stam
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golubTrain.cv  Examplar StAM Cross-Validation Results

Description
This data set has been generated by stam.cv. It has been computed on the Golub data set on leukemia, the classification task being to separate AML from ALL patients.
Usage

```r
data(golubTrain.cv)
```

Format

This is a `stamCV` object

Details

The original data set is drawn from the `golubEsets` library. The samples 1 to 38 have been fed to `stam.cv` to generate the data set at hand.

See Also

`stamCV-class, stam.cv, golubTrain`

---

golubTrain.fit  

### Examplar StAM Model Fit

Description

This data set has been generated by `stam.fit`. It has been computed on the Golub data set on leukemia, the classification task being to separate AML from ALL patients.

Usage

```r
data(golubTrain.cv)
```

Format

This is a `stamFit` object

Details

The original data set is drawn from the `golubEsets` library. The samples 1 to 38 have been fed to `stam.cv` to generate the `golubTrain.cv`. This set in turn was given to `stam.fit` to generate the data set at hand.

See Also

`stamFit-class, stam.fit, golubTrain.cv`
image.stamPrediction

Molecular Symptoms Image on StAM Prediction

Description

Shows prediction results for each sample of the class of interest and for all nodes of a StAM model fit as a color coded image.

Usage

## S3 method for class 'stamPrediction':
image(x, aclass = NULL, main = NULL,
      xlab = NULL, show.graph = TRUE,
      max.label.length = 40,
      sample.labels = FALSE, full.names = TRUE,
      outfile = NULL, ps = FALSE, res = 72,
      pointsize = 10, width = 11,
      minspec = NULL, minsens = 0.1,
      maxsens = 1, invert = FALSE,
      col = stam.rgb.colors(r0 = 0),
      bin.thresh = NULL, ...)

Arguments

x  stamPrediction object to be illustrated
aclass  the name of the phenotype class of interest, usually the disease class. If set to NULL the first class in lexicographical order is chosen
main  the main title of the plot, generated automatically if left blank
xlab  the label of the x-axis
show.graph  (default is T) whether or not to show GO relations between the nodes
max.label.length  the maximum string length for a GO term
sample.labels  whether or not to show sample names on the x-axis
full.names  whether or not to show GO terms instead of GO IDs
outfile  name of output file if postscript or PNG graphics is to be generated. The extension of the file is chosen automatically according to the ps argument. If no outfile is specified, an interactive plot is attempted
res  resolution in points per inch
ps  if set to TRUE postscript output is generated
pointsize  the standard fontsize
width  width of image in inches (the height is computed according to the number of nodes to be shown and the pointsize)
minspec  nodes to be shown must be at least this specific. If set to NULL this is chosen such that no more than 50 nodes are shown.
minsens  nodes to be shown must be at least this sensitive
maxsens  nodes to be shown must be at most this sensitive
invert   whether or not to invert background and foreground colors
col      the color gradient to code classifier output
bin.thresh  threshold for binary color coding, if left at the default NULL, classifier outputs
            are coded on a continuous scale.
...      additional options passed to image

Details

This image illustrates classifier outputs generated during a structured analysis of microarrays (StAM). The central part of the image shows the color coded classifier outputs for each sample in the class of interest (columns) and each node from StAM’s model fit (rows). Nodes can be restricted to those of minimal sensitivity or specificity. Also nodes with particularly high sensitivity can be excluded. The color code for the classifier outputs is shown in a color bar on the right hand side if a continuous is used. If a testset is specified only these samples are used to compute sensitivity and specificity. If the sample names are not displayed on the x-axis, the test samples are marked with capital letters or just vertical bars (bars if there are too many test samples). Sensitivity and specificity is shown to the left of the figure together with the relations between the GO nodes from the Gene Ontology. GO terms are printed on the right of the image and may be used for a clickable map in HTML.

Value

A string to be used on an HTML page to provide a clickable map for the GO terms.

Author(s)

Claudio Lottaz

See Also

stam.predict

Description

Plots performance and redundancy as well as number of remaining nodes and genes for a cross validation in structured analysis of microarray data.

Usage

```r
## S3 method for class 'stamCV':
plot(x, outfile = NULL, aclass = NULL, delta = NULL,
     main = NULL, which = 0, res = 72, ps = FALSE,
     pointsize = 16, ...)
```
Arguments

- **x** the object of type `stamCV` for which the plots are to be drawn
- **outfile** name of output file if postscript or PNG graphics is to be generated. The extension of the file is chosen automatically according to the `ps` argument. If no outfile is specified, an interactive plot is attempted
- **aclass** the name of the phenotype class of interest, usually the disease class
- **which** choose the plot to be generated. 1: performance plot, 2: genes/nodes plot, 0: both plots in interactive mode, non otherwise.
- **delta** if a delta is provided, vertical lines are added accordingly
- **main** the main title of the plot, generated automatically if left blank
- **res** resolution in points per inch
- **ps** if set to TRUE postscript output is generated
- **pointsize** the standard font size
- **...** additional arguments to be passed to `plot`

Details

This function generates two plots. The first one plots the root error rate, its performance and the graph redundancy versus the shrinkage level. The second plot depicts the number of remaining nodes and remaining accessible probesets for each shrinkage candidate.

Author(s)

Claudio Lottaz

See Also

- `stam.cv`

Description

Overall alpha vs. delta plot as well as nodewise scatter plots on performance/redundancy and specificity/sensitivity.

Usage

```r
## S3 method for class 'stamFit':
plot(x, outfile = NULL, aclass = NULL, main = NULL,
     which = 0, res = 72, ps = FALSE, pointsize = 12, ...)
```
Arguments

x  the object of type stamFit for which the plots are to be drawn
outfile  name of output file if postscript or PNG graphics is to be generated. The extension of the file is chosen automatically according to the ps argument. The filename is also augmented by suffixes to distinguish the plots generated. If no outfile is specified, an interactive plot is attempted
aclass  the name of the phenotype class of interest, usually the disease class
which  choose the plot to be generated. 1:alpha vs. delta, 2:nodewise evaluation, 0:both plots in interactive mode, non otherwise.
main  the main title of the plot, generated automatically if left blank
res  resolution in points per inch
ps  if set to TRUE postscript output is generated
pointsize  the standard fontsize
...  additional arguments to be passed to plot

Details

If several values for alpha are provided to stam.fit plot.stamFit generates a compound score vs. shrinkage level plot. For each alpha one line is drawn and the best shrinkage (where the minimum is achieved) is marked. In a second pane of the same plot alpha is plotted vs. these best shrinkage levels.

Additionally nodewise scatter plots comparing performance vs. redundancy and sensitivity vs. specificity are generated.

Author(s)

Claudio Lottaz

See Also

stam.fit

plot.stamPrediction

Scatter Plot on Node Results

Description

Plots nodewise performance vs. redundancy as well as nodewise sensitivity vs. specificity side by side.

Usage

## S3 method for class 'stamPrediction':
plot(x, outfile = NULL, aclass = NULL,
     main = NULL, minspec = 0.9,
     minsens = 0.1, maxsens = 1,
     res = 72, ps = FALSE, pointsize = 12,
     ...)
Arguments

x: object of class stamPrediction to be plotted
outfile: name of output file if postscript or PNG graphics is to be generated. The extension of the file is chosen automatically according to the ps argument. If no outfile is specified, an interactive plot is attempted
aclass: the name of the phenotype class of interest, usually the disease class. If set to NULL the first class in lexicographical order is chosen.
main: the main title of the plot, generated automatically if left blank
minsens: if given the corresponding horizontal line is drawn in the sensitivity vs. specificity plot
minsens: if given the corresponding vertical line is drawn in the sensitivity vs. specificity plot
maxsens: if given the corresponding vertical line is drawn in the sensitivity vs. specificity plot
res: resolution in points per inch
ps: if set to TRUE postscript output is generated
pointsize: the standard fontsize
... additional arguments to be passed to plot

Author(s)

Claudio Lottaz

See Also

stam.predict

stamCV-class Cross Validation Information Generated by StAM

Description

Objects if this class are generated by stam.cv. It contains results of cross validated model fits generated in structured analysis of microarrays in order to choose graph shrinkage levels.

Objects from the Class

Objects can be created by calls of the form new("stamCV", exprs, classifications, beta, chip, root), but it is recommended the use the function stam.cv.

Slots

sample.labels: Object of class "character", names of samples
class.sample.classes: Object of class "character", class names for each sample
class.labels: Object of class "character", one name for each class
prior: Object of class "numeric", prior class probabilities according to prevalence
beta: Object of class "numeric", class weights, one per class
**full.pamfit**: Object of class "nsc", PAM fit on all probesets

**probs**: Object of class "array", matrix of cross validated prediction probabilities [samples x classes x nodes]

**folds**: Object of class "list", buckets used in cross validation

**results**: Object of class "data.frame", cross-validated root error rate, root performance and mean redundancy as well as remaining nodes and the accessible probesets for each delta

**node.results**: Object of class "list", performance, redundancy, sensitivity and specificity per node

**max.leafdev**: Object of class "numeric", performance of worst leaf node

**deltas**: Object of class "numeric", shrinkage candidates

See stamNet-class for slots chip, root, chippkg, GOpkg, nodes, leafs, inodes and probes.

**Extends**

Class "stamNet", directly.

**Methods**

**print** signature(x = "stamCV"): print information on cross validation

**writeHTML** signature(x = "stamCV"): generate HTML information on cross validation. However, using stam.writeHTML is recommended.

**Author(s)**

Claudio Lottaz

**See Also**

*stam.cv, stamNet-class*

---

**stam.cv**

*Cross Validated Training for StAM*

**Description**

Determine classifiers in leaf nodes and weights in inner nodes as well as best graph shrinkage by cross validated model fitting.

**Usage**

```r
stam.cv(expression.matrix, classifications,
    chip = "hgu95av2", root = "GO:0008150",
    beta = NULL, deltas = NULL, ndeltas = 10,
    results.per.node = FALSE, old.cv = NULL,
    pamimagefile = NULL, verbose = FALSE)
```
Arguments

expression.matrix
holds the expression levels. It may be of classes exprSet or ExpressionSet, or a plain numeric matrix. In the first case exprs is used to extract the expression levels. The matrix is expected to hold one column per sample and one row per probeset.

classifications
This character vector must contain one entry per sample identifying the group it belongs to. Alternatively, if expression.matrix is an exprSet or ExpressionSet, this may be the name of a phenoData variable.

chip
the name of the microarray chip. A meta data package is expected to be found holding the needed annotation, namely the links between probesets and Gene Ontology nodes.

root
the GO node used as root of the classifier graph. Only successors of this node are considered during construction of the graph.

beta
holds class weights used when judging classifier quality. The default is to set class weights to the corresponding prevalence.

deltas
numeric vector holding graph shrinkage candidates. Default is to determine ndelta candidates between 0 and the lowest shrinkage level which removes all leaf nodes.

ndeltas
number of automatically determined graph shrinkage candidates determined if deltas is not defined.

results.per.node
whether results for each node should be returned

old.cv
stamCV object used to modify when PAM fits need not to be recomputed. E.g. used when only beta is adapted.

pamimagefile
When this parameter is specified stam.cv tries to read this file and extract a stamCV object to be used as old.cv. If the file does not yet exist, PAM fits are stored there after computation.

verbose
when set to TRUE reports summary on each leaf training, otherwise shows a progress bar.

Details

stam.cv uses stam.net to generate a classifier graph for the microarray chip at hand. It then fits a PAM classifier for each leaf node only considering the probesets annotated to the node. Afterwards, in each inner node, weights are attributed to each child according to the childs classification performance. Finally, the weights are shrunken such that most of them become zero. In fact, the best shrinkage level is chosen in a cross validation setting.

Classification performance is evaluated using an inverted deviance like measure which uses weights to overstate specificity of a classifier. Weights for nodes are chosen according to this measure and shrunken by an absolute shrinkage level. For each shrinkage candidate cross validated performance results in terms of graph heterogeneity and classification performance are stored.

Value

An object of class stamCV is returned. Use the methods print and plot to extract information about the cross validation.
Author(s)
Claudio Lottaz

See Also
stamCV-class, plot.stamCV, stam.writeHTML

Examples

```r
## Not run:
# load and prepare some data
library(golubEsets)
data(Golub_Merge)
golubTrain <- Golub_Merge[,1:38]

# classify into ALL and AML
# (root is chosen to yield results reasonably fast,
# consider GO:0008150 (biological process) to obtain
# meaningful results)
golubTrain.cv <- stam.cv(golubTrain, "ALL.AML", chip="hu6800",
                      root="GO:0005576", ndeltas=10)

# get further information
print(golubTrain.cv)
plot(golubTrain.cv, delta=0.6)
## End(Not run)
```

---

**stamEval-class**

Results of a Complete StAM Analysis

Description

Objects of this class are returned by `ctam.evaluate`. Results of all steps in a structured analysis of microarray data are stored.

Objects from the Class

Objects can be created by calls of the form `new("stamEval", exprs, cv, fit, pred, testset)`, but using `stam.evaluate` is recommended.

Slots

- **chip**: Object of class "character", the name of the chip for which the classifier net is generated.
- **exprs**: Object of class "matrix", the plain matrix of expression levels [probesets x samples], rownames and colnames are expected to be defined
- **cv**: Object of class "stamCV", store cross validation results
- **fit**: Object of class "stamFit", store model fit
- **pred**: Object of class "stamPrediction", store prediction results
- **testset**: Object of class "numeric", indices of samples treated as test set. The others are used for training.
Methods

**writeHTML** signature(x = "stamEval"): generate HTML information on a complete StAM analysis, but using stam.writeHTML is recommended.

Author(s)

Claudio Lottaz

See Also

stam.evaluate, stamCV-class, stamFit-class, stamPrediction-class, stam.writeHTML

---

stam.evaluate  
*StAM Evaluation Procedure*

Description

This performs a structured analysis of microarrays (StAM) from scratch to the end. It starts with a cross-validation, performs a model fit, predicts phenotypes and writes complete HTML code with images.

Usage

```r
stam.evaluate(expression.matrix, classifications,  
report.dir = getwd(), aclass =  
names(table(classifications))[1],  
titlestem = NULL, testset =  
stam.balanced.folds(classifications, 3)[[1]],  
chip = "hgu95av2", root = "GO:0008150",  
no.output = FALSE, alpha = seq(0, 1, 0.1),  
beta = NULL, deltas = NULL, ndeltas = 30,  
minspec = NULL, minsens = 0.1, maxsens = 1,  
pamimagefile = NULL)
```

Arguments

- `expression.matrix` holds the expression levels. It may be of class exprSet or ExpressionSet, or a plain numeric matrix. In the first case exprs is used to extract the expression levels. The matrix is expected to hold one column per sample and one row per probeset.

- `classifications` This character vector must contain one entry per sample identifying the group it belongs to.

- `aclass` the name of the phenotype class of interest, usually the disease class. If set to NULL the first class in lexicographical order is chosen.

- `testset` indeces of the columns in the expression.matrix representing test samples.

- `chip` the name of the microarray chip. A meta data package of the same name is expected to be found holding the needed annotation, namely the links between probesets and Gene Ontology nodes.
the GO node used as root of the classifier graph. Only successors of this node are considered during construction of the graph/model.

root performance vs. mean redundancy weight. If set to NULL the root error rate is used exclusively to determine the best shrinkage level. If a numeric vector is provided, all alternatives are computed and the user is given an interactive choice. Values between 0 and 1 are valid, 0 meaning exclusive weight on redundancy and 1 putting exclusive weight on performance.

beta holds class weights used when judging classifier quality. The default is to set class weights to the corresponding prevalence. Several combinations of class weights may be provided for testing one after the other. To do so a matrix is expected to hold one combination of weights per row and must thus have one column per class.

deltas numeric vector holding graph shrinkage candidates. Default is to determine ndelta candidates between 0 and the lowest shrinkage level which removes all leaf nodes.

ndeltas number of automatically determined graph shrinkage candidates determined if deltas is not defined.

titlestem the first part of the title of the HTML page to be written, is complemented by some of the parameters.

report.dir the directory where the HTML pages are to be written

no.output do not generate any HTML or images

minspec nodes to be shown in molecular symptoms image must be at least this specific

minsens nodes to be shown in molecular symptoms image must be at least this sensitive

maxsens nodes to be shown in molecular symptoms image must be at most this sensitive

pamimagefile When this parameter is specified stam.cv tries to read this file and extract a stamCV object to avoid recomputing PAM fits. If the file does not yet exist, PAM fits are stored there after computation.

Details

stam.evaluate executes all steps needed in a structured analysis of a microarray study and coherently generates HTML output including plots and images. In Firstly, a 10 fold cross validation is performed with the data not identified as test set. Secondly, using an adequate graph shrinkage level, a model fit is computed. Finally, all data is used for prediction to illustrate the performance. Furthermore, this method generates a set of HTML pages. One page reports on the analysis as a whole, while additional interlinked pages, one for each node in the model fit, contain information on the fit and results of each node. On the main page plots and images illustrate and summarize the analysis.Clickable maps make the exploration of the results convenient. All files are stored together with an R data containing the returned R object in the user specified report directory.

Value

Returns an object of class stamEval containing all results generated during the above described procedure. Use the methods defined on the class corresponding the slot you want to investigate further.

Author(s)

Claudio Lottaz
See Also

stamEval-class, stam.cv, stam.fit, stam.predict, stam.writeHTML

Examples

# load and normalize some data
## Not run:
library(golubEsets)
data(Golub_Merge)

# (root is chosen to yield results reasonably fast,
# consider GO:0008150 (biological process) to obtain
# meaningful results)

# demonstrate the use of several combinations of class weights
betas <- cbind(c(0.5, 0.8, 0.9), c(0.5, 0.2, 0.1))
golubNorm.eval.explore <- stam.evaluate(Golub_Merge, "ALL.AML",
chip="hu6800", root="GO:0005576",
alpha=seq(0, 1, 0.1), beta=betas, ndelta=10)

# demonstrate the use of testsets
golubNorm.eval.predict <- stam.evaluate(Golub_Merge, "ALL.AML", testset=39:72,
chip="hu6800", root="GO:0005576", ndelta=10)
## End(Not run)

stamFit-class  
Model Fit Generated by StAM

Description

Objects of this class hold a model fit as it is generated by structured analysis of microarray data. The
function stam.fit returns such objects. They are handed on to stam.predict for predictions.

Objects from the Class

Objects can be created by calls of the form new("stamFit", cv, exprs, alpha, delta,
max.nodes, collapse.scnodes), but it is recommended to use the function stam.fit.

Slots

sample.classes: Object of class "character", class names for each sample
class.labels: Object of class "character", one name for each class
prior: Object of class "numeric", prior class probabilities according to prevalence
full.pamfit: Object of class "nsc", PAM fit on all probesets
alpha: Object of class "numeric", performance vs. redundancy weight(s)
beta: Object of class "numeric", class weights, one per class
delta: Object of class "numeric", shrinkage level given by the user
best.delta: Object of class "numeric", shrinkage level used for computing
default.delta: Object of class "numeric", default shrinkage level suggested by stam.fit
scores: Object of class "matrix", compound scores weighted using the provided alpha(s)
**alpha.tab**: Object of class "matrix", results comparing alphas  

**node.results**: Object of class "list", performance, redundancy, sensitivity and specificity per node  

**collapse.scnodes**: Object of class "logical", whether single children nodes are removed after shrinkage  

See `stamNet-class` for slots chip, root, chippkg, GOpkg, nodes, leafs, inodes and probes.

**Extends**

Class "stamNet", directly.

**Methods**

- `print` signature(x = "stamFit"): print information on the model fit.  
- `writeHTML` signature(x = "stamFit"): generate HTML information on the model fit, but using `stam.writeHTML` is recommended.

**Author(s)**

Claudio Lottaz

**See Also**

- `stam.fit`, `stamNet-class`

---

**stam.fit**  

*Fit StAM Model to Training Data*

**Description**

Using the whole expression data provided fit one StAM model according to the chosen shrinkage level.

**Usage**

`stam.fit(cv, expression.matrix, collapse.scnodes = FALSE,  
alpha = 0.5, delta = NULL, max.nodes = 100)`

**Arguments**

- `cv`  
  result `stam.cv` on the same data, must be of class `stamCV`  
- `expression.matrix`  
  holds the expression levels. It may be of class `exprSet` or `ExpressionSet`, or a plain numeric matrix. In the first case `exprs` is used to extract the expression levels. The matrix is expected to hold one column per sample and one row per probeset.  
- `collapse.scnodes`  
  if set to TRUE replace single children nodes after shrinkage
**alpha**

root performance vs. mean redundancy weight. If set to NULL the root error rate is used exclusively to determine the best shrinkage level. If a numeric vector is provided, all alternatives are computed and the user is given an interactive choice. Values between 0 and 1 are valid, 0 meaning exclusive weight on redundancy and 1 putting exclusive weight on performance.

**delta**

overrule alpha and set shrinkage level explicitly.

**max.nodes**

choose default shrinkage level such that no more than this number of nodes remain after shrinkage.

**Details**

In a first step *stam.fit* must choose a shrinkage level. In order to do so it uses results stored in the *cv*. If the user provides a shrinkage level explicitly this *delta* is used. If he specifies a single weighting factor *alpha* the corresponding weighted score is used to determine the best shrinkage level. If *alpha* is set to a vector of values, the corresponding scores are computed and a default *delta* is suggested using the median value of the *alphas*. If this shrinkage level leads to more than *max.nodes* nodes remaining the shrinkage level is increase until no more than *max.nodes* remain after shrinkage.

Using the thus determined shrinkage level a weighting of nodes is computed using the leaf node results from *cv*. Thereby, the whole dataset supplied is used.

**Value**

An object of class *stamFit* is returned. You may use the print and plot methods to further investigate the returned value.

**Author(s)**

Claudio Lottaz

**See Also**

*stam.cv*, *stamFit-class*, *plot.stamFit*, *stam.graph.plot*, *stam.writeHTML*

**Examples**

```r
## Not run:
# prepare data
library(golubEsets)
data(Golub_Merge)

# load and prepare some data
golubTrain <- Golub_Merge[,1:38]
data(golubTrain.cv)

# compute fit
golubTrain.fit <- stam.fit(golubTrain.cv, golubTrain, alpha=seq(0, 1, 0.1))

# investigate
print(golubTrain.fit)
plot(golubTrain.fit)
## End(Not run)

# show clickable web-page
```

stam.graph.plot

Draw StAM Model Fit Graph

Description

This function uses graphviz to layout a graph plot of a model fit. In addition a client side clickable map is returned to added to an HTML page.

Usage

stam.graph.plot(x, outfile = "", pointsize = 10, width = 9, height = 6)

Arguments

x
outfile
pointsize
width
height

the stamFit object holding the model to be drawn
name of output file without extension.
the standard font size
width of plot in inches
height of plot in inches

Details

This function generates a file in the dot language for graphviz. It uses the dot program to produce the layout of the graph and png as well as postscript files of this layout. Moreover, a client-side clickable map is generated which can be included in an HTML page.

Value

A character string containing HTNL code for a clickable map.

Note

This function only works on unix systems with graphviz installed.

Author(s)

Claudio Lottaz

References

stamINode-class

See Also

stam.fit

stamINode-class  Inner Nodes in Classifier Nets by StAM

Description

Objects of this class represent inner nodes of classifier nets as they are generated by structured analysis of microarray data. These nodes only contain children but never hold direct annotations of genes.

Objects from the Class

Objects can be created by calls of the form `new("stamINode", ID, GOidx, children)`.

- **ID** is the GO identifier as character string
- **GOidx** environment attributing indices to all GO identifiers
- **children** indeces of all children

Exactly one of GOidx and children must be defined, the other set to NULL (default)

Slots

- **children**: Object of class "numeric", indeces of the node’s direct children in the Gene Ontology.
- **weights**: Object of class "numeric", weights attributed to the node’s direct children.

See stamNode-class for ID, category, replacedParents and supNode.

Extends

Class "stamNode", directly.

Methods

- **print** signature(x = "stamINode"): print information on the inner node.
- **writeHTML** signature(x = "stamINode"): generate HTML information on the inner node.

Author(s)

Claudio Lottaz

See Also

stamNode-class, stamLeaf-class
stam-internal  

StAM Internal Function

Description

Function for internal use only.

Details

No details given. This function is subject to change without further notice.

Author(s)

Claudio Lottaz

stamLeaf-class  

Leaf Nodes in Classifier Nets by StAM

Description

Objects of this class represent leaf nodes in a classifier net as it is used by structured analysis of microarray data. These are the only nodes which have genes annotated.

Objects from the Class

Objects can be created by calls of the form `new("stamLeaf", ID, chip, probesidx)`.

- **ID**: The GO identifier as character string
- **chip**: The name of the chip for which the classifier net is generated.
- **probes**: A character vector holding the identifiers of the probesets in the order as they occur in the expression matrix

Slots

- **chip**: Object of class "character", the name of the chip for which the classifier net is generated.
- **probes**: Object of class "numeric", for each probeset holds its position in the expression matrices to be analyzed
- **pamfit**: Object of class "nsc", holds the results of pamr.train restricted to the genes annotated to the current leaf node.
- **delta**: Object of class "numeric", stores the best delty for the local pamfit as determined by pamr.cv.

See `stamNode-class` for `ID`, `category`, `replacedParents` and `supNode`.

Extends

Class "stamNode", directly.
Methods

**print** signature(x = "stamLeaf"): print information on the leaf node.

**writeHTML** signature(x = "stamLeaf"): generate HTML information on the leaf node.

Author(s)

Claudio Lottaz

See Also

stamNode-class, stamINode-class

---

**stamNet-class**  
Classifer Net for StAM

Description

Objects of this class describe a network of classifiers as it is used by structured analysis of microarray data.

Objects from the Class

Objects can be created by calls of the form `new("stamNet", chip, root, probes)`, or by a call to `stam.net`.

- **chip**: the name of the chip for which the classifier net is generated.
- **root**: the GO identifier of the node where the generation of the classifier net is started.
- **probes**: a character vector holding the identifiers of the probesets in the order as they occur in the expression matrix.

Slots

- **chip**: Object of class "character", the name of the chip for which the classifier net is generated.
- **root**: Object of class "character", the GO identifier of the node where the generation of the classifier net is started.
- **chippkg**: Object of class "character", information on the version of the meta data package for the chip.
- **GOpkg**: Object of class "character", information on the version of the meta data package on the Gene Ontology.
- **nodes**: Object of class "list", elements are of class stamINode or stamLeaf, one for each node in the classifier net.
- **leafs**: Object of class "numeric", indices of all leaf nodes in slot nodes.
- **inodes**: Object of class "numeric", indices of all inner nodes in slot nodes.
- **probes**: Object of class "environment", the corresponding index for each probeset in the expression matrices to be analyzed.
Methods

**print**
signature(x = "stamNet"):

print information on the classifier net

**writeHTML**
signature(x = "stamNet"):
generate HTML information on the classifier net. However, using stam.writeHTML is recommended.

Author(s)

Claudio Lottaz

See Also

stam.net, stamLeaf-class, stamINode-class

---

**stam.net**

*Generate a Classifier Graph for StAM*

Description

Generates a classifier graph for structured analysis of microarray data based on the Gene Ontology.

Usage

```r
stam.net(chip = "hgu95av2", root = "GO:0008150", probes = character(0))
```

Arguments

- **chip**
  A character string representing the microarray chip from which data has been generated. A meta data package of the same name is used to obtain the annotation data, namely for the Gene Ontology.

- **root**
  The identifier of the GO node to be used as the root of the classifier graph. Only successors of this node are considered to generate the graph.

- **probes**
  The probe names in the order as they occur in the expression matrix. Indices according to this character array are used to index probes/probesets.

Details

stam.net crawls through the Gene Ontology starting with the root node specified. It collects all successors of the root into a classifier graph according to the parent-children relations defined in the Gene Ontology. Probesets of the microarray chip at hand are attributed to GO nodes according to the Bioconductor annotation meta data package for the chip.

For any node i which has GO annotations AND successors an additional node i’ is introduced to the classifier graph. The new node is added as an additional child to i and all probesets annotated to i are moved to i’, such that only leaf nodes hold probesets. Nodes which neither themselves nor any of their successors hold probesets are discarded. stam.net also replaces inner nodes with only one child by their successor.

Value

An object of class stamNet is returned. You may use the print method to obtain detailed information about the classifier graph. You may further investigate any element of the ’nodes’ list using the print method.
stamNode-class

Author(s)
Claudio Lottaz

See Also
stamNet-class, stam.writeHTML

Examples

```r
## Not run:
# load some data
library(golubEsets)
data(Golub_Merge)
emat <- exprs(Golub_Merge)

# determine classifier graph for chip "hu6800" on
# biological processes, taking the positions of
# probesets in Golub's expression matrix into account
net <- stam.net(chip="hu6800", root="GO:0003674", probes=rownames(emat))

# have a look
print(net)
print(net@nodes[[16]])
print(net@nodes[["GO:0007638")
## End(Not run)
```

stamNode-class  

Nodes in a Classifier Net by StAM

Description

Objects of this class and its subclasses represent single nodes in a classifier net as it is used by structured analysis of microarray data.

Objects from the Classes

Objects can be created by calls of the form `new("stamNode", ID, supNode).

ID is the GO identifier as character string

supNode indicates whether the node is a supplementary node to avoid nodes which have both, genes directly annotated and children in the stamNet.

Slots

ID: Object of class "character", holds the GO identifier as character string
category: Object of class "character", represents the GO ontology the node belongs (MF: molecular function, BP: biological process, CC: cellular component).
replacedParents: Object of class "character", holds all GO identifiers of nodes which have been removed, because the had only one child.
supNode: Object of class "logical", indicates whether the nodes has been added to avoid nodes which have both directly annotated genes as well as children in the classifier graph
Methods

getGOchildren signature(x = "stamNode"): returns the identifiers of the direct children of node x in the Gene Ontology.

getGOchildren signature(x = "character"): returns the identifiers of the direct children of the node with identifier x in the Gene Ontology.

getGOparents signature(x = "stamNode"): returns the identifiers of the direct parents of node x in the Gene Ontology.

getGOparents signature(x = "character"): returns the identifiers of the direct parents of the node with identifier x in the Gene Ontology.

getGOterm signature(x = "stamNode"): returns the GO term represented by node x.

getGOterm signature(x = "character"): returns the GO term represented by the node with identifier x.

print signature(x = "stamNode"): print information on the node.

writeHTML signature(x = "stamNode"): generate HTML information on the node.

Author(s)

Claudio Lottaz

See Also

stamLeaf-class, stamINode-class

stamPrediction-class

Results of Predictions by StAM

Description

Object of this class are returned by stam.predict and contain prediction results as they are computed by structured analysis of microarray data.

Objects from the Class

Objects can be created by calls of the form new("stamPrediction", fit, expr.mat, cls, testset), but using stam.predict is recommended.

Slots

chip: Object of class "character", the name of the chip for which the classifier net is generated.

nodes: Object of class "list", elements are of class stamINode or stamLeaf, one for each remaining node in the classifier net.

class.labels: Object of class "character", one name for each class

best.delta: Object of class "numeric", shrinkage level used for computing

cls: Object of class "character", class names for each sample

probs: Object of class "array", matrix of prediction probabilities [samples x classes x nodes]
**stam.predict**

**Predictions:** Object of class "character", overall prediction for each sample

**Testset:** Object of class "numeric", indexes of samples which belong to the test set. The other samples are assumed to be the training set.

**Node results:** Object of class "list", performance, redundancy, sensitivity and specificity per node

**Methods**

- `image` signature(x = "stamPrediction"): molecular symptoms image, see `image.stamPrediction` for details
- `print` signature(x = "stamPrediction"): print information on prediction
- `writeHTML` signature(x = "stamPrediction"): generate HTML information on prediction, but using `stam.writeHTML` is recommended

**Author(s)**

Claudio Lottaz

**See Also**

`stam.predict, image.stamPrediction, stam.writeHTML`

---

**stam.predict**

*Predict Classifications of New Data*

**Description**

StAM analysis on new data using a given model fit.

**Usage**

```r
stam.predict(fit, expression.matrix, classifications = NULL, testset = NULL)
```

**Arguments**

- **fit**: stamFit object containing a trained model
- **expression.matrix**: matrix or exprSet or ExpressionSet, containing new data
- **classifications**: character vector specifying class names per sample. You may either specify one class per sample in the `expression.matrix`, or one class per training sample (all but the testset).
- **testset**: indexes of samples not used in training

**Details**

`stam.predict` uses an object returned by `stam.fit` to perform a structured analysis of the new expression data provided. Thereby, it uses all classifiers in the leaf nodes to provide classification results in these for each sample. In addition, weighted sums in inner nodes are computed to provide classification results for the whole graph.
Value

An object of type stamPrediction is returned. You may use print, plot and image functions to further investigate the results. Information on node classifiers are obtained through the print methods on elements of the nodes slot.

Author(s)

Claudio Lottaz

See Also

stamPrediction-class, plot.stamPrediction, image.stamPrediction, stam.fit, stam.writeHTML

Examples

```r
## Not run:
# load and prepare data
library(golubEsets)
data(Golub_Merge)
golubTest <- Golub_Merge[,39:72]
data(golubTrain.fit)

# compute predictions
golubTest.pred <- stam.predict(golubTrain.fit, golubTest,
pData(golubTest)[,"ALL.AML"])
golubMerge.pred <- stam.predict(golubTrain.fit, Golub_Merge,
pData(Golub_Merge)[,"ALL.AML"], testset=39:72)

# further investigate
print(golubTest.pred)
plot(golubTest.pred)
## End(Not run)
## Not run:
map <- image(golubMerge.pred, outfile="golubMerge")
cat("<HTML><BODY><MAP NAME='image_map'>", map,
"</MAP>",
"<IMG SRC='golubMerge_pred_img.png' USEMAP='#image_map'></BODY></HTML>\n",
file="pred_img.html")
browseURL(paste("file://", getwd(), "/pred_img.html", sep=""))
## End(Not run)
```

stam.rgb.colors

Generate a Color Gradient

Description

Generates a color gradient starting from one color given by the user and ending in another going through black.

Usage

```r
stam.rgb.colors(n = 12, m = 1, r0 = 1, g0 = 0, b0 = 0,
    r1 = 0, g1 = 1, b1 = 0)
```
Arguments

n  Number of shades generated in the color gradient
m  skewing factor. m=1 produces a linear gradient from start to black and from
    black to end. High m decreases the amount of black, low m's increase it.
r0 red component of the starting color
g0 green component of starting color
b0 blue component of the starting color
r1 red component of the ending color
g1 green component of the ending color
b1 blue component of the ending color

Value

A character vector of length 2*n. Each element represents the code of an RGB color.

Author(s)

Claudio Lottaz

See Also

image.stamPrediction

Examples

red.to.blue <- stam.rgb.colors(10, 1, 1, 0, 0, 0, 0, 1)
fake.data <- matrix(seq(0, 1, 0.01), nrow=10, ncol=10)
image(fake.data, col=red.to.blue)

stam.serve  StAM server launch and installation

Description

stam.serve installs StAM's server feature if it has not been installed before. Moreover it launches
the StAM server needed to use StAM output interactively through the internet.

Usage

stam.serve(tmp.path = NULL, cgi.path = NULL, cgi.url = NULL)

Arguments

tmp.path  the path to the directory where files are stored for communication between your
           WWW-server and the StAM-server.
cgi.path   the path to the directory in which the StAM-related CGI scripts are to be stored. Make
           sure that your WWW-server can execute CGI scripts from here and that access rights
           are set correctly.
cgi.url    the URL prefix needed to access the CGI scripts, i.e. the directory specified in
c           cgi.path.
Details

The stam package provides a feature to manipulate some parameters interactively using HTML forms. This feature needs a WWW browser which is able to execute CGI scripts. The HTML output used to work with interactively must be generated with the stam.write.forms option turned on.

StAM provides a set of CGI scripts which are called from the forms written into the HTML code when the above mentioned option is turned on. This scripts write task files into the directory given by tmp.path. This directory must have write permission for the WWW server. The StAM server regularly checks this directory for such tasks and executes them.

stam.serve installs the StAM server feature when it is called the first time after the installation of the stam package. Firstly, this consists of registering the three parameters of stam.serve into the package installation such that they can be reloaded when the packages is loaded into R the next time. For this purpose a dataset is written into the installation, thus you need write permissions in the corresponding directory. Secondly, the cgi.url is written into StAM’s CGI scripts and these scripts are written into the cgi.path.

When the installation of StAM’s server feature is complete, stam.serve starts checking the directory where it expects the tasks written by the CGI scripts and is thus ready for operation. After the server feature has been installed the server can be launched simply by calling stam.serve() without parameters. A second call with parameters modifies the StAM server installation accordingly.

Note

- You need write permission in the stam installation directory in order to install StAM’s server feature.
- You need write permission in the directory where the CGI scripts are to be deposited in order to install StAM’s server feature.
- You must regenerate all HTML you want to work with through the stam server after installation of the server feature.
- You must turn on the stam.write.forms option when generating HTML for use with the server feature.

Author(s)

Claudio Lottaz

See Also

stam.writeHTML

Examples

```r
## Not run:
# make sure subsequent calls to stam.writeHTML generate forms
options(stam.write.forms=TRUE)

# first call to stam.serve after installation of the stam package
stam.serve(tmp.path = "~/home/myhome/upload",
cgi.path = "~/home/myhome/cgi-bin/stam",
cgi.url = "http://www.myserver.com/cgi-bin/stam")

# subsequent calls to launch StAM server without modifying the installation
stam.serve()
## End(Not run)
```
stam.writeHTML

Write StAM Output in HTML

Description

Write HTML output for various pieces of a structured analysis of microarray data for further interactive exploration.

Usage

```
stam.writeHTML(x, title = NULL, align = "left",
               outfile = "index.html", nonodes = FALSE, ...)
```

Arguments

- **x**: the piece of the analysis for which HTML is to be generated. It may be of any of the following classes: stamNode, stamNet, stamCV, stamFit, stamPrediction or stamEval
- **title**: the title to be used on the generated main page. A default is generated according to a few important parameters of x
- **align**: alignment of the title (left, right, center)
- **outfile**: the file where to store the generated HTML code unless x is of class stamEval. In the latter case, outfile is a directory where the collection of files is stored.
- **nonodes**: whether to generate (possibly lengthy) output for all nodes
- **...**: further arguments passed to writeHTML calls.

Details

Use this function to generate HTML pages for further investigation of StAM results. The pages are interlinked and contain links to external resources such as the Gene Ontology and the Affymetrix website. Clickable maps are generated for the illustration of the model fit as well as the molecular symptoms image.

Author(s)

Claudio Lottaz

Examples

```r
## Not run:
data(golubTrain.cv)
data(golubTrain.fit)

stam.writeHTML(golubTrain.cv)
stam.writeHTML(golubTrain.fit, nonodes=TRUE)
## End(Not run)
```
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