesian Hierarchical Model for Identifying the Most Harmful Medication Errors*

---

**Description**

This function implements the transformation needed to apply the importance link function resampling methodology based on the Markov Chain Monte Carlo simulations obtained with the `bhm.mcmc` command (see the References).

**Usage**

```
bhm.resample(model, dat, p.resample = 0.1, k, eta)
```

**Arguments**

model	an object of <a href="#">class "mederrFit"</a> .
dat	an object of <a href="#">class "mederrData"</a> .
p.resample	proportion of simulations resampled from the <code>model</code> argument.
k	required vector of $k$ values to be used in the resampling process.
eta	required vector of $\eta$ values to be used in the resampling process.

**Value**

`bhm.resample` returns an object of the class "[mederrResample](#)".

**Author(s)**

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**References**

- MacEachern, S. and Peruggia, M. (2000), "Importance Link Function Estimation for Markov Chain Monte Carlo Methods", *Journal of Computational and Graphical Statistics*, 9, 99-121.  
 Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[mederrData](#), [mederrFit](#), [bhm.mcmc](#).

**Examples**

```
## Not run:
data("simdata", package = "mederrRank")
summary(simdata)

fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
resamp <- bhm.resample(fit, simdata, p.resample = .1,
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)
plot(fit, fit2, simdata)

theta0 <- c(10, 6, 100, 100, .1)
ans <- mixnegbinom.em(simdata, theta0, 50000, 0.01, se = TRUE,
stratified = TRUE)

summary(fit2, ans, simdata)
```

---

```
## End(Not run)
```

---

**dmixnegbinom***The Negative Binomial Mixture Distribution*

## Description

Density function for a mixture of two negative binomial distributions.

## Usage

```
dmixnegbinom(x, theta, E, log.p = FALSE)
```

## Arguments

<code>x</code>	vector of (non-negative integer) quantiles.
<code>theta</code>	vector of parameters for the negative binomial distribution mixture.
<code>E</code>	vector of (non-negative integer) expected counts.
<code>log.p</code>	logical; if TRUE, probabilities p are given as log(p).

## Details

The mixture of two negative binomial distributions has density

$$P(N = x) = \text{theta}[5]f(x; \text{theta}[1], \text{theta}[2], E) + (1 - \text{theta}[5])f(x; \text{theta}[3], \text{theta}[4], E),$$

where

$$f(x; \alpha, \beta, E) = \frac{\Gamma(\alpha + x)}{\Gamma(\alpha)x!} \frac{1}{(1 + \beta/E)^x} \frac{1}{(1 + E/\beta)^\alpha}$$

for  $x = 0, 1, \dots, \alpha, \alpha, \beta, E > 0$  and  $0 < \text{theta}[5] \leq 1$ . The mixture of two negative binomial distributions represents the marginal distribution of the counts  $N$  coming from Poisson data with parameter  $\lambda$  and a mixture of two gamma distributions as its prior. For details see the paper by Dumouchel (1999).

## Value

`dmixnegbinom` gives the density corresponding to the `E` and `theta` values provided.

## Author(s)

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 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

- DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.
- Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[dnbinom](#), [rmixnegbinom](#).

## Examples

```
## Not run:
data("simdata", package = "mederrRank")
ni <- simdata@numi
theta0 <- c(10, 6, 100, 100, .1)
ans <- mixnegbinom.em(simdata, theta0, 50000, 0.01,
se = FALSE, stratified = TRUE)
theta <- ans$theta.hat
N.E <- cbind(ans$N[1:ni], ans$E[1:ni])[sort(ans$N[1:ni], index.return = TRUE)$ix, ]
N.ix <- match(unique(N.E[, 1]), N.E[, 1])
N <- N.E[N.ix, 1]
E <- N.E[N.ix, 2]
dens <- dmixnegbinom(N, theta, E)
hist(N.E[, 1], breaks = 40, freq = FALSE)
points(N, dens)

## End(Not run)
```

**dnegbinom**

*The Negative Binomial Distribution*

## Description

Density function for the negative binomial distribution with parameters alpha and prob.

## Usage

`dnegbinom(x, alpha, prob, log.p = FALSE)`

## Arguments

- |                    |   |
|--------------------|---|
| <code>x</code>     | vector of (non-negative integer) quantiles.   |
| <code>alpha</code> | target for number of successful trials. Must be strictly positive, need not be integer. |
| <code>prob</code>  | probability of success in each trial. $0 < \text{prob} \leq 1$ .                        |
| <code>log.p</code> | logical; if TRUE, probabilities p are given as log(p).                                  |

## Details

The negative binomial distribution with parameters `alpha = alpha` and `prob = prob` has density

$$\frac{\Gamma(x + \alpha)}{\Gamma(\alpha)x!} p^\alpha (1 - p)^x$$

for  $x = 0, 1, \dots, \alpha > 0$  and  $0 < p \leq 1$ . This represents the number of failures which occur in a sequence of Bernoulli trials before a target number of successes is reached.

## Value

`dnegbinom` gives the density corresponding to the `alpha` and `prob` values provided.

## Author(s)

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 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[dmixnegbinom](#), [dnbinom](#).

dst

*The Skewed Student t Distribution*

## Description

Density function for the skewed t distribution with `k` degrees of freedom, scale parameter `sigma` and skewness `eta`.

## Usage

`dst(x, sigma, k, eta)`

## Arguments

<code>x</code>	vector of quantiles.
<code>sigma</code>	scale parameter ( $> 0$ ).
<code>k</code>	degrees of freedom ( $> 0$ , maybe non-integer). <code>df = Inf</code> is allowed.
<code>eta</code>	skewness parameter ( $> 0$ ).

## Details

This distribution is based on introducing skewing into the symmetric scaled t distribution, as described in Fernandez and Steel (1998). The parameters characterizing the center (here set at 0) and the spread (*sigma*) refer to the mean and standard deviation of the underlying symmetric distribution. In the skewed t distribution, the centrality parameter defines the mode of the distribution, but it is no longer either the mean or the median. Similarly, in the skewed t distribution, *sigma* still characterizes the spread, but it can no longer be interpreted directly as the standard deviation of the distribution.

## Value

*dst* gives the density corresponding to the *sigma*, *k* and *eta* values provided.

## Author(s)

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Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

Fernandez, C. and Steel, M. (1998), "On Bayesian Modeling of Fat Tails and Skewness". Journal of the American Statistical Association, 93, 359-371.

Lee, K. and Thompson, S. (2008), "Flexible Parametric Models for Random-Effects Distributions". Statistics in Medicine, 27, 418-434.

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[dt](#).

EBGM

*Geometric Mean of the Relative Risk Empirical Bayes Posterior Distribution*

## Description

This function computes the geometric mean of the empirical Bayes posterior distribution for the observed vs. expected count relative risk.

## Usage

`EBGM(eb.result)`

## Arguments

`eb.result` output of the [mixnegbinom.em](#) or [negbinom.em](#) commands.

## Details

For further details see DuMouchel (1999).

## Value

EBGM returns the vector of geometric means.

## Author(s)

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 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[mixnegbinom.em](#), [negbinom.em](#).

## Examples

```
## Not run:
data("simdata", package = "mederrRank")
summary(simdata)

fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
resamp <- bhm.resample(fit, simdata, p.resample = .1,
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)

theta0 <- c(10, 6, 100, 100, .1)
ans <- mixnegbinom.em(simdata, theta0, 50000, 0.01,
se = FALSE, stratified = TRUE)

ni <- simdata@numi
rank(EBGM(ans)[1:ni])
summary(fit2, ans, simdata)
## End(Not run)
```

**llDiffD***Log-Likelihood Difference for the  $\delta_j$  Parameters***Description**

This function computes the log-likelihood difference for the candidate  $\delta_j$  random effects. It is a helper function and not meant to be used on its own.

**Usage**

```
llDiffD(dat, deltaj, cand, thetai, gamma, tau2)
```

**Arguments**

<code>dat</code>	data frame containing the observed sample counts.
<code>deltaj</code>	vector of previous accepted values for the $\delta_j$ random effects.
<code>cand</code>	vector of candidate values for the $\delta_j$ random effects.
<code>thetai</code>	vector of previous accepted values for the $\theta_i$ random effects.
<code>gamma</code>	last sampled value for the $\gamma$ parameter.
<code>tau2</code>	last sampled value for the $\tau^2$ parameter.

**Details**

For further details see Myers et al. (2011).

**Value**

`llDiffD` returns the vector of log-likelihood differences.

**Author(s)**

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 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[`bhm.constr.resamp`](#), [`bhm.mcmc`](#), [`bhm.resample`](#).

---

llDiffT*Log-Likelihood Difference for the  $\theta_i$  Parameters*

---

**Description**

This function computes the log-likelihood difference for the candidate  $\theta_i$  random effects. It is a helper function and not meant to be used on its own.

**Usage**

```
llDiffT(dat, thetai, cand, deltaj, gamma, sigma2)
```

**Arguments**

dat	data frame containing the observed sample counts.
thetai	vector of previous accepted values for the $\theta_i$ random effects.
cand	vector of candidate values for the $\delta_j$ random effects.
deltaj	vector of previous accepted values for the $\delta_j$ random effects.
gamma	last sampled value for the $\gamma$ parameter.
sigma2	last sampled value for the $\sigma^2$ parameter.

**Details**

For further details see Myers et al. (2011).

**Value**

llDiffT returns the vector of log-likelihood differences.

**Author(s)**

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**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bhm.constr.resamp](#), [bhm.mcmc](#), [bhm.resample](#).

---

**logp***Negative Log-Posterior Function of the Bayesian Hierarchical Model  
for Identifying the Most Harmful Medication Errors*

---

## Description

This function computes the negative log-posterior distribution of the Bayesian hierarchical model described in Myers et al (2011). It is a helper function and not meant to be used on its own.

## Usage

```
logp(theta, deltaj, sigma2, i, k, eta, dat)
```

## Arguments

theta	value of the error profile random effect at which the log.posterior distribution is calculated.
deltaj	vector of hospital random effect values.
sigma2	scale parameter ( $> 0$ ).
i	error profile index for which the calculate of the log.posterior distribution is needed.
k	degrees of freedom ( $> 0$ , maybe non-integer). df = Inf is allowed.
eta	skewness parameter ( $> 0$ ).
dat	an object of class "mederrData".

## Details

For further details see Myers et al. (2011).

## Value

logp returns a vector of log-posterior values.

## Author(s)

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Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[bhm.constr.resamp](#), [bhm.mcmc](#), [bhm.resample](#).

---

logunpost

*Unnormalized Marginal Posterior Distributions for k and  $\eta$*

---

## Description

This function computes the unnormalized marginal posterior distributions for the  $k$  and  $\eta$  parameters as described in Myers et al (2011).

## Usage

```
logunpost(resample)
```

## Arguments

resample      an object of class "mederrResample".

## Details

logunpost is used in the plot method for a `mederrResample` object.

## Value

logunpost returns an array with the posterior distribution values.

## Author(s)

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

`bhm.constr.resamp`, `bhm.mcmc`, `bhm.resample`, `mederrResample`.

---

<b>mederrData-class</b>	<i>Class "mederrData". Data Specification for Identifying the Most Harmful MEDication Errors using a Bayesian Hierarchical Model.</i>
-------------------------	---

---

## Description

This class encapsulates the data specification for a Bayesian Hierarchical Model used to identify the most harmful medication errors as described in Myers et al. (2011).

## Objects from the Class

Objects can be created by calls of the form `new("mederrData", data)`, where the `data` argument has to be a matrix or a data frame object that contains the following (*numeric*) information for each error profile/hospital combination:

1. the number of times ( $y$ ) that profile  $i$  in hospital  $j$  was reported with harm;
2. the total number of times ( $N$ ) that the error profile  $i$  is cited on a report from hospital  $j$ ,
3. the error profile  $i$  identification code,
4. the hospital  $j$  identification code.

## Slots

**data:** Object of class "data.frame"; data in the standard [data.frame](#) form.  
**size:** Object of class "numeric"; total number of observations in the data set.  
**numi:** Object of class "numeric"; number of error profiles available in the data set.  
**numj:** Object of class "numeric"; number of hospitals available in the data set.

## Methods

**plot** `signature(x = "mederrData", y = "missing")`: Provides a pictorial representation for a sample of error profiles reported by some hospitals.  
**summary** `signature(object = "mederrData")`: Summarizes information about an [mederrData](#) object.

## Author(s)

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bayes.rank](#), [bhm.mcmc](#), [bhm.resample](#), [mixnegbinom.em](#).

**Examples**

```
ng <- 50
i <- rep(1:ng, ng)
j <- rep(1:ng, each = ng)
N <- rpois(ng^2, 3 + .05*i - .01*j) + 1

theta_i <- rgamma(ng, 5, 5) - 4/5
delta_j <- rnorm(ng, 0, .2)
logit <- -3 + theta_i[i] + delta_j[j]
y <- rbinom(ng^2, N, exp(logit)/(1 + exp(logit)))

simdata <- new("mederrData", data = cbind(y, N, i, j))
```

**mederrFit-class**

*Class "mederrFit". Simulated Monte Carlo Chains (Step 1) for the Bayesian Hierarchical Model Used to Identify the Most Harmful Medication Errors.*

**Description**

This class encapsulates the simulated Monte Carlo chains for the Bayesian Hierarchical Model as described in Myers et al. (2011) forcing a symmetric normal distribution on the  $\theta_i$ ,  $i = 1, \dots, n$ .

**Objects from the Class**

Objects can be created by calls of the form `new("mederrFit", thetai, deltaj, gamma, sigma2, tau2, p.acc.i, p.acc.j, tune.theta, tune.delta, k, eta)`, but most often as the result of a call to [bhm.mcmc](#) or to [bhm.constr.resamp](#).

**Slots**

- thetai:** Object of class "matrix"; simulated chains for the  $\theta_i$ ,  $i = 1, \dots, n$ , error profiles random effects; see [bhm.mcmc](#).
- deltaj:** Object of class "matrix"; simulated chains for the  $\delta_j$ ,  $i = j, \dots, J$ , hospitals random effects; see [bhm.mcmc](#).
- gamma:** Object of class "numeric"; simulated chain for the  $\gamma$  parameter; see [bhm.mcmc](#).
- sigma2:** Object of class "numeric"; simulated chain for the  $\sigma^2$  parameter; see [bhm.mcmc](#).
- tau2:** Object of class "numeric"; simulated chain for the  $\tau^2$  parameter; see [bhm.mcmc](#).
- p.acc.i:** Object of class "numeric"; acceptance rates for the error profiles random effects.
- p.acc.j:** Object of class "numeric"; acceptance rates for the hospitals random effects.
- tune.theta:** Object of class "numeric"; last updated values of the  $\theta_i$  working variances for the Metropolis step.

**tune.delta:** Object of class "numeric"; last updated values of the  $\delta_j$  working variances for the Metropolis step.

**k:** Object of class "numeric";  $k$  value used in the simulation.

**eta:** Object of class "numeric";  $\eta$  value used in the simulation.

## Methods

**plot** signature(x = "mederrFit", y = "mederrFit"): Provides a graphical representation of the estimates.

**summary** signature(object = "mederrFit"): Summarizes the information regarding the estimates.

## Author(s)

Sergio Venturini <sergio.venturini@unicatt.it>,

Jessica A. Myers <jmyers6@partners.org>

## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[bayes.rank](#), [bhm.constr.resamp](#), [bhm.mcmc](#).

**mederrResample-class** *Class "mederrResample". Simulated Monte Carlo Chains (Step 2) for the Bayesian Hierarchical Model Used to Identify the Most Harmful Medication Errors.*

## Description

This class encapsulates the information needed to resample the Monte Carlo chains for the Bayesian Hierarchical Model as described in Myers et al. (2011) using user defined values for  $k$  and  $\eta$ .

## Objects from the Class

Objects can be created by calls of the form `new("mederrResample", log.ir, samp, A, t.new, t.old, grd)`, but most often as the result of a call to [bhm.resample](#).

**Slots**

- log.ir:** Object of class "array"; logarithm of the importance ratio for each pair of  $(k, \eta)$  values.
- samp:** Object of class "array"; resampled MCMC simulation indexes.
- A:** Object of class "array"; transformation ratio for each pair of  $(k, \eta)$  values.
- t.new:** Object of class "array";  $\theta_i$  posterior modes using  $(k = \infty, \eta = 1)$ .
- t.old:** Object of class "numeric";  $\theta_i$  posterior modes using user defined  $(k, \eta)$  values.
- grd:** Object of class "list"; grid of required  $(k, \eta)$  values.

**Methods**

- plot** signature(x = "mederrResample", y = "missing"): : Provides a graphical representation of a [mederrResample](#) object.

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bayes.rank](#), [bhm.constr.resamp](#), [bhm.mcmc](#).

MEDMARX

*Subset of the MEDMARX Data***Description**

Subset of the MEDMARX data included for illustrative purposes only in the [mederrRank](#) package.

**Usage**

```
data(MEDMARX)
```

**Format**

An object of class [mederrData](#).

**Details**

The data contained in this object are reproduced by gentle permission of Quantros, Inc., 690 N. McCarthy Blvd., Suite 200, Milpitas, CA 95035.

**Author(s)**

Sergio Venturini <sergio.venturini@unicatt.it>,  
 Jessica A. Myers <jmyers6@partners.org>

**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bhm.mcmc](#), [mederrData](#), [mederrFit](#).

**Examples**

```
data("MEDMARX", package = "mederrRank")
summary(MEDMARX)
plot(MEDMARX, nbins.err = 20, nbins.hosp = 10)
```

**mixnegbinom.em**

*Expectation-Maximization Algorithm for the Mixture of Negative Binomial Distributions*

**Description**

This function provides the empirical Bayes estimates for the parameters theta of a mixture of two negative binomial distributions (see [dmixnegbinom](#)) using an Expectation-Maximization algorithm.

**Usage**

```
mixnegbinom.em(dat, theta0, maxiter = 50000, toler = 0.01,
se = TRUE, stratified = FALSE)
```

**Arguments**

<b>dat</b>	an object of class "mederrData".
<b>theta0</b>	initial values for the parameters to be optimized over.
<b>maxiter</b>	a positive integer specifying the maximum number of iterations to be performed before the program is terminated.
<b>toler</b>	a positive scalar giving the tolerance at which the change in the log-likelihood is considered close enough to zero to terminate the algorithm.
<b>se</b>	logical; if TRUE the standard errors of the estimates are also returned.
<b>stratified</b>	logical; if TRUE the analysis will be performed by stratifying on the hospitals.

## Details

For further details see Myers et al. (2011).

## Value

`mixnegbinom.em` returns a list with components:

<code>theta.hat</code>	The best set of parameters found.
<code>final.err</code>	The last change in the log-likelihood; it has to be smaller than the <code>toler</code> argument.
<code>final.ll</code>	The likelihood value corresponding to <code>theta.hat</code> .
<code>final.score</code>	The log-likelihood score value corresponding to <code>theta.hat</code> .
<code>num.iter</code>	The number of iterations performed to find the proposed solution.
<code>se</code>	Only if argument <code>se</code> is true. A vector of estimates standard errors for the solution found.
<code>N</code>	The vector of observed error profiles counts.
<code>E</code>	The vector of expected error profiles counts.
<code>prior</code>	A character string giving the prior used; for this function is set to "mixgamma", i.e. a mixture of two gamma distributions as in DuMouchel (1999).

## Author(s)

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

- DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.
- Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[dmixnegbinom](#), [EBGM](#), [negbinom.em](#).

## Examples

```
## Not run:
data("simdata", package = "mederrRank")
summary(simdata)

fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
resamp <- bhm.resample(fit, simdata, p.resample = .1,
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)
```

```

plot(fit, fit2, simdata)

theta0 <- c(10, 6, 100, 100, .1)
ans <- mixnegbinom.em(simdata, theta0, 50000, 0.01,
se = TRUE, stratified = TRUE)
ans$theta
ans$se

summary(fit2, ans, simdata)
## End(Not run)

```

**mixnegbinom.loglik***Log-Likelihood Function for the Mixture of Negative Binomial Distributions*

## Description

This function computes the log-likelihood function for the mixture of two negative binomial distributions as described in [dmixnegbinom](#).

## Usage

```
mixnegbinom.loglik(theta, N, E)
```

## Arguments

theta	vector of parameter values.
N	vector of observed error profiles counts.
E	vector of expected error profiles counts.

## Details

For further details see Myers et al. (2011).

## Value

`mixnegbinom.loglik` returns the log-likelihood value for the negative binomial mixture.

## Author(s)

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

- DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.
- Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[dmixnegbinom](#), [mixnegbinom.em](#), [mixnegbinom.score](#).

`mixnegbinom.score`

*Log-Likelihood Score Function for the Mixture of Negative Binomial Distributions*

**Description**

This function computes the log-likelihood score for the mixture of two negative binomial distributions as described in [dmixnegbinom](#).

**Usage**

```
mixnegbinom.score(theta, N, E)
```

**Arguments**

<code>theta</code>	vector of parameter values.
<code>N</code>	vector of observed error profiles counts.
<code>E</code>	vector of expected error profiles counts.

**Details**

For further details see Myers et al. (2011).

**Value**

`mixnegbinom.score` returns the vector of log-likelihood score values for the negative binomial mixture.

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[dmixnegbinom](#), [mixnegbinom.em](#), [mixnegbinom.loglik](#).

negbinom.em

*Expectation-Maximization Algorithm for the Negative Binomial Distribution*

## Description

This function provides the empirical Bayes estimates for the parameters theta of a negative binomial distribution (see [dnegbinom](#)) using an Expectation-Maximization algorithm.

## Usage

```
negbinom.em(dat, theta0, maxiter = 50000, toler = 0.01,
se = TRUE, stratified = FALSE)
```

## Arguments

<code>dat</code>	an object of <a href="#">class "mederrData"</a> .
<code>theta0</code>	initial values for the parameters to be optimized over.
<code>maxiter</code>	a positive integer specifying the maximum number of iterations to be performed before the program is terminated.
<code>toler</code>	a positive scalar giving the tolerance at which the scaled gradient is considered close enough to zero to terminate the algorithm.
<code>se</code>	logical; if TRUE the standard errors of the estimates are also returned.
<code>stratified</code>	logical; if TRUE the analysis will be performed by stratifying on the hospitals.

## Details

For further details see Myers et al. (2011).

## Value

`negbinom.em` returns a list with components:

<code>theta.hat</code>	The best set of parameters found.
<code>final.err</code>	The last change in the log-likelihood; it has to be smaller than the <code>toler</code> argument.
<code>final.ll</code>	The likelihood value corresponding to <code>theta.hat</code> .
<code>final.score</code>	The log-likelihood score value corresponding to <code>theta.hat</code> .
<code>num.iter</code>	The number of iterations performed to find the proposed solution.
<code>se</code>	Only if argument <code>se</code> is true. A vector of estimates standard errors for the solution found.
<code>N</code>	The vector of observed error profiles counts.
<code>E</code>	The vector of expected error profiles counts.
<code>prior</code>	A character string giving the prior used; for this function is set to "gamma", i.e. a gamma distribution.

**Author(s)**

Sergio Venturini <sergio.venturini@unicatt.it>,  
 Jessica A. Myers <jmyers6@partners.org>

**References**

- DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.
- Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[dnegbinom](#), [EBGM](#), [mixnegbinom.em](#).

**Examples**

```
data("simdata", package = "mederrRank")
summary(simdata)

## Not run:
fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
resamp <- bhm.resample(fit, simdata, p.resample = .1,
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)
plot(fit, fit2, simdata)

## End(Not run)

theta0 <- runif(2, 0, 5)
ans <- negbinom.em(simdata, theta0, 50000, 0.01,
se = TRUE, stratified = TRUE)
ans$theta
ans$se

## Not run:
summary(fit2, ans, simdata)

## End(Not run)
```

**negbinom.loglik**

*Log-Likelihood Function for the Mixture of Negative Binomial Distributions*

**Description**

This function computes the log-likelihood function for the mixture of two negative binomial distribution as described in [dmixnegbinom](#).

**Usage**

```
negbinom.loglik(theta, N, E)
```

**Arguments**

theta	vector of parameter values.
N	vector of observed error profiles counts.
E	vector of expected error profiles counts.

**Details**

For further details see Myers et al. (2011).

**Value**

`negbinom.loglik` returns the log-likelihood value for the negative binomial distribution.

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

- DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.
- Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[dnegbinom](#), [negbinom.em](#), [negbinom.score](#).

[negbinom.score](#)

*Log-Likelihood Score Function for the Negative Binomial Distribution*

**Description**

This function computes the log-likelihood score for the negative binomial distribution as described in [dmixnegbinom](#).

**Usage**

```
negbinom.score(theta, N, E)
```

**Arguments**

theta	vector of parameter values.
N	vector of observed error profiles counts.
E	vector of expected error profiles counts.

**Details**

For further details see Myers et al. (2011).

**Value**

`negbinom.score` returns the vector of log-likelihood score values for the negative binomial distribution.

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

- DuMouchel W. (1999), "Bayesian Data Mining in Large Frequency Tables, with an Application to the FDA Spontaneous Reporting System". *The American Statistician*, 53, 177-190.
- Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[dnegbinom](#), [negbinom.em](#), [negbinom.loglik](#).

---

*p.value*

*Posterior Predictive Test statistics*

---

**Description**

This function computes posterior predictive test statistics as described in Myers et al. (2011).

**Usage**

`p.value(reps)`

**Arguments**

reps	list of replications created with the <code>post.rep</code> function.
------	---

## Details

For further details see Myers et al. (2011).

## Value

`p.value` creates a list of p-values.

## Author(s)

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[bhm.constr.resamp](#), [bhm.mcmc](#), [bhm.resample](#), [post.rep](#).

## Examples

```
## Not run:
data("simdata", package = "mederrRank")
summary(simdata)

fit <- bhm.mcmc(simdata, nsim = 1000, burnin = 500, scale.factor = 1.1)
resamp <- bhm.resample(fit, simdata, p.resample = .1,
k = c(3, 6, 10, 30, 60, Inf), eta = c(.5, .8, 1, 1.25, 2))
fit2 <- bhm.constr.resamp(fit, resamp, k = 3, eta = .8)
reps <- post.rep(fit2, simdata)
pvalues <- p.value(reps)

## End(Not run)
```

## Description

Methods for function `plot` in Package ‘graphics’ to be used with “[mederrData](#)”, [mederrFit](#) and “[mederrResample](#)” objects.

## Methods

```
signature(x = "mederrData", y = "missing") Pictorial representation for a "mederrData" object.  
signature(x = "mederrFit", y = "mederrFit") Graphical representation of Markov Chain Monte Carlo simulations for a "mederrFit" object.  
signature(x = "mederrResample", y = "missing") Graphical representation of the resampling transformation for a "mederrResample" object.
```

## Author(s)

Sergio Venturini <sergio.venturini@unicatt.it>,  
Jessica A. Myers <jmyers6@partners.org>

## References

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

## See Also

[bhm.mcmc](#), [bhm.resample](#).

---

post.rep

*Posterior Predictive Data Replications*

---

## Description

This function creates a list of replicated data for posterior predictive checking as described in Myers et al. (2011).

## Usage

```
post.rep(model, dat)
```

## Arguments

model	an object of class "mederrFit".
dat	an object of class "mederrData".

## Details

For further details see Myers et al. (2011).

## Value

post.rep returns a list of replicated data.

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bhm.constr.resamp](#), [bhm.mcmc](#), [bhm.resample](#), [p.value](#).

*rmixnegbinom*

*The Negative Binomial Mixture Distribution*

**Description**

Random generation for a mixture of two negative binomial distributions.

**Usage**

```
rmixnegbinom(n, theta, E)
```

**Arguments**

- |       |  |
|-------|--|
| n     | number of observations.  |
| theta | vector of parameters for the negative binomial distribution mixture. |
| E     | vector of (non-negative integer) expected counts.                    |

**Value**

*rmixnegbinom* generates random deviates corresponding to the E and theta values provided.

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
 Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[dmixnegbinom](#), [rnbinom](#).

---

simdata

*Simulated Data*

---

**Description**

Simulated data to use for illustrative purposes in the [mederrRank](#) package.

**Usage**

```
data(simdata)
```

**Format**

An object of class [mederrData](#).

**Author(s)**

Sergio Venturini <[sergio.venturini@unicatt.it](mailto:sergio.venturini@unicatt.it)>,  
Jessica A. Myers <[jmyers6@partners.org](mailto:jmyers6@partners.org)>

**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), "Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database". Technical Report.

**See Also**

[bhm.mcmc](#), [mederrData](#), [mederrFit](#).

**Examples**

```
data("simdata", package = "mederrRank")
summary(simdata)
plot(simdata)
```

**Description**

Methods for function `summary` in Package ‘base’ to be used with “[mederrData](#)” and “[mederrFit](#)” objects.

**Methods**

`signature(object = "mederrData")` Extracts summary information about the slots of a “[mederrData](#)” object.

`signature(object = "mederrFit")` Extracts summary information about the slots of a “[mederrFit](#)” object.

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**References**

Myers, J. A., Venturini, S., Dominici, F. and Morlock, L. (2011), “Random Effects Models for Identifying the Most Harmful Medication Errors in a Large, Voluntary Reporting Database”. Technical Report.

**See Also**

[bayes.rank](#), [bhm.mcmc](#).

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