macat

April 19, 2009

buildMACAT

Create MACAT list from objects in workspace

Description

This is a wrapper around the preprocessedLoader function. Use it, when you want to build a MACAT-list structure from objects already in your workspace.

Usage

```
buildMACAT(matrix, chip, labels = NULL, chromLocObj = NULL)
```

Arguments

matrix expression matrix with rows=genes and columns=samples; Rownames have to

match chip; Columnnames are not mandatory.

chip Identifier for used microarray

labels Classlabels for samples, has to have length=number of columns in matrix

chromLocObj Object of class chromLocation specifying the genomic position, each probe

on the array is mapped to. If not provided, it is build in the function using

annotate's function buildChromLocation.

Details

This is only a convenience wrapper around the function preprocessedLoader for the case, that you want to build a MACAT-list from objects in your workspace.

Value

A MACAT-list structure. For an example and a description of the format see data stjude in package 'stjudem'.

Author(s)

MACAT development team

See Also

preprocessedLoader, stjude in package 'stjudem'

2 compute.sliding

Examples

compute.sliding

Compute and plot smoothing of expression values or scores along the chromosome

Description

'compute.sliding' computes a smoothing of the expression data or scores along the chromosome using the specified kernel function. This function is also used within the 'evalScoring' function. 'plotSliding' creates a plot of the smoothed expression values / scores.

Usage

```
compute.sliding(data, chromosome, sample, kernel, kernelparams=NULL, step.width plotSliding(data, chromosome, sample, kernel, kernelparams=NULL, step.width=1000000, ...)
```

Arguments

data	A MACATData list holding the Expression values and gene locations
chromosome	the chromosome to be smoothed
sample	the sample (patient) whose expression values are smoothed
kernel	a kernel function (one of rbf, kNN, basePairDistance or your own)
kernelparams	a list of named parameters for the kernel (by default estimated from the data)
step.width	the smoothing is computed stepwise every step. width basepairs (default is 100000) $$
	further graphical parameters passed on to plot.default

Value

for compute.sliding: a matrix of dimension (steps x 2) with in the first column the locations in basepairs where an interpolation is computed, and in the second column the smoothed values. plotSliding does not return anything and is merely called for its side-effect producing the plot.

Author(s)

MACAT development team

```
kernelize, evalScoring
```

discreteKernelize 3

Examples

discreteKernelize Discretize and smooth expression values

Description

returns discretized kernelized expression values and saves them to a file if argument 'saveToFile' is TRUE. For details on discretization see discretize.

Usage

Arguments

data	MACATData Object
chrom	chromosome to kernelize
margin	symmetric qunatile in percent
step.width	size of the interpolation steps
kernel	kernel function one of rbf, kNN, basePairDistance or your own
kernelparams	list of named kernel parameters
saveToFile	logicval indicating whether to write a flatfile or not; default is FALSE

Details

Filename of the flatfile is: discrete_kernelized_seq_margin_<margin>_chrom_<chrom>.py where <margin> is the discretization parameter and <chrom> the name of the chromosome.

Value

discretized and kernelized expression matrix

Author(s)

The MACAT Development team

```
pydata, kernelizeAll
```

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Examples

```
#loaddatapkg("stjudem")
data(stjd)
discretizedKernelized = discreteKernelize(stjd, 13)
```

discretize

Discretize expression values

Description

'discretize' returns the discretized expression data for all chromosomes in chrom and all samples that have a label listed in label. Discretization is performed by comparing the value gene-wise (location-wise) with the symmetric upper and lower quantile given by margin (in percent margin/2 lower and upper quantile).

Usage

```
discretize(data, chrom, label, margin = 10)
discretizeChromosome(data, chrom, margin=10)
discretizeOne(data, chrom, sample, margin=10)
```

Arguments

data	MACATData object
chrom	list of chromosomes
label	list of labels
margin	symmetric quantile in percent
sample	the sample for which you want discretized expression data

Value

returns a discretized expression matrix for all genes on the chromosomes in 'chrom' and all samples that have a label in 'label'.

Author(s)

MACAT development team

```
discretizeAll
```

discretize.tscores 5

```
discretize.tscores Discretize regularized t-scores
```

Description

discretize.tscores returns a discretized version of the scores in the MACATevalScoring object. Discretization is performed by comparing the value gene-wise (location-wise) with the symmetric upper and lower quantile given by margin (in percent margin/2 lower and upper quantile). discretizeAllClasses produces a flatfile readable by PYTHON.

Usage

```
discretize.tscores(scores)
discretizeAllClasses.tscores(data, chrom, nperms=10, kernel=rbf, kernelparams=NU
```

Arguments

scores	a MACATevalScoring object obtained from evalScoring
data	a MACATData Object containing all expression values, geneLocations and labels (obtained from preprocessedLoader)
chrom	chromosome that is discretized
nperms	number of permutations for the computation of empirical p values (evalScoring)
kernel	$kernel\ function\ used\ for\ smoothing\ one\ of\ rbf,\ kNN,\ base Pair Distance\ or\ your\ own$
kernelparams	list of parameters for the kernels
step.width	size of a interpolation step in basepairs

Details

The filename for the python flat files are discrete_chrom_<chrom>_class_<label>.py where <chrom> and <label> are the names of the chromosome and class label.

Value

```
discretize.tscores

a vector of discretized tscores

discretizeAllClasses.tscores

creates python flatfiles (see details)
```

Author(s)

The MACAT development team

```
evalScoring, kernels, pythondata
```

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Examples

```
#loaddatapkg("stjudem")
data(stjd)
# simple scoring with short running time
scores = evalScoring(stjd, "T", 1, nperms=100, cross.validate=FALSE)
discrete = discretize.tscores(scores)
```

evalScoring

Score differential expression, assess significance, and smooth scores along the chromosome

Description

This function computes for all genes on one chromosome the regularized t-statistic to score differential gene expression for two given groups of samples. Additionally these scores are computed for a number of permutations to assess significance. Afterwards these scores are smoothed with a given kernel along the chromosome to give scores for chromosomal regions.

Usage

```
evalScoring(data, class, chromosome, nperms=1000, permute="labels", subset=NULL,
    newlabels=NULL, kernel=rbf, kernelparams=NULL, cross.validate=TRUE,
    paramMultipliers=2^(-4:4), ncross=10, step.width=100000,
    memory.limit=TRUE, verbose=TRUE)
```

Arguments

data	Gene expression data in the MACAT list format. See data(stjude) for an example.
class	Which of the given class labels is to be analyzed
chromosome	Chromosome to be analyzed
nperms	Number of permutations
permute	Method to do permutations. Default 'labels' does permutations of the class labels, which is the common and faster way to assess significance of differential expression. The altenative 'locations' does permutations of gene locations, is much slower and right only preliminary.
subset	If a subset of samples is to be used, give vector of column- indices of these samples in the original matrix here.
newlabels	If other labels than the ones in the MACAT-list-structure are to be used, give them as character vector/factor here. Make sure argument 'class' is one of them.
kernel	Choose kernel to smooth scores along the chromose. Available are 'kNN' for k-Nearest-Neighbors, 'rbf' for radial-basis-function (Gaussian), 'basePairDistance' for a kernel, which averages over all genes within a given range of base pairs around a position.
kernelparams	Additional parameters for the kernel as list, e.g., kernelparams=list($k=5$) for taking the 5 nearest neighbours in the kNN-kernel. If NULL some defaults are set

within the function.

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cross.validate

Logical. Should the paramter settings for the kernel function be optimized by a

cross-validation?

paramMultipliers

Numeric vector. If you do cross-validation of the kernel parameters, specify the multipliers of the given (standard) parameters to search over for the optimal one.

ncross Integer. If you do cross-validation, specify how many folds.

step.width Defines the resolution of smoothed scores on the chromosome, is in fact the

distance in base pairs between 2 positions, for which smoothed scores are to be

calculated.

memory.limit If you have a computer with lots of RAM, setting this to FALSE will increase

speed of computations.

verbose logical; should function's progress be reported to STDOUT?; default: TRUE.

Details

Please see the package vignette for more details on this function.

Value

List of class 'MACATevalScoring' with 11 components:

original.geneid

Gene IDs of the genes on the chosen chromosome, sorted according to their position on the chromosome

original.loc Location of genes on chromosome in base pairs from 5'end

original.score

Regularized t-score of genes on chromosome

original.pvalue

Empirical p-value of genes on chromosome. How often was a higher score observed than this one with random permutations? In other words, how significant

seems this score to be?

steps Positions on the chromosome in bp from 5', for which smoothed scores have

been computed.

sliding.value

Smoothed regularized t-scores at step-positions.

lower.permuted.border

Smoothed scores from permutations, lower significance border, currently 2.5%-

quantile of permutation scores.

upper.permuted.border

Smoothed scores from permutations, upper significance border, currently 97.5%-

quantile of permutation scores.

chromosome, which has been analyzed

class Class, which has been analyzed chip Identifier for used microarray

Author(s)

MACAT development team

8 evaluateParameters

See Also

```
scoring,plot.MACATevalScoring,getResults
```

Examples

```
data(stjd) # load example data
# if you have the data package 'stjudem' installed,
# you should work on the full data therein, of which
  the provided example data, is just a piece
#loaddatapkg("stjudem")
# T-lymphocyte versus B-lymphocyte on chromosome 1,
  smoothed with k-Nearest-Neighbours kernel(k=15),
  few permutations for higher speed
chrom1Tknn <- evalScoring(stjd,"T",chromosome="1",permute="labels",</pre>
nperms=100, kernel=kNN, kernelparams=list(k=15), step.width=100000)
# plotting on x11:
if (interactive())
   plot(chrom1Tknn)
# plotting on HTML:
if (interactive())
   plot(chrom1Tknn, "html")
```

evaluateParameters Evaluate Performance of Kernel Parameters by Cross-validation

Description

For a given data set, chromosome, class, and kernel function, this function helps in determining optimal settings for the kernel parameter(s). The performance of individual parameter setting is assessed by cross-validation.

Usage

```
evaluateParameters(data, class, chromosome, kernel, kernelparams = NULL,
                   paramMultipliers = 2^{(-4:4)}, subset = NULL,
                   newlabels = NULL, ncross = 10, verbose = TRUE)
```

Arguments

data Gene expression data in the MACAT list format. See data(stjude) for an exam-

Sample class to be analyzed class Chromosome to be analyzed chromosome

kernel Choose kernel to smooth scores along the chromosome. Available are 'kNN' for

k-Nearest-Neighbors, 'rbf' for radial-basis-function (Gaussian), 'basePairDistance' for a kernel, which averages over all genes within a given range of base

pairs around a position.

evaluateParameters 9

kernelparams Additional parameters for the kernel as list, e.g., kernelparams=list(k=5) for taking the 5 nearest neighbours in the kNN-kernel. If NULL some defaults are set within the function.

paramMultipliers

Numeric vector. If you do cross-validation of the kernel parameters, specify these as multipliers of the given (standard) kernel parameter, depending on your kernel choice (see page 5 of the vignette). The multiplication results are the kernel argument settings, among which you want to search for the optimal one

using cross-validation.

subset If a subset of samples is to be used, give vector of column- indices of these

samples in the original matrix here.

newlabels If other labels than the ones in the MACAT-list-structure are to be used, give

them as character vector/factor here. Make sure argument 'class' is one of them.

ncross Integer. Specify how many folds in cross-validation.

verbose Logical. Should progress be reported to STDOUT?

Value

A list of class 'MACATevP' with 4 components:

[parameterName]

List of assessed settings for the parameter [parameterName].

avgResid Average Residual Sum of Squares for the parameter settings in the same order

as the first component.

multiplier Multiplier of the original parameters in the same order as the first components.

best List of parameter settings considered optimal by cross-validation. Can be di-

rectly inserted under the argument 'kernelparams' of the 'evalScoring' function.

Author(s)

MACAT development team

See Also

```
evalScoring
```

Examples

10 getResults

getResults	Access results of 'evalScoring'	
------------	---------------------------------	--

Description

This function processes the result of the evalScoring function and returns a list of probe sets within chromosome regions deemed significant by MACAT. Additional annotation for these probe sets is provided along with their identifiers.

Usage

```
getResults (MACATevalScoringOBJ)
```

Arguments

MACATevalScoringOBJ

Object of class MACATevalScoring, usually the result from evalScoring

Details

The p-values have been computed individually for probe sets (genes), not for whole chromosome regions. Thus, regions deemed significant by sliding window approach do not have to consist only of probe sets with low p-values. These probe-set p-values are not used to determine whether a region is considered significant or not. Instead the comparison between actual and interpolated scores to actual and interpolated boundaries determines whether a region is considered significant.

This function is called within the plot function for the results of evalScoring, when HTML output is desired.

Value

A list with the following components, describing probe sets within chromosome regions deemed significant:

probeID	IDs of probe sets within these chromosome regions	
cytoband	chromosomal bands these probe sets have been annotated to	
geneSYM	gene symbols these probe sets have been annotated to	
pvalue	p-values for probe sets; see details	
locusid	EntrezGene-(formerly LocusLink) IDs of these probe sets	
genedescription		
	Description of genes the probe sets have been annotated to	
probeScore	the differential expression scores for the probe sets	
chromosome	chromosome, the analysis has been done for	
class	sample class, the analysis has been done for	

Author(s)

MACAT development team

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See Also

```
evalScoring
```

Examples

kernelize

Smooth expression values or scores

Description

'kernelize' uses a kernel to smooth the data given in geneLocations by computing a weighted sum of the values vector. The weights for each position are given in the kernelweights matrix. A kernelweights matrix can be obtained by using the kernelmatrix function.

Usage

```
getsteps(geneLocations, step.width)
kernelmatrix(steps, geneLocations, kernel, kernelparams)
kernelize(values, kernelweights)
```

Arguments

geneLocations

a list of gene locations (length n)

step.width the width of steps in basepairs

steps a list of locations where the kernelization shall be computed

kernel kernel function one of rbf, kNN or basePairDistance (or your own)
kernelparams a list of named parameters for the kernel (default is fitted to the data)
values vector of length n or matrix (m x n) of values that are to be smoothed

kernelweights

a matrix of (n x steps) where n is the length of the values vector and steps is the

number of points where you wish to interpolate

Value

getsteps a list of locations starting at min(genLocations) going to max(geneLocations)

with steps of size step.width

kernelmatrix a matrix of (n x steps) containing the kernel weights for each location in steps kernelize a vector of length steps or a matrix (m x steps) containing the smoothed values

Author(s)

MACAT Development team

See Also

```
compute.sliding, evalScoring
```

Examples

```
data(stjd)
genes = seq(100)
geneLocations = abs(stjd$geneLocation[genes])
geneExpression = stjd$expr[genes,]
step.width = 100000
steps = getsteps(geneLocations, step.width)
weights = kernelmatrix(steps, geneLocations, rbf, list(gamma=1/10^13))
kernelized = kernelize(geneExpression, weights)
plot(steps, kernelized[1,])
```

```
plot.MACATevalScoring
```

Plot function for MACATevalScoring objects.

Description

Function plots scores, 0.025 and 0.975 quantiles of the permuted scores (grey lines), and sliding average score (red line) along the chromosome. Yellow dots highlight regions, in which the smoothed absolute scores exceed the permutation-derived quantile boundaries.

Usage

Arguments

X	MACATevalScoring object.
output	plot "x11" or create a "html" -file with further information. HTML-page will open automatically.
HTMLfilename	HTML-filename, default:Results <chomosome>_<class>.html</class></chomosome>
mytitle	Title of HTML-page, default: "Results of class <class> on chromosome <chromosome>"</chromosome></class>
new.device	if FALSE: Possibility to plot several plots in one device
	further arguments passed on to generic function plot

Details

One can create a HTML-page on-the-fly if argument output='html'. The HTML-page provides informations about highlighted regions in the plot. Furthermore there are click-able Entrezgene-IDs for further analysis.

Author(s)

MACAT development team

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See Also

```
evalScoring, getResults
```

Examples

```
# see function 'evalScoring' for an example
```

preprocessedLoader Read in data and produce MACAT list

Description

This function reads expression data either from a saved R-file (.RData,.rda), or from a tab-separated text-file (.xls). For building a MACAT-list structure from objects in your workspace, you can either use this function or the convenience wrapper 'buildMACAT'.

Usage

```
preprocessedLoader(rdatafile, chip, labels = NULL, chromLocObj = NULL,
rdafile = TRUE, tabfile = FALSE, labelfile = FALSE)
```

Arguments

rdatafile	Complete name of the expression data file, or the expression matrix
chip	Identifier of the used microarray. To date only commercial Affymetrix microarrays are supported by MACAT
labels	Classlabels of the samples, vector of same length as number of columns in expression matrix; alternatively complete name of textfile with one label per line
chromLocObj	Object of class chromLocation specifying the genomic position, each probe on the array is mapped to. If not provided, it is build in the function using annotate's function buildChromLocation.
rdafile	Logical; is first argument a saved R-file?
tabfile	Logical; is first argument a tab-separated text file?
labelfile	Logical; is third argument a file with one label per line?

Value

List of class 'MACATData' with 6 components:

geneName	Identifiers of genes/probe sets in expression data
geneLocation	Location of genes on their chromosome as distance from 5'end in base pairs Negative numbers denote genes on the antisense strand.
chromosome	Chromosome of the respective gene. Components 'geneName', 'geneLocation', and 'chromosome' are in the same order.
expr	expression matrix with rows = genes and columns = samples/patients
labels	(disease) subtype of each sample, has length = number of columns of expression matrix
chip	Identifier for Microarray used for the experiments

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Note

At present, macat can only work with Affymetrix microarrays, for which an annotation package is installed on your system. Such annotation packages can either be obtained from the Bioconductor annotation packages repository or be constructed using the Bioconductor package AnnBuilder. For an example, see the common annotation package hgu95av2.

Author(s)

MACAT development team

See Also

```
buildMACAT, read. table, stjd, stjude in package 'stjudem'
```

Examples

```
## Not run:
    # assume you have your HG-U95Av2 expression values in a
    # tab-separated text file, called 'foo.txt'
    mydata <- preprocessedLoader("foo.txt","hgu95av2",rdafile=FALSE,tabfile=TRUE)
## End(Not run)</pre>
```

scoring

Compute (regularized) t-scores for gene expression data

Description

This function computes for all genes in an expression matrix the (regularized) t-scores (statistics) with the given class labels and a number of permutations of these labels. Each gene is also assigned a p-value either empirically from the permutation scores or from a t-distribution.

Usage

Arguments

data	Expression matrix with rows = genes and columns = samples
labels	Vector or factor of class labels; Scoring works only with two classes!
method	Either "SAM" to compute regularized t-scores, or "t.test" to compute Student's t-statistic
pcompute	Method to compute p-values for each genes, either "empirical" to do permutations and compute p-values from them, or "tdist" to compute p-values based on respective t-distribution
nperms	Number of permutations of the labels to be investigated, if argument 'pcompute="empirical"'
memory.limit	Logical, if you have a really good computer (>2GB RAM), setting this FALSE will increase speed of computations
verbose	Logical, if progress should be reported to STDOUT

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Details

If 'pcompute="empirical"', the statistic is computed based on the given class labels, afterwards for 'nperms' permutations of the labels. The p-value for each gene is then the proportion of permutation statistics that are higher or equal than the statistic from the real labels. For each gene the 2.5%- and the 97.5%-quantile of the permutation statistics are also returned as lower and upper 'significance threshold'.

If 'pcompute="tdist", the statistic is computed only based on the given class labels, and the p-value is computed from the t-distribution with (Number of samples - 2) degrees of freedom.

Value

A list, with four components:

```
observed (Regularized) t-scores for all genes based on the given labels

pvalues P-values for all genes, either from permutations or t-distribution

expected.lower

2.5%-quantile of permutation test-statistics, supposed to be a lower 'significance border' for the gene; or NULL if p-values were computed from t-distribution

expected.upper

97.5%-quantile of permutation test-statistics, supposed to be an upper 'signifi-
```

97.5%-quantile of permutation test-statistics, supposed to be an upper 'significance border' for the gene; or NULL if p-values were computed from t-distribution

Note

In package macat, this function is only called internally by the function evalScoring

Author(s)

MACAT development team

References

Regarding the regularized t-score please see the macat vignette.

See Also

```
evalScoring
```

Examples

```
data(stjd)
# compute gene-wise regularized t-statistics for
# T- vs. B-lymphocyte ALL:
isT <- as.numeric(stjd$labels=="T")
TvsB <- scoring(stjd$expr,isT,method="SAM",pcompute="none")
summary(TvsB$observed)</pre>
```

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stjd Subset Microarray Data from St.Jude Children Research Hospital (USA)

Description

Example for list-structure used by many functions in MACAT. It's based on the gene expression data published by Yeoh et al. (2002) The data has been preprocessed using 'vsn' on probe level and the probe values have been summed up to probe set values using the 'median polish' procedure. This is a subset of the data, containing only the data for the 5000 probe sets with the highest variance across the samples and for 10 exemplary samples, 5 from T-lymphocytic Acute Lymphocytic Leukemia (ALL) and 5 from B-lymphocytic ALL.

Usage

```
data(stjd)
```

Format

List of class 'MACATData' with 6 components:

geneName: Identifiers of genes/probe sets in expression data

geneLocation: Location of genes on their chromosome as distance from 5'end in base pairs Negative numbers denote genes on the antisense strand.

chromosome: Chromosome of the respective gene. Components 'geneName', 'geneLocation', and 'chromosome' are in the same order.

expr: expression matrix with rows = genes and columns = samples/patients

labels: (disease) subtype of each sample, has length = number of columns of expression matrix

chip: Identifier for Microarray used for the experiments (here for the Affymetrix HG-U95av2 Oligonucleotide GeneChip)

Note

For the full data package see the Bioconductor data package stjudem. If it is not already installed on your system, try source ("http:\www.bioconductor.org\biocLite.R"); biocLite("stjudem")

References

Yeoh et al. Classification, subtype discovery, and prediction of outcome in pediatric acute lymphoblastic leukemia by gene expression profiling. Cancer Cell. March 2002. 1: 133-143.

See Also

buildMACAT, st jude in package 'stjudem' for the complete expression data

Examples

```
data(stjd)
summary(stjd)
```

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