

Package: MRS (via r-universe)

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Type Package

Title Multi-Resolution Scanning for Cross-Sample and Cross-Group Differences

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Description Implements the multi-resolution scanning (MRS) method for cross-sample distribution comparisons, as described in Soriano and Ma (2017) <[doi:10.1111/rssb.12180](https://doi.org/10.1111/rssb.12180)>, and the analysis of distributional variation (ANDOVA) method for cross-group comparisons introduced in Ma and Soriano (2018) <[doi:10.1080/10618600.2017.1402774](https://doi.org/10.1080/10618600.2017.1402774)>. Both methods use nonparametric models on multi-resolution partition trees to detect and characterize differences among distributions, with tools for visualizing the results.

License GPL (>= 3)

URL <https://github.com/MaStatLab/MRS>

BugReports <https://github.com/MaStatLab/MRS/issues>

Encoding UTF-8

Imports Rcpp (>= 1.0.0)

Suggests KernSmooth, testthat (>= 3.0.0)

LinkingTo Rcpp, RcppArmadillo

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andova	<i>Multi Resolution Scanning for one-way ANDOVA using the multi-scale Beta-Binomial model</i>
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Description

This function executes the Multi Resolution Scanning algorithm to detect differences across the distributions of multiple groups having multiple replicates.

Usage

```
andova(  
  X,  
  G,  
  H,  
  n_groups = length(unique(G)),  
  n_subgroups = NULL,  
  Omega = "default",  
  K = 6,  
  init_state = c(0.8, 0.2, 0),  
  beta = 1,  
  gamma = 0.07,  
  delta = 0.4,  
  eta = 0,  
  alpha = 0.5,  
  nu_vec = 10^(seq(-1, 4)),  
  return_global_null = TRUE,  
  return_tree = TRUE,  
  n_post_samples = 0,  
  baseline = 0,  
  method = "newton",  
  n_grid_theta = 4  
)
```

Arguments

X	Matrix of the data. Each row represents an observation.
G	Numeric vector of the group label of each observation. Labels are integers starting from 1.
H	Numeric vector of the replicate label of each observation. Labels are integers starting from 1.
n_groups	Number of groups.
n_subgroups	Vector indicating the number of replicates for each group.
Omega	Matrix defining the vertices of the sample space. The "default" option defines a hyperrectangle containing all the data points. Otherwise the user can define a matrix where each row represents a dimension, and the two columns contain the associated lower and upper limit.
K	Depth of the tree. Default is $K = 6$, while the maximum is $K = 14$.
init_state	Initial state of the hidden Markov process. The three states are <i>null</i> , <i>alternative</i> and <i>prune</i> , respectively.
beta	Spatial clustering parameter of the transition probability matrix. Default is $\beta = 1.0$.
gamma	Parameter of the transition probability matrix. Default is $\gamma = 0.07$.
delta	Parameter of the transition probability matrix. Default is $\delta = 0.4$.
eta	Parameter of the transition probability matrix. Default is $\eta = 0.0$.
alpha	Pseudo-counts of the Beta random probability assignments.
nu_vec	The support of the discrete uniform prior on ν .
return_global_null	Boolean indicating whether to return the marginal posterior probability of the global null.
return_tree	Boolean indicating whether to return the posterior representative tree.
n_post_samples	Number of posterior trees to sample. Default is zero.
baseline	Group used as the baseline for effect sizes. A value of zero uses the pooled mean as the baseline.
method	Numerical integration method, either "newton" or "riemann".
n_grid_theta	Number of grid points used by the numerical integration. The default of four uses the same midpoint quadrature rule as version 1.2.6.

Value

An mrs object.

References

Ma L. and Soriano J. (2018). Analysis of distributional variation through multi-scale Beta-Binomial modeling. *Journal of Computational and Graphical Statistics*, 27(3), 529–541. doi:10.1080/10618600.2017.1402774

Examples

```

set.seed(12345)
n = 1000
M = 5
class_1 = sample(M, n, prob= 1:5, replace=TRUE )
class_2 = sample(M, n, prob = 5:1, replace=TRUE )

Y_1 = rnorm(n, mean=class_1, sd = .2)
Y_2 = rnorm(n, mean=class_2, sd = .2)

X = matrix( c(Y_1, Y_2), ncol = 1)
G = c(rep(1,n),rep(2,n))
H = sample(3,2*n, replace = TRUE )

ans = andova(X, G, H)
ans$PostGlobNull
plot1D(ans)

```

mrs

Multi Resolution Scanning

Description

This function executes the Multi Resolution Scanning algorithm to detect differences across multiple distributions.

Usage

```

mrs(
  X,
  G,
  n_groups = length(unique(G)),
  Omega = "default",
  K = 6,
  init_state = NULL,
  beta = 1,
  gamma = 0.3,
  delta = NULL,
  eta = 0.3,
  alpha = 0.5,
  return_global_null = TRUE,
  return_tree = TRUE,
  n_post_samples = 0,
  baseline = 0,
  min_n_node = 0
)

```

Arguments

X	Matrix of the data. Each row represents an observation.
G	Numeric vector of the group label of each observation. Labels are integers starting from 1.
n_groups	Number of groups.
Omega	Matrix defining the vertices of the sample space. The "default" option defines a hyperrectangle containing all the data points. Otherwise the user can define a matrix where each row represents a dimension, and the two columns contain the associated lower and upper limits for each dimension.
K	Depth of the tree. Default is $K = 6$, while the maximum is $K = 14$.
init_state	Initial state of the hidden Markov process. The three states are <i>null</i> , <i>alternative</i> and <i>prune</i> , respectively.
beta	Spatial clustering parameter of the transition probability matrix. Default is $\beta = 1$.
gamma	Parameter of the transition probability matrix. Default is $\gamma = 0.3$.
delta	Optional parameter of the transition probability matrix. Default is $\delta = \text{NULL}$.
eta	Parameter of the transition probability matrix. Default is $\eta = 0.3$.
alpha	Pseudo-counts of the Beta random probability assignments. Default is $\alpha = 0.5$.
return_global_null	Boolean indicating whether to return the posterior probability of the global null hypothesis.
return_tree	Boolean indicating whether to return the posterior representative tree.
n_post_samples	Number of posterior trees to sample. Default is zero.
baseline	Group used as the baseline for effect sizes. A value of zero uses the pooled mean as the baseline.
min_n_node	Node in the tree is returned if there are more than <code>min_n_node</code> data-points in it.

Value

An `mrs` object.

References

Soriano J. and Ma L. (2017). Probabilistic multi-resolution scanning for two-sample differences. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*. doi:[10.1111/rssb.12180](https://doi.org/10.1111/rssb.12180)

Examples

```
set.seed(12345)
n = 20
p = 2
X = matrix(c(runif(p*n/2),rbeta(p*n/2, 1, 4)), nrow=n, byrow=TRUE)
G = c(rep(1,n/2), rep(2,n/2))
ans = mrs(X=X, G=G)
```

plot1D

Plot regions of the representative tree in 1D

Description

This function visualizes the regions of the representative tree of the output of the [mrs](#) function. For each region the posterior probability of difference (PMAP) or the effect size is plotted.

Usage

```
plot1D(
  ans,
  type = "prob",
  group = 1,
  dim = 1,
  regions = rep(1, length(ans$RepresentativeTree$Levels)),
  legend = FALSE,
  main = "default",
  abs = TRUE,
  eff_scale = "lor"
)
```

Arguments

ans	An mrs object.
type	What is represented at each node. The options are type = c("eff", "prob"). Default is type = "prob".
group	If type = "eff", which group effect size is used. Default is group = 1.
dim	If the data are multivariate, dim is the dimension plotted. Default is dim = 1.
regions	Binary vector indicating the regions to plot. The default is to plot all regions.
legend	Color legend for type. Default is legend = FALSE.
main	Overall title for the plot.
abs	If TRUE, plot the absolute value of the effect size. Only used when type = "eff".
eff_scale	Scale used for effect-size legend labels: "lor" for log odds ratios or "odds-ratio" for odds ratios.

Value

Invisibly returns NULL. The function is called for its side effect of drawing a one-dimensional multiresolution map on the current graphics device. The plotted regions encode posterior probabilities of a difference when type = "prob", or the selected group's effect sizes when type = "eff".

References

- Soriano J. and Ma L. (2017). Probabilistic multi-resolution scanning for two-sample differences. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*. doi:10.1111/rssb.12180
- Ma L. and Soriano J. (2018). Analysis of distributional variation through multi-scale Beta-Binomial modeling. *Journal of Computational and Graphical Statistics*, 27(3), 529–541. doi:10.1080/10618600.2017.1402774

Examples

```
set.seed(12345)
p = 1
n1 = 200
n2 = 200
mu1 = matrix( c(0,10), nrow = 2, byrow = TRUE)
mu2 = mu1; mu2[2] = mu1[2] + .01
sigma = c(1,.1)

Z1 = sample(2, n1, replace=TRUE, prob=c(0.9, 0.1))
Z2 = sample(2, n2, replace=TRUE, prob=c(0.9, 0.1))
X1 = mu1[Z1] + matrix(rnorm(n1*p), ncol=p)*sigma[Z1]
X2 = mu2[Z2] + matrix(rnorm(n2*p), ncol=p)*sigma[Z1]
X = rbind(X1, X2)
G = c(rep(1, n1), rep(2,n2))

ans = mrs(X, G, K=10)
plot1D(ans, type = "prob")
plot1D(ans, type = "eff")
```

plot1DSigWindows	<i>Plot one-dimensional windows selected by Bayesian FDR</i>
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Description

Selects representative-tree windows whose posterior alternative probabilities satisfy a requested Bayesian false discovery rate, then displays them with [plot1D](#).

Usage

```
plot1DSigWindows(ans_mrs, fdr = 0.1, precision = 0.001, ...)
```

Arguments

ans_mrs	An mrs object.
fdr	Desired Bayesian false discovery rate between zero and one.
precision	Step size used to search posterior-probability thresholds.
...	Additional arguments passed to plot1D .

Value

Invisibly, a list containing the selected node indices and threshold.

plot2D

Plot regions of the representative tree in 2D

Description

This function visualizes the regions of the representative tree of the output of the [mrs](#) function.

Usage

```
plot2D(
  ans,
  type = "prob",
  data.points = "all",
  background = "none",
  group = 1,
  dim = c(1, 2),
  levels = sort(unique(ans$RepresentativeTree$Levels)),
  regions = rep(1, length(ans$RepresentativeTree$Levels)),
  legend = FALSE,
  main = "default",
  abs = TRUE
)
```

Arguments

ans	An mrs object.
type	Different options on how to visualize the rectangular regions. The options are <code>type = c("eff", "prob", "empty", "none")</code> . Default is <code>type = "prob"</code> .
data.points	Different options on how to plot the data points. The options are <code>data.points = c("all", "differential", "none")</code> . Default is <code>data.points = "all"</code> .
background	Different options on the background. The options are <code>background = c("smeared", "none")</code> .
group	If <code>type = "eff"</code> , which group effect size is used. Default is <code>group = 1</code> .
dim	If the data are multivariate, <code>dim</code> are the two dimensions plotted. Default is <code>dim = c(1, 2)</code> .
levels	Vector with the level of the regions to plot. The default is to plot regions at all levels.
regions	Binary vector indicating the regions to plot. The default is to plot all regions.
legend	Color legend for type. Default is <code>legend = FALSE</code> .
main	Overall title for the legend.
abs	If TRUE, plot the absolute value of the effect size. Only used when <code>type = "eff"</code> .

Value

Invisibly returns NULL. The function is called for its side effect of drawing one panel for each selected tree level on the current graphics device. Rectangles encode posterior probabilities or effect sizes; optional points and a smoothed background show the observed data.

References

Soriano J. and Ma L. (2017). Probabilistic multi-resolution scanning for two-sample differences. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*. doi:[10.1111/rssb.12180](https://doi.org/10.1111/rssb.12180)

Ma L. and Soriano J. (2018). Analysis of distributional variation through multi-scale Beta-Binomial modeling. *Journal of Computational and Graphical Statistics*, 27(3), 529–541. doi:[10.1080/10618600.2017.1402774](https://doi.org/10.1080/10618600.2017.1402774)

Examples

```
set.seed(12345)
p = 2
n1 = 200
n2 = 200
mu1 = matrix( c(9,9,0,4,-2,-10,3,6,6,-10), nrow = 5, byrow=TRUE)
mu2 = mu1; mu2[2,] = mu1[2,] + 1

Z1 = sample(5, n1, replace=TRUE)
Z2 = sample(5, n2, replace=TRUE)
X1 = mu1[Z1,] + matrix(rnorm(n1*p), ncol=p)
X2 = mu2[Z2,] + matrix(rnorm(n2*p), ncol=p)
X = rbind(X1, X2)
colnames(X) = c(1,2)
G = c(rep(1, n1), rep(2,n2))

ans = mrs(X, G, K=8)
plot2D(ans, type = "prob", legend = TRUE)

plot2D(ans, type="empty", data.points = "differential",
  background = "none")

plot2D(ans, type="none", data.points = "differential",
  background = "smeared", levels = 4)
```

plotTree

Plot nodes of the representative tree

Description

This function visualizes the representative tree of the output of the [mrs](#) function. For each node of the representative tree, the posterior probability of difference (PMAP) or the effect size is plotted. Each node in the tree is associated to a region of the sample space. All non-terminal nodes have two children obtained by partitioning the parent region with a dyadic cut along a given direction. The numbers under the vertices represent the cutting direction.

Usage

```
plotTree(
  ans,
  type = "prob",
  group = 1,
  legend = FALSE,
  main = "",
  node.size = 5,
  abs = TRUE
)
```

Arguments

ans	A mrs object.
type	What is represented at each node. The options are <code>type = c("eff", "prob")</code> .
group	If <code>type = "eff"</code> , which group effect size is used.
legend	Color legend for type. Default is <code>legend = FALSE</code> .
main	Main title. Default is <code>main = ""</code> .
node.size	Size of the nodes. Default is <code>node.size = 5</code> .
abs	If <code>TRUE</code> , plot the absolute value of the effect size. Only used when <code>type = "eff"</code> .

Value

Invisibly returns `NULL`. The function is called for its side effect of drawing the representative binary partition tree on the current graphics device. Node size and color encode posterior probabilities or effect sizes, and labels beneath non-terminal nodes identify split directions.

References

Soriano J. and Ma L. (2017). Probabilistic multi-resolution scanning for two-sample differences. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*. doi:10.1111/rssb.12180

Ma L. and Soriano J. (2018). Analysis of distributional variation through multi-scale Beta-Binomial modeling. *Journal of Computational and Graphical Statistics*, 27(3), 529–541. doi:10.1080/10618600.2017.1402774

Examples

```
set.seed(12345)
p = 2
n1 = 200
n2 = 200
mu1 = matrix( c(9,9,0,4,-2,-10,3,6,6,-10), nrow = 5, byrow=TRUE)
mu2 = mu1; mu2[2,] = mu1[2,] + 1

Z1 = sample(5, n1, replace=TRUE)
Z2 = sample(5, n2, replace=TRUE)
X1 = mu1[Z1,] + matrix(rnorm(n1*p), ncol=p)
X2 = mu2[Z2,] + matrix(rnorm(n2*p), ncol=p)
```

```
X = rbind(X1, X2)
colnames(X) = c(1,2)
G = c(rep(1, n1), rep(2,n2))

ans = mrs(X, G, K=8)
plotTree(ans, type = "prob", legend = TRUE)
```

```
print.summary.mrs      Print summary of a mrs object
```

Description

This function prints a summary of the output of the [mrs](#) function. It provides the marginal prior and posterior of the null and the top regions of the representative tree.

Usage

```
## S3 method for class 'summary.mrs'
print(x, ...)
```

Arguments

x	A <code>summary.mrs</code> object
...	Additional print parameters.

Value

Invisibly returns NULL. The method is called for its side effect of printing the global prior and posterior null probabilities and, when present, the selected differential regions with their posterior alternative probabilities, effect sizes, region bounds, and split directions.

References

Soriano J. and Ma L. (2017). Probabilistic multi-resolution scanning for two-sample differences. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*. doi:[10.1111/rssb.12180](#)

Ma L. and Soriano J. (2018). Analysis of distributional variation through multi-scale Beta-Binomial modeling. *Journal of Computational and Graphical Statistics*, 27(3), 529–541. doi:[10.1080/10618600.2017.1402774](#)

Examples

```
set.seed(12345)
n = 100
p = 2
X = matrix(c(runif(p*n/2),rbeta(p*n/2, 1, 4)), nrow=n, byrow=TRUE)
G = c(rep(1,n/2), rep(2,n/2))
x = mrs(X=X, G=G)
fit = summary(x, rho = 0.95, abs_eff = 1)
print(fit)
```

summary.mrs	<i>Summary of a mrs object</i>
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Description

This function summarizes the output of the [mrs](#) function. It provides the marginal prior and posterior null and the top regions of the representative tree.

Usage

```
## S3 method for class 'mrs'
summary(object, rho = 0.5, abs_eff = 0, sort_by = "eff", ...)
```

Arguments

object	A mrs object
rho	Threshold for the posterior alternative probability. All regions with posterior alternative probability larger than rho are reported. Default is rho = 0.5.
abs_eff	Threshold for the effect size. All regions with effect size larger than abs_eff in absolute value are reported. Default is abs_eff = 0.
sort_by	Define in which order the regions are reported. The options are sort_by = c("eff", "prob") and the default is sort_by = "eff".
...	Additional summary parameters.

Value

A list with information about the top regions.

References

Soriano J. and Ma L. (2017). Probabilistic multi-resolution scanning for two-sample differences. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*. doi:[10.1111/rssb.12180](#)

Ma L. and Soriano J. (2018). Analysis of distributional variation through multi-scale Beta-Binomial modeling. *Journal of Computational and Graphical Statistics*, 27(3), 529–541. doi:[10.1080/10618600.2017.1402774](#)

Examples

```
set.seed(12345)
n = 100
p = 2
X = matrix(c(runif(p*n/2),rbeta(p*n/2, 1, 4)), nrow=n, byrow=TRUE)
G = c(rep(1,n/2), rep(2,n/2))
object = mrs(X=X, G=G)
fit = summary(object, rho = 0.5, abs_eff = 0.1)
```

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