# Package 'MAQCsubset'

October 16, 2018

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Title Experimental Data Package: MAQCsubset					
Version 1.18.0					
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<b>Description</b> Data Package automatically created on Sun Nov 19 15:59:29 2006.					
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<b>Depends</b> R (>= 2.10.0), affy (>= 1.23.4), Biobase (>= 2.5.5), lumi, methods					
Suggests genefilter, codelink					
License Artistic-2.0					
biocViews ExperimentData, MicroarrayData, GEO					
LazyLoad true					
git_url https://git.bioconductor.org/packages/MAQCsubset					
git_branch RELEASE_3_7					
git_last_commit d40614d					
git_last_commit_date 2018-04-30					
Date/Publication 2018-10-16					
R topics documented:					
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gehMAQCs	uhDef
gennaves	ubbei

Excerpt from GE Codelink array contributions to MAQC

## **Description**

Excerpt from GE Codelink contributions to MAQC

# Usage

```
data(gehSubRaw)
data(gehMAQCsubDef)
```

#### **Details**

 $\label{lem:gehSubRaw} gehSubRaw is a codelink: :Codelink instance based on reading the raw GEO files: "GSM123122\_GEH\_1\_A1.TXT" "GSM123123\_GEH\_1\_A2.TXT" "GSM123127\_GEH\_1\_B1.TXT" "GSM123128\_GEH\_1\_B2.TXT" "GSM123132\_GEH\_1\_C1.TXT" "GSM123133\_GEH\_1\_C2.TXT" "GSM123137\_GEH\_1\_D1.TXT" "GSM123138\_GEH\_1\_D2.TXT" \\ \label{eq:gehSubRaw} gehSuble = GSM123138 - GEH\_1\_D2.TXT" \\ \label{eq:gehSubRaw} gehSuble = GSM123128 - GEH\_1\_D2.TXT" \\ \label{eq:gehSuble} gehSuble = GSM123128 - GEH\_1\_1\_D2.TXT" \\ \label{eq:gehSuble} gehSuble = GSM123128 - GEH\_1\_1$ 

gehMAQCsubDef is an ExpressionSet instance based on default background correction and normalization of the codelink package. The original feature names include duplicates; these were made unique by make.names with unique=TRUE.

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# **Examples**

```
data(gehMAQCsubDef)
gehMAQCsubDef
```

gnfCerebHi

Gene lists for hi or low abundance in cerebellum according to Novartis GNF symatlas

# Description

Data frames with gene lists for hi or low abundance in cerebellum according to Novartis GNF symatlas

## Usage

```
data(gnfCerebHi)
data(gnfCerebLow)
```

#### **Details**

The symatlas.gnf.org database was searched using the gcrma version of human gene atlas for genes having expression in cerebellum at least 3 times (or at most 1/3 times) median expression over all organs surveyed. The resulting gene lists were intersected with genes present on GE codelink (version used in MAQCsubset) and hgu95a.

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#### Value

data.frame instances with columns providing gene name, affy probe set identifier, codelink probe identifier, illuminaHumanv1 identifier.

## Author(s)

Vince Carey <stvjc@channing.harvard.edu>

## **Examples**

```
data(gnfCerebHi)
gnfCerebHi[1:3,]
```

MAQCsubset

Experimental Data Package: MAQCsubset

# **Description**

selected data from the MAQC project (Nature Biotechnology, Sept. 2006)

## Usage

```
data(afxsubRMAES)
data(afxsubRMA)
data(afxsub)
```

## **Format**

The format is: An ExpressionSetObject with covariates:

• site: from cel

• samp: rna src/mixture code

• repl: replicate

#### Note

afxsubRMA is an exprSet (deprecated) and afxsub is an AffyBatch. afxsubRMAES is a proper ExpressionSet instance.

ilmMAQCsubR is the result of applying lumiR to the files in the vicinity of GEO GSM122901 with filename suffixes matching those of the sampleNames in the set.

## **Examples**

```
data(afxsubRMAES)
```

4 proboscis

proboscis Produce a plot similar to Figure 2 of the Shippy MAQC paper (PMID 16964226).	proboscis	Produce a plot similar to Figure 2 of the Shippy MAQC paper (PMID 16964226).	
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# Description

Produce a plot similar to Figure 2 of the Shippy MAQC paper (PMID 16964226).

## Usage

```
proboscis(es, site=1, ABp=0.001, CDp=0.01, mmrad=100)
```

## **Arguments**

es	ExpressionSet instance with MAQC assay results
site	numeric code – site to be assessed
АВр	$ABp$ – p-value threshold to declare concentration of gene in sample $\boldsymbol{A}$ to be different from ehe concentration in sample $\boldsymbol{B}$
CDp	$\mbox{\rm CDp}-\mbox{\rm p-value}$ threshold to declare concentration of gene in sample C to be different from the concentration in sample D
mmrad	numeric radius of the moving mean used to smooth the proportions differentially expressed

## **Details**

Figure 2 of the Shippy paper consists of a collection of plots of estimated probabilities of self-consistent monotone titration – briefly, samples are such that A has 100% USRNA, B has 100% Ambion brain, C has 75% USRNA+25% brain, D has 25% USRNA, 75% brain. Self-consistent monotone titration holds for gene g if microarray measures for that gene satisfy A > C > D > B or B > C > D > A. The estimated probability functions look like a creature sticking its nose over a wall, thus the name of this function.

#### Value

an instance of proboStruct, for which a plot and lines method are available.

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

#### References

PMID 16964226

## **Examples**

```
data(afxsubRMAES)
NN2 = proboscis(afxsubRMAES, site=2)
plot(NN2)
```

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proboStruct-class

Class "proboStruct"

# **Description**

structure for managing proboscis plot data

## **Objects from the Class**

Objects can be created by calls of the form new("proboStruct", ...).

#### **Slots**

```
.Data: Object of class "list" ~~ call: Object of class "call" ~~
```

## **Extends**

Class "list", from data part. Class "vector", by class "list", distance 2. Class AssayData-class, by class "list", distance 2.

#### Methods

plot

# Note

The proboscis plot shows how the probability of self-consistent monotone titration (SCMT) varies with the spiked difference in concentrations of two mRNA preparations in an MAQC dataset.

## Author(s)

V Carey <stvjc@channing.harvard.edu>

## References

For Figure 2 of Shippy et al., Using RNA sample titrations... (Nat Biotech, 24(9):1123-1131, Sep 2006)

# **Examples**

```
data(afxsubRMAES)
NN1 = proboscis(afxsubRMAES)
plot(NN1)
showClass("proboStruct")
```

# **Index**

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