

edd

October 5, 2010

`centerScale` *center and scale a vector to zero median, unit mad*

Description

center and scale a vector to zero median, unit mad

Usage

```
centerScale(x)
```

Arguments

`x` a numeric vector

Value

a shifted and scaled version of `x` with zero median, unit mad

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

Examples

```
summary(centerScale(runif(200)))
```

edd

*new expression density diagnostics interface***Description**

this will replace edd.unsupervised; has more sensible parameters

Usage

```
edd(eset, distList=eddDistList, tx=c(sort, flatQQNormY)[[1]],
    refDist=c("multiSim", "theoretical")[1],
    method=c("knn", "nnet", "test")[1], nRowPerCand=100, ...)
```

Arguments

eset	eset – instance of Biobase ExpressionSet class
distList	distList – list comprised of eddDist objects
tx	tx – transformation of data and reference prior to classification
refDist	refDist – type of reference distribution system to use
method	method – type of classifier to use. knn is k-nearest neighbors, nnet is neural net, test is max p-value from ks.test
nRowPerCand	nRowPerCand – number of realizations for a multiSim reference system
...	... – parameters to classifiers

Details

Classifies genes according to distributional shape, by comparing observed expression distributions to a collection of references, which may be simulated or evaluated theoretically.

The distList argument is important. It enumerates the catalog of distributions for classification of gene expression vectors by distributional shape. See the HOWTO-edd vignette for information on how this list is constructed and how it can be extended.

The tx argument specifies how the data are processed for comparison to the reference catalog. This is a function on a vector returning a vector, but the input and the output need not have the same length. The default value of tx is sort, which entails that the order statistics are treated as multivariate data for classification.

The refDist argument selects the type of reference catalog. Options are 'multiSim', for which the reference consists of nRowPerCand realizations of each catalog entry, and 'theoretical', for which the reference consists of one vector of quantiles for each catalog entry.

The method argument selects the type of classifier. It would be desirable to allow this to be a function, but there is insufficient structure on classifier argument and return value structure to permit this at present; see the e1071 package for some work on handling various classifiers programmatically (e.g., tune).

Value

a character vector or factor depending on the classifier

Author(s)

Vince Carey <stvjc@channing.harvard.edu>

See Also

[ExpressionSet](#)

Examples

```
require(Biobase)
data(sample.ExpressionSet)
# should filter to genes with reasonable variation
table( edd(sample.ExpressionSet, meth="nnet", size=10, decay=.2) )
library(golubEsets)
data(Golub_Merge)
madvec <- apply(exprs(Golub_Merge),1,mad)
minvec <- apply(exprs(Golub_Merge),1,min)
keep <- (madvec > median(madvec)) & (minvec > 300)
gmfilt <- Golub_Merge[keep==TRUE,]
ALL <- gmfilt$ALL.AML=="ALL"
gall <- gmfilt[,ALL==TRUE]
gaml <- gmfilt[,ALL==FALSE]
alldists <- edd(gall, meth="nnet", size=10, decay=.2)
amldists <- edd(gaml, meth="nnet", size=10, decay=.2)
table(alldists,amldists)
amldists2 <- edd(gaml, meth="nnet", refDist="theoretical", size=10, decay=.2)
table(amldists,amldists2)
```

eddDist-class

Class "eddDist"

Description

objects from this class can be used to simulate or tabulate reference distributions for edd

Objects from the Class

Objects can be created by calls of the form `new("eddDist", ...)`. These objects include a simple stub (like "norm", which can be modified to obtain the name of a generator (prepend "r"), cdf (prepend "p"), etc.) in R.

Slots

stub: Object of class "character" stub of a distribution system in R, to which "r" is prepended to get a generator, "p" to get a cdf, "q" to get a quantile function...

parms: Object of class "numeric" named vector of parameters for a member of the family

median: Object of class "numeric" median of the distribution (sometimes has to be computed by simulation)

mad: Object of class "numeric" MAD of the distribution (sometimes has to be computed by simulation)

tag: Object of class "character" an informative character tag

latexTag: Object of class "character" an informative character tag in latex format, which can use subscripts, greek letters, etc.

plotlim: Object of class "numeric" an informative numeric indicating the upper and lower bounds of the distribution.

Methods

CDFname signature(x = "eddDist"): prepend "p" to stub(x)

genName signature(x = "eddDist"): prepend "r" to stub(x)

Mad signature(x = "eddDist"): accessor

med signature(x = "eddDist"): accessor

parms signature(x = "eddDist"): accessor

qfName signature(x = "eddDist"): prepend "q" to stub(x)

qfun signature(e = "eddDist"): construct a quantile function from the object

stub signature(x = "eddDist"): accessor

tag signature(x = "eddDist"): accessor

latexTag signature(x = "eddDist"): accessor

plotlim signature(x = "eddDist"): accessor

testVec signature(x = "numeric", eddd = "eddDist", is.centered = "logical"): apply ks.test to a given vector x against the dist specified by the eddDist components

Examples

```
library(edd)
eddDistList[[1]]
qfun(eddDistList[[1]])
```

eddObsolete

Expression Density Diagnostics

Description

Classify cohort distributions of gene expression values.

Usage

```
eddObsolete(eset,
  ref=c("multiCand", "uniCand", "test", "nnet")[1],
  k=10, l=6, nnsite=6, nniter=200)
```

Arguments

eset	instance of Biobase class ExpressionSet .
ref	one of 'multiCand', 'uniCand', 'test' or 'nnet'. see details.
k	k setting for knn – number of nearest neighbors to poll.
l	l setting for knn – minimum number of concordant assents.
nnsite	size parameter for nnet.
nniter	iter setting for nnet.

Details

Four options are available for classifying expression densities. Data on each gene are shifted and scaled to have median zero and mad 1. They are then compared to shapes of reference distributions (standard Gaussian, $\text{chisq}(1)$, $\text{lognorm}(0,1)$, $t(3)$, $.75N(0,1)+.25N(4,1)$, $.25N(0,1)+.75N(4,1)$, $\text{Beta}(2,8)$, $\text{Beta}(8,2)$, $U(0,1)$) after each of these has been transformed to have median 0 and mad 1. Classification proceeds by one of four methods, selected by setting of the 'ref' argument. Suppose there are S samples in the ExpressionSet.

multiCand – 100 samples of size S are drawn from each reference distribution and then scaled to med 0, mad 1. The knn(k,l) procedure is used to classify the genes based on proximity to representatives in this set.

uniCand – one representative of size S is created from each reference distribution, using the theoretical quantiles. knn($1,0$) is used to classify genes based on proximity to these representatives.

test – classification of each gene is based on maximum p-value of Kolmogorov-Smirnov tests vs each reference distribution. If the p-value never exceeds .1, 'doubt' is declared.

nnet – 100 samples of size S are drawn from each reference distribution and then scaled to med 0, mad 1. A neural net is fit to this dataset and the associated labels. The net is then applied to the scaled gene expression data and the predictions are used for classification.

Value

the vector of classifications, with NAs for nonclassifiable genes

Author(s)

VJ Carey

Examples

```
require(Biobase)
data(sample.ExpressionSet)
print(summary(eddObsolete(sample.ExpressionSet,k=10,l=2)))

# 6 x 20 x 50 test problem
set.seed(1234)
test <- matrix(NA,nr=120,nc=50)
test[1:20,] <- rnorm(1000)
test[21:40,] <- rt(1000,3)
test[41:60,] <- rexp(1000,4)
test[61:80,] <- rmixnorm(1000,.750,0,1,4,1)
test[81:100,] <- runif(1000)
test[101:120,] <- rlnorm(1000)
labs <- c(rep("n01",20),rep("t3",20),
rep("exp",20),rep("mix1",20),rep("u01",20),rep("ln01",20))

phenoData <- new("AnnotatedDataFrame")
pData(phenoData) <- data.frame(1:50)
varLabels(phenoData) <- list("Col1")
TT <- new("ExpressionSet", exprs=test, phenoData = phenoData)

multrun <- eddObsolete(TT, k=10, l=2)
print(table(given=labs, multiCand=multrun))
netrun <- eddObsolete(TT, ref="nnet")
print(table(given=labs, netout=netrun))
```

```
newrun <- edd(TT, meth="nnet", size=10, decay=.2)
print(table(given=labs, newout=newrun))
newrun <- edd(TT, meth="test")
print(table(given=labs, newout=newrun))
```

flatQQNorm	<i>QQ difference plot</i>
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Description

standard normal transforms to horizontal line at 0

Usage

```
flatQQNorm(y)
```

Arguments

y numeric vector

Value

list with elements *x* and *y* describing the trace of the qq difference plot

Examples

```
set.seed(1234)
plot(flatQQNorm(rnorm(40)),ylim=c(-3,3),ylab="QQNorm - line of identity ")
```

latEDtable	<i>use latex tags for dimnames of table created from edd classification</i>
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Description

use latex tags for dimnames of table created from edd classification

Usage

```
latEDtable(x, baselist=eddDistList, reorder=NULL)
```

Arguments

x *x* – a table (possibly two dimensional) of results of applying edd
baselist *baselist* – a list of eddDist objects
reorder *reorder* – a numeric vector describing how to re order the table rows/columns

Details

for use with xtable rendering. table will give results with margin names in lexicographic order; reorder can be used to get a different order.

Value

a matrix with dimnames computed from the latexTag slots of the corresponding eddDist objects

Author(s)

Vince Carey <stvjc@channing.harvard.edu>

Examples

```
require(Biobase)
data(sample.ExpressionSet)
# should filter to genes with reasonable variation
rawTab <- table( edd(sample.ExpressionSet, meth="nnet", size=10, decay=.2) )
latEDtable(rawTab)
library(xtable)
xtable(latEDtable(rawTab))
#
realTags <- sapply(eddDistList, tag)
reo <- match(realTags, names(rawTab))
xtable(latEDtable(rawTab, reorder=reo))
```

makeCandmat.raw *create and store reference distributions for edd*

Description

create and store reference distributions for edd

Usage

```
makeCandmat.raw (nPerRow=20, nRowPerCand=20, dists=
  eddDistList, centerScale=TRUE)
```

Arguments

nPerRow	size of each reference sample
nRowPerCand	number of samples per candidate
dists	list of eddDist objects specifying reference distributions
centerScale	logical indicating that simulated data should be centered and scaled

Value

matrix with rows realized from reference distributions

Author(s)

VJ Carey <stvjc@channing.harvard.edu>

Examples

```
makeCandmat.raw(5, 2, eddDistList[1:3])
```

maxKSp	<i>classify densities by maximum KS p-value</i>
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Description

compares a sample to reference distributions, identifying the closest reference by maximum KS p

Usage

```
maxKSp( x, is.centered=TRUE, dists=eddDistList, thresh=.1 )
```

Arguments

x	matrix of samples, distributions to be classified by row
is.centered	have data been brought to median 0, mad 1
dists	list of instances of class eddDist
thresh	p-value above which some test must lie, or else 'outlier' is declared

Value

a vector of classifications

Examples

```
X <- rbind( rnorm(30), runif(30), rchisq(30,1) )
tX <- t(apply(X,1,centerScale))
apply(tX,1,maxKSp)
```

plotED	<i>visualize an eddDist object</i>
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Description

plots an eddDist model; can superimpose density sketch from transformed data

Usage

```
plotED(x, y, data=NULL, is.centered=FALSE, ...)
```

Arguments

x	eddDist object
y	not used
data	optional vector of data to be superimposed in the form of a kernel density estimate
is.centered	is.centered: logical indicating that the data vector has zero median and unit mad
...	...: options passed to curve

Author(s)

Vince Carey <stvjc@channing.harvard.edu>

Examples

```
#
# show the first 8 supplied reference dists
if (interactive()){
omf <- par()$mfrow
oas <- par()$ask
on.exit(par(mfrow=omf,ask=oas))
par(mfrow=c(4,2))
par(ask=TRUE)
}
set.seed(1234)
for (i in 1:8) plotED(eddDistList[[i]])
# illustrate the superposition
if (interactive()) par(mfrow=c(1,1))
x <- rnorm(30,3,4)
plotED(N01,data=x) # relocates/scales x
y <- 12*rbeta(30,2,8)+4
plotED(B28,data=y)
```

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