Package ‘LPS’

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BugReports https://github.com/maressyl/R.LPS/issues
Description An implementation of the Linear Predictor Score approach, as initiated by Radmacher et al. (J Comput Biol 2001) and enhanced by Wright et al. (PNAS 2003) for gene expression signatures. Several tools for unsupervised clustering of gene expression data are also provided.
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clusterize

Hierarchical clustering heat maps

Description

This function draws a heat map ordered according to hierarchical clusterings, similarly to heatmap. It offers more control on layout and allows multiple row annotations.

hclust.ward is derivated from 'stats' package hclust, with an alternative default (as arguments can not be passed to it).

dist.COR mimics 'stats' package dist, computing distances as 1 - Pearson’s correlation coefficient.

Usage

```r
clusterize(expr, side = NULL, cex.col = NA, cex.row = NA, mai.left = NA, mai.bottom = NA, mai.right = 0.1, mai.top = 0.1, side.height = 1, side.col = NULL, side.srt = 0, side.cex = 1, col.heatmap = heat(), zlim = "0 centered", zlim.trim = 0.02, norm = c("rows", "columns", "none"), norm.clust = TRUE, norm.robust = FALSE, customLayout = FALSE, getLayout = FALSE, plot = TRUE, widths = c(1, 4), heights = c(1, 4), order.genes = NULL, order.samples = NULL, fun.dist = dist.COR, fun.hclust = hclust.ward, clust.genes = NULL, clust.samples = NULL)
cdist.COR(input)
hclust.ward(input)
```

Arguments

- `expr`: A numeric matrix, holding features (genes) in columns and observations (samples) in rows. Rows and columns will be ordered according to hierarchical clustering results.
- `side`, `cex.col`, `cex.row`, `mai.left`, `mai.bottom`, `mai.right`, `mai.top`, `side.height`, `side.col`, `side.srt`, `side.cex`: To be passed to heat.map.
- `col.heatmap`, `zlim`, `zlim.trim`, `norm`, `norm.clust`, `norm.robust`, `customLayout`, `getLayout`, `plot`, `widths`, `heights`: To be passed to heat.map.
clusterize invisibly returns the same list as heat.map, plus:

genes  The gene dendrogram.
samples The sample dendrogram.

See hclust and dist respectively for the other functions.
**Author(s)**

Sylvain Mareschal

**See Also**

heat.map, heatmap, hclust, dist

**Examples**

```r
# Data with features in columns
data(rosenwald)
group <- rosenwald$group
expr <- t(rosenwald.expr)[,1:100]

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[is.na(x)] <- round(mean(x, na.rm=TRUE), 3); x }
expr <- apply(expr, 2, f)

# Simple heat map
clusterize(expr)

# With annotation (row named data.frame)
side <- data.frame(group, row.names=rownames(expr))
clusterize(expr, side=side)
```

---

**heat**

Heatmap palette generation

**Description**

This function generates a ramp of colors for heat.map derivated functions.

**Usage**

```r
heat(colors = c("#8888FF", "#000000", "#FF4444"), n = 256, shapeFun = heat.exp, ...)
heat.exp(n, part, base = 1.015)
heat.lin(n, part)
```

**Arguments**

- `colors` Character vector of length 3, determining starting, middle and final colors.
- `n` Single integer value, amount of colors / values to generate.
- `shapeFun` Function taking at least 2 arguments: `n` and `part`. heat.exp and heat.lin are provided as examples.
- `...` Further arguments to heat will be passed to shapeFun.
- `part` Single integer, defined as 1 while generating colors between the first two boundaries, and 2 otherwise.
- `base` Single numeric value, base for exponential slope.
**Value**

heat returns a character vector of colors in hexadecimal representation.

heat.lin and heat.exp return n numeric values, defining a curve whose slope will be mimiced during color interpolation.

**Author(s)**

Sylvain Mareschal

**See Also**

colorRampPalette

heat.map, clusterize, predict.LPS

**Examples**

```
# Classical heatmap colors
palette <- heat(c("green", "black", "red"))
heat.scale(zlim=c(-2,2), col.heatmap=palette)

# Two distinct shapes provided
heat.scale(zlim=c(-2,2), col.heatmap=heat(shapeFun=heat.lin))
heat.scale(zlim=c(-2,2), col.heatmap=heat(shapeFun=heat.exp))
```

**Description**

This function draws a heatmap from a matrix, similarly to image. It also offers normalization and annotation features, with more control than heatmap.

side can provide multiple sample annotations, and are handled differently depending on their class:

- **numeric** are attributed grey shades from the minimum to the maximum, which are provided in the legend.
- **factor** have their levels attributed colors using a default or custom palette. Hexadecimal color codes starting with # and color names known by R are used "as is".
- **character** are printed as is in a blank cell. Hexadecimal color codes starting with # and color names known by R are used as background colors instead of text.
- **logical** are ploted in dark (TRUE) or light (FALSE) gray, leaving NAs in white.
heat.map(expr, side = NULL, cex.col = NA, cex.row = NA, mai.left = NA, mai.bottom = NA, mai.right = 0.1, mai.top = 0.1, side.height = 1, side.col = NULL, side.srt = 0, side.cex = 1, col.heatmap = heat(), zlim = "0 centered", zlim.trim = 0.02, norm = c("rows", "columns", "none"), norm.robust = FALSE, customLayout = FALSE, getLayout = FALSE, font = c(1, 3), xaxt = "s", yaxt = "s")

Arguments

expr A numeric matrix, holding features (genes) in columns and observations (samples) in rows. Column and row order will not be altered.

side An annotation data.frame for expr, or NULL. Must contain at least a row for each expr row, and one or many annotation column. Merging is performed on row names, so rows must be named following the same conventions as expr. Hexadecimal color definitions will be used "as is", other values will be attributed colors according to side.col.

cex.col Single numeric value, character expansion factor for column names. NA will compute a value from expr size, similarly to heatmap.

cex.row Single numeric value, character expansion factor for row names. NA will compute a value from expr size, similarly to heatmap.

mai.left Single numeric value, left margin in inches (for row names). Use NA for an automatic value computed from row name lengths. See par.

mai.bottom Single numeric value, bottom margin in inches (for column names). Use NA for an automatic value computed from column name lengths. See par.

mai.right Single numeric value, right margin in inches (for higher level functions). See par.

mai.top Single numeric value, top margin in inches. See par.

side.height Single numeric value, scaling factor for annotation track.

side.col A function returning as many colors as requested by its sole argument, defining the colors to be used for side legend. Default uses a custom palette for few values, and a derivative of rainbow if more than 8 colors are needed.

side.srt Single numeric value, determining the string rotation angle when writing character side columns (default is 0, horizontal, 90 is suggested for vertical text on busy heat maps).

side.cex Single numeric value, the character expansion factor to use for character side columns.

col.heatmap Character vector of colors, to be used for the cells of the heat map.

zlim Numeric vector of length two, defining minimal and maximal expr values that will be mapped to colors in col.heatmap. Values outside of this range will be rounded to the nearest boundary. Two special values are also allowed: "0 centered" to get a symmetrical range around 0 (with the default palette, it enforces 0 as the center color), and "range" to get expr range after normalization.

zlim.trim Single numeric value between 0 and 1, defining the proportion of extreme values (equally split on both sides) to remove before computing "0 centered" or "range" zlim.
heat.map

```
norm
norm.robust
customLayout
getLayout
font
xaxt
yaxt

Value

Invisibly returns a named list:

- `zlim`: Final value of the `zlim` argument.
- `col.heatmap`: Final value of the `col.heatmap` argument.
- `legend`: If `side` is used, a named character vector of colors used for annotation.
- `cex.col`: Final value of the `cex.col` argument.
- `cex.row`: Final value of the `cex.row` argument.
- `mai.left`: Final value of the `mai.left` argument.
- `mai.bottom`: Final value of the `mai.bottom` argument.

Author(s)

Sylvain Mareschal

See Also

clusterize, heatmap

Examples

```r
# Data with features in columns
data(rosenwald)
group <- rosenwald.cli$group
eexpr <- t(rosenwald.expr)[,1:100]

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[is.na(x)] <- round(mean(x, na.rm=TRUE), 3); x }
eexpr <- apply(eexpr, 2, f)
```
### Description

This function plots a color scale using a custom color palette, to legend `heat.map` derivated functions.

### Usage

```r
heat.scale(zlim, col.heatmap, at = -10:10, labels = NULL, horiz = TRUE, robust = FALSE, customMar = FALSE, title=NA)
```

### Arguments

- `zlim` Numeric vector of length 2, minimum and maximum of values in the palette. Should correspond to `zlim` in `heat.map`, consider to use `heat.map` invisible return to get special values.
- `col.heatmap` Character vector of colors used in the heat map. Should correspond to `col.heatmap` in `heat.map`, consider to use `heat.map` invisible return to get special values.
- `at` Numeric vector, values shown in the axis.
- `labels` Character vector as long as `at`, defining the values to show at `at`.
- `horiz` Single logical value, whether to plot an horizontal or a vertical scale.
- `robust` Single logical value, whether to legend `median` and `mad` or `mean` and `sd`. Should correspond to `heat.map norm.robust` value.
- `customMar` Single logical value, whether to skip the call to `par` to set `mar` or not.
- `title` Single logical value, whether to skip the call to `par` to set `mar` or not.

### Author(s)

Sylvain Mareschal

### See Also

`heat.map`, `clusterize`, `predict.LPS`
Description

This function trains a Linear Predictor Score model, given pre-computed coefficients. It uses data with known classes to fit the model.

It has numerous way to be called, and all the arguments are not mandatory. See the 'Examples' section.

Usage

LPS(data, coeff, response, k, threshold, formula, method = "fdr", ...)

Arguments

data: Continuous data used to retrieve classes, as a data.frame or matrix, with samples in rows and features (genes) in columns. Rows and columns should be named. Some precautions must be taken concerning data normalization, see the corresponding section below.

coeff: Pre-computed coefficients for the model, as returned by LPS.coeff (see there for format details).

response: Already known classes for the samples provided in data, preferably as a two-level factor. Can be missing if a formula with a response element is provided, but this argument precedes.

k: Single integer value, amount of features to include in the model, in decreasing order of coefficient. Can be missing if threshold or formula are provided, but this argument precedes other both of them.

threshold: Single numeric value, p-value threshold to apply for feature selection. Can be missing if k or formula are provided, but k precedes on it and it precedes on formula.

formula: A formula object, describing the model to fit (several templates are handled, see 'Examples'). The formula response element (before the "~" sign) can replace the response argument if it is not provided. The variables (after the "~" sign) can be a single integer (standing for the k argument), a single numeric (standing for the threshold argument) or a sum of feature names to use directly. "." is also handled in the usual way (all data columns), and "1" is a more efficient way to refer to all numeric columns of data.

method: Single character value, to be passed to p.adjust when threshold is provided.

...: Further arguments are passed to model.frame if response is missing (thus defined via formula). subset and na.action may be particularly useful for cross-validation schemes, see model.frame.default for details. subset is always handled but masked in "..." for compatibility reasons.
Value

An object of (S3) class "LPS":

- `coeff`: Named numeric vector, the coefficients used in the model.
- `classes`: Character vector, the labels of the two groups to be predicted.
- `scores`: List of two numeric vectors, training dataset scores sorted by group.
- `means`: Numeric vector, score means of each group in the training dataset.
- `sds`: Numeric vector, score standard deviations of each group in the training dataset.
- `ovl`: Numeric value, overlapping coefficient as returned by `OVL`.
- `k`: Integer value, amount of features selected in the model (if relevant).
- `p.threshold`: Numeric value, threshold used for feature selection (if relevant).
- `p.method`: Character value, p-value correction used for feature selection (if relevant).

Normalization

As expression values are directly used in the score, gene centering and scaling are strongly recommended. For Affymetrix raw expression values (strictly positive, linear and absolute), Wright et al. suggests a multiplicative centering on a median of 1000 followed by a log2 transformation. For log-ratio, gene centering and scaling should not be necessary, as they are naturally 0-centered.

Time efficiency

Using a numeric matrix as `data` and a factor as `response` is the fastest way to compute coefficients, if time consumption matters (as in cross-validation schemes). `formula` is there only for consistency with R modeling functions, and to provide `response`, `k` or `threshold` in a single way.

Author(s)

Sylvain Mareschal

References


See Also

LPS.coeff
Examples

# Data with features in columns
data(rosenwald)
group <- rosenwald.cli$group
expr <- t(rosenwald.expr)

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[is.na(x)] <- round(mean(x, na.rm=TRUE), 3); x }
expr <- apply(expr, 2, f)

# Coefficients
coeff <- LPS.coeff(data=expr, response=group)

# 10 best features (straightforward)
m <- LPS(data=expr, coeff=coeff, response=group, k=10)

# 10 best features (formula)
### 'k' MUST be an integer, or will be understood as a 'threshold'
### Numbers are "numeric", enforce integer with "L" or "as.integer"
m <- LPS(data=as.data.frame(expr), coeff=coeff, formula=group~10L)
k <- as.integer(10)
m <- LPS(data=as.data.frame(expr), coeff=coeff, formula=group~k)

# FDR threshold
thr <- 0.01
m <- LPS(data=expr, coeff=coeff, response=group, threshold=thr)
m <- LPS(data=as.data.frame(expr), coeff=coeff, formula=group~0.01)
m <- LPS(data=as.data.frame(expr), coeff=coeff, formula=group~thr)

# Custom model
m <- LPS(data=expr, coeff=coeff[c("27481","17013")], response=group, k=2)
m <- LPS(data=as.data.frame(expr), coeff=coeff, formula=group~"27481+17013")
### Notice backticks in formula for syntactically invalid names

# Complete model
m <- LPS(data=expr, coeff=coeff, response=group, k=ncol(expr))
m <- LPS(data=expr, coeff=coeff, response=group, threshold=1)
### m <- LPS(data=as.data.frame(expr), coeff=coeff, formula=group~.)
### The last is correct but (really) slow on large datasets

---

**LPS.coeff**  
Linear Predictor Score coefficient computation

**Description**

As Linear Predictor Score coefficients are genuinely t statistics, this function provides a faster implementation for large datasets than using `t.test`. 
Usage

LPS.coeff(data, response, formula = ~1, type = c("t", "limma"),
  p.value = TRUE, log = FALSE, weighted = FALSE, ...)

Arguments

data
Continuous data used to retrieve classes, as a data.frame or matrix, with samples in rows and features (genes) in columns. Rows and columns should be named. NA values are silently ignored. Some precautions must be taken concerning data normalization, see the corresponding section in LPS manual page.

response
Already known classes for the samples provided in data, preferably as a two-level factor. Can be missing if a formula with a response element is provided, but this argument precedes.

formula
A formula object, describing the features to consider in data. The formula response element (before the "~" sign) can replace the response argument if it is not provided. The features can be enumerated in the variable section of the formula (after the "~" sign). "." is also handled in the usual way (all data columns), and "1" is a more efficient way to refer to all numeric columns of data.

type
Single character value, "t" to compute genuine t statistics (unequal variances and unpaired samples) or "limma" to use the lmFit() and eBayes() t statistics from this microarray oriented Bioconductor package.

p.value
Single logical value, whether to compute (two-sided) p-values or not.

log
Single logical value, whether to log-transform t or not (sign will be preserved). Original description of the LPS does not include log-transformation, but it may be useful to not over-weight discriminant genes in large series. Values between -1 and 1 are transformed to 0 to avoid sign shifting, as it generally comes with non significant p-values.

weighted
Single logical value, whether to divide t (or log-transformed t) by gene mean or not. We recommend to normalize data only by samples and use weighted = TRUE to include gene centering in the model, rather than centering and scaling genes by normalizing independantly each series as Wright et al. did.

...
Further arguments are passed to model.frame if response is missing (thus defined via formula). subset and na.action may be particularly useful for cross-validation schemes, see model.frame.default for details. subset is always handled but masked in "..." for compatibility reasons.

Value

Always returns a row named numeric matrix, with a "t" column holding statistics computed. If p.value is TRUE, a second "p.value" column is added.

Note

Using a numeric matrix as data and a factor as response is the fastest way to compute coefficients, if time consumption matters (as in cross-validation schemes). formula was added only for consis-
tency with other R modeling functions, and eventually to subset features to compute coefficients for.

Author(s)
Sylvain Mareschal

References


See Also

LPS

Examples

# Data with features in columns
data(rosenwald)
group <- rosenwald$group
eqr <- t(rosenwald$expr)

# All features, all samples
k <- LPS.coeff(data=eqr, response=group)
k <- LPS.coeff(formula=group~1, data=as.data.frame(eqr))
### LPS.coeff(formula=group~., data=as.data.frame(eqr), na.action=na.pass)
### The last is correct but (really) slow on large datasets

# Feature subset, all samples
k <- LPS.coeff(data=eqr[, c("27481","17013") ], response=group)
k <- LPS.coeff(formula=group~`27481`+`17013`, data=as.data.frame(eqr))
### Notice backticks in formula for syntactically invalid names

# All features, sample subset
training <- rosenwald$set == "Training"
### training <- sample.int(nrow(eqr), 10)
### training <- which(rosenwald$set == "Training")
### training <- rownames(subset(rosenwald, set == "Training"))
k <- LPS.coeff(data=eqr, response=group, subset=training)
k <- LPS.coeff(formula=group~1, data=as.data.frame(eqr), subset=training)

# NA handling by model.frame()
k <- LPS.coeff(formula=group~1, data=as.data.frame(eqr), na.action=na.omit)
Description

Quantify the overlap between gaussian distributions of the two group scores, to assess model efficiency (best models should not overlap, to prevent from false discovery).

Usage

OVL(means, sds, cutoff=1e-4, n=1e4)

Arguments

means     Numeric vector of two values, the means of the gaussian distributions.
sds       Numeric vector of two values, the standard deviations of the gaussian distributions.
cutoff    Single numeric value, minimal quantile for integration range definition (distributions will be considered between their cutoff and 1 - cutoff quantiles only). The lesser it is, the more precise the returned value will be.
n         Single integer value, the amount of equi-distant points to use for the computation. The greater it is, the more precise the returned value will be.

Value

Returns the proportion of the overlap between the two gaussian distributions N1 and N2, i.e. min(N1,N2) / (N1 + N2).

Author(s)

Sylvain Mareschal

See Also

LPS-class, LPS, link{dnorm}

Examples

# Full overlap between identical distributions
OVL(c(0,0), c(1,1))

# Increasing shift
OVL(c(0,1), c(1,1))
OVL(c(0,2), c(1,1))
OVL(c(0,3), c(1,1))
OVL(c(0,10), c(1,1))
**plot.LPS**

*Plot method for LPS objects*

---

**Description**

This function plots the distributions of the LPS scores in each group for a fitted `LPS` object.

**Usage**

```r
## S3 method for class 'LPS'
plot(x, y, method = c("Wright", "Radmacher", "exact"), threshold = 0.9,
     values = FALSE, col.classes = c("#FFCC00", "#1144CC"), xlim, yaxt = "s",
     xlab = "LPS", ylab, las = 0, lwd = 2,...)
```

**Arguments**

- `x` An object of class "LPS", as returned by `LPS`.
- `y` Single character value defining y axis: "density" or (bayesian) "probability".
- `method` Single character value, the method to use for predictions. See `predict.LPS`.
- `threshold` Single numeric value, the confidence threshold to use for the "gray zone" (scores for which none of the two groups can be assigned with a probability greater than this threshold). See `predict.LPS`.
- `values` Single logical value, whether to plot individual scores from the training series or not.
- `col.classes` Character vector of two values giving to each class a distinct color.
- `xlim` To be passed to `plot`, see `plot.default`.
- `yaxt` To be passed to `plot`, see `par`.
- `xlab` To be passed to `plot`, see `plot.default`.
- `ylab` To be passed to `plot`, see `plot.default`.
- `las` To be passed to `plot`, see `par`.
- `lwd` To be passed to `plot`, see `par`.
- `...` Further arguments to be passed to `plot` or `par`.

**Author(s)**

Sylvain Mareschal

**See Also**

`LPS`
Examples

# Data with features in columns
data(rosenwald)
group <- rosenwald$group
eexpr <- t(rosenwald.expr)

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[is.na(x)] <- round(mean(x, na.rm=TRUE), 3); x }
eexpr <- apply(eexpr, 2, f)

# Coefficients
coeff <- LPS.coeff(data=eexpr, response=group)

# 10 best features model
m <- LPS(data=eexpr, coeff=coeff, response=group, k=10)

# Distributions of scores in each group
plot(m, "density")

# Probability for each group along the score axis
plot(m, "probability", yaxt="s")

predict.LPS  
Predict method for LPS objects

Description

This function allows predictions to be made from a fitted LPS model and a new dataset.

It can also plot a gene expression heatmap to visualize results of the prediction.

Usage

## S3 method for class 'LPS'
predict(object, newdata, type=c("class", "probability", "score"),
  method = c("Wright", "Radmacher", "exact"), threshold = 0.9, na.rm = TRUE,
  subset = NULL, col.lines = "#FFFFFF", col.classes = c("#FFCC00", "#1144CC"),
  plot = FALSE, side = NULL, cex.col = NA, cex.row = NA, mai.left = NA,
  mai.bottom = NA, mai.right = 1, mai.top = 0.1, side.height = 1, side.col = NULL,
  col.heatmap = heat(), zlim = "0 centered", norm = c("rows", "columns", "none"),
  norm.robust = FALSE, customLayout = FALSE, getLayout = FALSE, ...)

Arguments

object  
An object of class "LPS", as returned by LPS.

newdata  
Continuous data used to retrieve classes, as a data.frame or matrix, with samples in rows and features (genes) in columns. Rows and columns should be named. It can also be a named numeric vector of already computed scores.
### predict.LPS

Some precautions must be taken concerning data normalization, see the corresponding section in LPS manual page.

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>type</strong></td>
<td>Single character value, return type of the predictions to be made (&quot;class&quot;, &quot;probability&quot; or &quot;score&quot;). See 'Value' section.</td>
</tr>
<tr>
<td><strong>method</strong></td>
<td>Single character value, the method to use to make predictions (&quot;Wright&quot;, &quot;Radmacher&quot; or &quot;exact&quot;). See 'Details' section.</td>
</tr>
<tr>
<td><strong>threshold</strong></td>
<td>Threshold to use for class prediction. &quot;Wright&quot; method was designed with 0.9, &quot;Radmacher&quot; method makes no use of the threshold.</td>
</tr>
<tr>
<td><strong>na.rm</strong></td>
<td>Single logical value, if TRUE samples with one or many NA features will be scored too (concerned feature is removed for the concerned sample, which might be discutable).</td>
</tr>
<tr>
<td><strong>subset</strong></td>
<td>A subsetting vector to apply on newdata rows. See [] for handled values.</td>
</tr>
<tr>
<td><strong>col.lines</strong></td>
<td>If graph is TRUE, a single character value to be used for line drawing on the heatmap.</td>
</tr>
<tr>
<td><strong>col.classes</strong></td>
<td>If graph is TRUE, a character vector of two values giving to each class a distinct color.</td>
</tr>
<tr>
<td><strong>plot</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>side</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>cex.col</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>cex.row</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>mai.left</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>mai.bottom</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>mai.right</strong></td>
<td>To be passed to heat.map (used to plot score coefficients).</td>
</tr>
<tr>
<td><strong>mai.top</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>side.height</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>side.col</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>col.heatmap</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>zlim</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>norm</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>norm.robust</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>customLayout</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>getLayout</strong></td>
<td>To be passed to heat.map.</td>
</tr>
<tr>
<td><strong>...</strong></td>
<td>Ignored, just there to match the predict generic function.</td>
</tr>
</tbody>
</table>

**Details**

The "Compound covariate predictor" from Radmacher et al. (method = "Radmacher") simply assign each sample to the closest group (comparing the sample score to the mean scores of each group in the training dataset).

The "Linear Predictor Score" from Wright et al. (method = "Wright") modelizes scores in each training sub-group with a distinct gaussian distribution, and computes the probability for a sample to be in one of them or the other using a bayesian rule.

The "exact" mode is still under development and should not be used.
Value

For a "class" type, returns a character vector with group assignment for each new sample (possibly NA), named according to data row names.

For a "probability" type, returns a numeric matrix with two columns (probabilities to be in each group) and a row for each new sample, row named according to data row names and column named according to the group labels.

For a "score" type, returns a numeric vector with LPS score for each new sample, named according to data row names. Notice the score is the same for all methods.

If plot is TRUE, returns the list returned by heat.map, with data described above in the first unamed element.

Author(s)

Sylvain Mareschal

References


See Also

LPS

Examples

# Data with features in columns
data(rosenwald)
group <- rosenwald.cli$group
erpr <- t(rosenwald.expr)

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[is.na(x)] <- round(mean(x, na.rm=TRUE), 3); x }
erpr <- apply(erpr, 2, f)

# Coefficients
coeff <- LPS.coeff(data=erpr, response=group)

# 10 best features model
m <- LPS(data=erpr, coeff=coeff, response=group, k=10)

# Class prediction plot
predict(m, erpr, plot=TRUE)

# Wright et al. class prediction
Rosenwald dataset

Rosenwald et al. Lymphochip data

Description

This dataset contains 60 Diffuse Large B-Cell Lymphomas analysed on Lymphochip microarrays, as published by Rosenwald et al. The "Germinal Center B-cell like" and "Activated B-Cell like" subtypes, as determined by hierarchical clustering, were predicted by a LPS approach in Wright et al.

To minimize package size, values were rounded at 3 decimals and only 60 DLBCL from the 240 series were randomly selected (40 from the "Training" set, 20 from the "Validation" set), excluding "Type III" sub-types.

Usage

data(rosenwald)
**Format**

`rosenwald.expr` is a numeric matrix of expression values, with probes in rows and samples in columns. Both dimensions are named, probes by their "UNIQID" and samples by their "LYM numbers". Many NA values are present.

`rosenwald.cli` is a data.frame with a row for each sample, and 4 factor columns described below. Rows are named by samples "LYM numbers", in the same order than `rosenwald.expr`.

- `set` the "Training" or "Validation" set the sample comes from.
- `group` the DLBCL sub-type that is to be predicted ("GCB" or "ABC").
- `follow.up` follow-up of the patient, in years.
- `status` status of the patient at the end of the follow-up ("Dead" or "Alive").

**Source**

http://llmpp.nih.gov/DLBCL/

**References**


---

**surv.colors**

*Produces visual representation of survival data*

**Description**

This function generates color shades for each individual, according to their respective right-censored survival data (event occurred or not, after which follow-up time). This can prove useful to annotate heat maps with survival data.

Two color scales are used, one for right-censored individuals (lost of sight before the event occurs, yellow with default colors) and another for individual with observed events (death, relapse ... black in default colors). Shades are generated according to their impact: fast events and long follow-ups without event have strong colors, while late events and short follow-up without event are light-colored.

**Usage**

```r
surv.colors(time, event, eventColors = c("#000000", "CCCCCC"),
            censColors = c("#FFFFEE", "#FFDD00"))
```
**surv.colors**

**Arguments**

- **time**: Numeric vector, the follow-up times of each individual (see `Surv` in the survival package).
- **event**: Logical vector, whether an event (death, relapse ...) occurred at the end of each individual follow-up or not (see `Surv` in the survival package).
- **eventColors**: Character vector of length 2, the boundaries of the color scale to generate for individuals with events.
- **censColors**: Character vector of length 2, the boundaries of the color scale to generate for right-censored individuals.

**Value**

Returns a character vector, named according to `time` names.

**Author(s)**

Sylvain Mareschal

**See Also**

- `surv.scale`, `heat.map`

**Examples**

```r
# Rosenwald's dataset (hand-picked prognostic probes)
data(rosenwald)
probes <- c("30580", "16006", "32315", "16978", "26588")
expr <- t(rosenwald.expr[, probes])

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[is.na(x)] <- round(mean(x, na.rm=TRUE), 3); x }
expr <- apply(expr, 2, f)

# Survival colors
surv <- with(rosenwald.cli, surv.colors(time=follow.up, event=status=="Dead"))

# Color scale legend
with(rosenwald.cli, surv.scale(time=follow.up, event=status=="Dead"))

# Annotated clustering
side <- data.frame(OS=surv, row.names=rownames(rosenwald.cli))
clusterize(expr, side=side)
```
surv.scale  

Plots a survival color scale, for legend

Description
This function plots a color scale using a custom color palette, to legend surv.colors annotations.

Usage

```r
surv.scale(time, event, eventColors = c("#000000", "#CCCCCC"),
            censColors = c("#FFFFEE", "#FFDD00"))
```

Arguments

time  Numeric vector, the follow-up times of each individual (see Surv in the survival package).

event  Logical vector, whether an event (death, relapse ...) occured at the end of each individual follow-up or not (see Surv in the survival package).

eventColors  Character vector of length 2, the boundaries of the color scale to generate for individuals with events.

censColors  Character vector of length 2, the boundaries of the color scale to generate for right-censored individuals.

Author(s)

Sylvain Mareschal

See Also

surv.colors, survival::Surv

Examples

```r
# Rosenwald's dataset (hand-picked prognostic probes)
data(rosenwald)
probes <- c("30580", "16006", "32315", "16978", "26588")
expr <- t(rosenwald.expr[ probes ,])

# NA imputation (feature's mean to minimize impact)
f <- function(x) { x[ is.na(x) ] <- round(mean(x, na.rm=TRUE), 3); x }
expr <- apply(expr, 2, f)

# Survival colors
 surv <- with(rosenwald.cli, surv.colors(time=follow.up, event=status=="Dead"))

# Annotated clustering
 side <- data.frame(OS=surv, row.names=rownames(rosenwald.cli))
clusterize(expr, side=side)
```
# Color scale legend
with(rosenwald.cli, surv.scale(time=follow.up, event=status=="Dead"))
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