

# Package: semisup (via r-universe)

June 27, 2024

**Version** 1.29.0

**Title** Semi-Supervised Mixture Model

**Description** Implements a parametric semi-supervised mixture model. The permutation test detects markers with main or interactive effects, without distinguishing them. Possible applications include genome-wide association analysis and differential expression analysis.

**biocViews** SNP, GenomicVariation, SomaticMutation, Genetics, Classification, Clustering, DNaseq, Microarray, MultipleComparison

**Depends** R (>= 3.0.0)

**Imports** VGAM

**Suggests** knitr, testthat, SummarizedExperiment

**VignetteBuilder** knitr

**License** GPL-3

**LazyData** true

**RoxygenNote** 7.0.0

**URL** <https://github.com/rauschenberger/semisup>

**BugReports** <https://github.com/rauschenberger/semisup/issues>

**Repository** <https://bioc.r-universe.dev>

**RemoteUrl** <https://github.com/bioc/semisup>

**RemoteRef** HEAD

**RemoteSha** 2856f5cc370e9fe2bd6e9f2172f06696800c033b

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`semisup-package`      *Semi-supervised mixture model*

## Description

This R package implements the semi-supervised mixture model. Use `mixtura` for model fitting, and `scrutor` for hypothesis testing.

## Getting started

Please type the following commands:

```
utils::vignette("semisup")
?semisup::mixtura
?semisup::scrutor
```

## More information

A Rauschenberger, RX Menezes, MA van de Wiel, NM van Schoor, and MA Jonker (2020). "Semi-supervised mixture test for detecting markers associated with a quantitative trait", *Manuscript in preparation*.

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`mixtura`      *Model fitting*

## Description

This function fits a semi-supervised mixture model. It simultaneously estimates two mixture components, and assigns the unlabelled observations to these.

## Usage

```
mixtura(y, z, dist = "norm",
        phi = NULL, pi = NULL, gamma = NULL,
        test = NULL, iter = 100, kind = 0.05,
        debug = TRUE, ...)
```

## Arguments

<code>y</code>	<b>observations:</b> numeric vector of length n
<code>z</code>	<b>class labels:</b> integer vector of length n, with entries 0, 1 and NA
<code>dist</code>	<b>distributional assumption:</b> character "norm" (Gaussian), "nbinom" (negative binomial), or "zinb" (zero-inflated negative binomial)
<code>phi</code>	<b>dispersion parameters:</b> numeric vector of length q, or NULL

pi	zero-inflation parameter(s): numeric vector of length q, or NULL
gamma	offset: numeric vector of length n, or NULL
test	resampling procedure: character "perm" (permutation) or "boot" (parametric bootstrap), or NULL
iter	(maximum) number of resampling iterations : positive integer, or NULL
kind	resampling accuracy: numeric between 0 and 1, or NULL; all p-values above kind are approximate
debug	verification of arguments: TRUE or FALSE
...	settings EM algorithm: starts, it.em and epsilon (see <a href="#">arguments</a> )

## Details

By default, phi and pi are estimated by the maximum likelihood method, and gamma is replaced by a vector of ones.

## Value

This function fits and compares a one-component ( $H_0$ ) and a two-component ( $H_1$ ) mixture model.

posterior	probability of belonging to class 1: numeric vector of length n
converge	path of the log-likelihood: numeric vector with maximum length it.em
estim0	parameter estimates under $H_0$ : data frame
estim1	parameter estimates under $H_1$ : data frame
loglik0	log-likelihood under $H_0$ : numeric
loglik1	log-likelihood under $H_1$ : numeric
lrts	likelihood-ratio test statistic: positive numeric
p.value	$H_0$ versus $H_1$ : numeric between 0 and 1, or NULL

## Reference

A Rauschenberger, RX Menezes, MA van de Wiel, NM van Schoor, and MA Jonker (2020). "Semi-supervised mixture test for detecting markers associated with a quantitative trait", *Manuscript in preparation*.

## See Also

Use [scrutor](#) for hypothesis testing. All other functions are [internal](#).

## Examples

```
# data simulation
n <- 100
z <- rep(0:1,each=n/2)
y <- rnorm(n=n,mean=2,sd=1)
z[(n/4):n] <- NA
```

```
# model fitting
mixtura(y,z,dist="norm",test="perm")
```

scrutor

*Hypothesis testing*

## Description

This function tests whether the unlabelled observations come from a mixture of two distributions.

## Usage

```
scrutor(Y, Z, dist = "norm",
       phi = NULL, pi = NULL, gamma = NULL,
       test = "perm", iter = NULL, kind = NULL,
       debug = TRUE, ...)
```

## Arguments

Y	<b>observations:</b> numeric vector of length n, or numeric matrix with n rows (samples) and q columns (variables)
Z	<b>class labels:</b> numeric vector of length n, or numeric matrix with n rows (samples) and p columns (variables), with entries 0 and NA
dist	distributional assumption: character "norm" (Gaussian), "nbinom" (negative binomial), or "zinb" (zero-inflated negative binomial)
phi	dispersion parameter(s): numeric vector of length q, or NULL (norm: none, nbinom: MLE)
pi	zero-inflation parameter(s): numeric vector of length q, or NULL (norm: none,nbinom: MLE)
gamma	offset: numeric vector of length n, or NULL
test	resampling procedure: character "perm" (permutation) or "boot" (parametric bootstrap), or NULL
iter	(maximum) number of resampling iterations : positive integer, or NULL
kind	resampling accuracy: numeric between 0 and 1, or NULL; all p-values above kind are approximate
debug	verification of arguments: TRUE or FALSE
...	settings EM algorithm: starts, it.em and epsilon (see <a href="#">arguments</a> )

## Details

By default, phi and pi are estimated by the maximum likelihood method, and gamma is replaced by a vector of ones.

**Value**

This function tests a one-component ( $H_0$ ) against a two-component mixture model ( $H_1$ ).

y	index observations
z	index class labels
lrts	test statistic
p.value	p-value

**Reference**

A Rauschenberger, RX Menezes, MA van de Wiel, NM van Schoor, and MA Jonker (2020). "Semi-supervised mixture test for detecting markers associated with a quantitative trait", *Manuscript in preparation*.

**See Also**

Use [mixtura](#) for model fitting. All other functions are [internal](#).

**Examples**

```
# data simulation
n <- 100
z <- rep(0:1,each=n/2)
y <- rnorm(n=n,mean=2*z,sd=1)
z[(n/4):n] <- NA

# hypothesis testing
scrutor(y,z,dist="norm")
```

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