xps

April 19, 2010

addData-methods Import additional CEL files into a DataTreeSet

Description

Import additional CEL files into a DataTreeSet and update ROOT data file.

Usage

```
addData(object, celdir = NULL, celfiles = "", celnames = NULL, project
= NULL, verbose = TRUE)
```

Arguments

| object | object of class DataTreeSet. |
|----------|---|
| celdir | system directory containing the CEL-files for corresponding scheme. |
| celfiles | optional vector of CEL-files to be imported. |
| celnames | optional vector of names which should replace the CEL-file names. |
| project | optional class ProjectInfo. |
| verbose | logical, if TRUE print status information. |

Details

Import additional CEL-files and update ROOT data file rootfile.

To import CEL-files from different directories, vector celfiles must contain the full path for each CEL-file and celdir must be celdir=NULL.

Value

A DataTreeSet object.

Author(s)

Christian Stratowa

See Also

import.data,root.data

Examples

AnalysisTreeSet-class

Class AnalysisTreeSet

Description

This class provides the link to the ROOT analysis file and the ROOT trees contained therein. It extends class ProcesSet.

Objects from the Class

Objects are currently created using function unifilter.

Slots

- fltrset: Object of class "FilterTreeSet" providing indirect access to the ExprTreeSet
 used and the UniFilter settings.
- scheme: Object of class "SchemeTreeSet" providing access to ROOT scheme file.
- data: Object of class "data.frame". The data.frame contains the data of the unitest stored in ROOT data trees.
- params: Object of class "list" representing relevant parameters.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT trees are stored, currently 'UniFilterSet'.
- rootfile: Object of class "character" representing the name of the ROOT file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

attachBgrd-methods

Extends

Class "ProcesSet", directly. Class "TreeSet", by class "ProcesSet", distance 2.

Methods

```
filterTreeSet signature(object = "AnalysisTreeSet"): extracts slot fltrset.
getTreeData signature(object = "AnalysisTreeSet"): exports tree data and returns
    a data.frame.
validData signature(object = "AnalysisTreeSet"): extracts data.frame data.
validFilter signature(object = "AnalysisTreeSet"): extracts data.frame data.
filtrset.
```

```
volcanoplot signature(x = "AnalysisTreeSet"): creates a volcano-plot.
```

Author(s)

Christian Stratowa

See Also

related classes FilterTreeSet.

Examples

showClass("AnalysisTreeSet")

attachBgrd-methods Attach/Remove Background Intensities

Description

Attach/remove background intensities to/from DataTreeSet.

Usage

```
attachBgrd(object, treenames = "*")
removeBgrd(object)
```

Arguments

| object | Object of class "DataTreeSet". |
|-----------|---|
| treenames | Object of class "list" representing the names of the ROOT background trees. |

Details

Whenever one of the bgcorrect methods will be applied to raw CEL intensities, the background intensities will be stored in ROOT background trees. However, the background intensities will not be saved as data.frame bgrd, thus avoiding memory problems. Function attachBgrd allows to fill slot bgrd on demand.

attachBgrd exports intensities from background trees from ROOT data file and saves as data.frame bgrd. treenames is a vector of tree names to attach; for treenames="*" all trees from slot treenames will be exported and background intensities attached as data.frame bgrd.

removeBgrd removes background intensities from DataTreeSet and replaces data.frame bgrd with an empty data.frame of dim(0,0).

Value

A DataTreeSet object.

Note

Do not use attachBgrd unless you know that your computer has sufficient RAM, especially when using exon arrays. It may be advisible to use a subset of treenames only.

Author(s)

Christian Stratowa

See Also

attachInten, removeInten

attachCall-methods Attach/Remove Detection Call Measures

Description

Attach/remove detection call and detection p-value to/from CallTreeSet.

```
Usage
```

```
attachCall(object, treenames = "*")
attachPVal(object, treenames = "*")
removeCall(object)
removePVal(object)
```

Arguments

| object | Object of class "CallTreeSet". |
|-----------|--|
| treenames | Object of class "list" representing the names of the ROOT call trees |

Details

By default detection calls will be saved in class CallTreeSet in slots data and detcall, respectively, since usually the data.frames obtained as result of e.g. mas5.call are of reasonable size. However, when computing many arrays, especially exon arrays at probeset levels, it may be better to compute detection calls with slot add.data=FALSE thus avoiding memory problems. In this case, functions attachCall and attachPVal allow to fill slots detcall and data, respectively, on demand.

attachCall exports detection calls from call trees from ROOT call file and and saves as data.frame detcall. treenames is a vector of tree names to attach; for treenames="*" all trees from slot treenames will be exported and detection calls attached as data.frame detcall.

attachPVal exports detection p-values from call trees from ROOT call file and and saves as data.frame data.treenames is a vector of tree names to attach; for treenames="*" all trees from slot treenames will be exported and detection p-values attached as data.frame data.

removeCall removes detection calls from CallTreeSet and replaces data.frame detcall with an empty data.frame of dim(0,0).

attachExpr-methods

removePVal removes detection p-values from CallTreeSet and replaces data.frame data with an empty data.frame of dim(0,0).

Value

A CallTreeSet object.

Note

Do not use attachCall and attachPVal unless you know that your computer has sufficient RAM, especially when using exon arrays. It may be advisible to use a subset of treenames only.

Author(s)

Christian Stratowa

See Also

attachExpr, removeExpr

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/'</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.</pre>
## MAS5 detection call
call.mas5 <- mas5.call(data.test3,"tmp_Test3Call0",tmpdir="",add.data=FALSE,verbose=FALSE
## attach data
call.mas5 <- attachPVal(call.mas5)</pre>
call.mas5 <- attachCall(call.mas5)</pre>
## get data.frames
pval.mas5 <- pvalData(call.mas5)</pre>
pres.mas5 <- presCall(call.mas5)</pre>
head(pval.mas5)
head(pres.mas5)
## remove data
call.mas5 <- removePVal(call.mas5)</pre>
call.mas5 <- removeCall(call.mas5)</pre>
rm(scheme.test3, data.test3)
gc()
```

attachExpr-methods Attach/Remove Expression Measures

Description

Attach/remove expression levels to/from ExprTreeSet.

Usage

```
attachExpr(object, treenames = "*")
removeExpr(object)
```

Arguments

| object | Object of class "ExprTreeSet". |
|-----------|---|
| treenames | Object of class "list" representing the names of the ROOT expression trees. |

Details

By default expression levels will be saved in class ExprTreeSet as slot data, since usually the data.frame obtained as result of e.g. rma normalization is of reasonable size. However, when normalizing many arrays, especially exon arrays at probeset levels, it may be better to compute rma with slot add.data=FALSE thus avoiding memory problems. In this case, function attachExpr allows to fill slot data on demand.

attachExpr exports expression levels from expression trees from ROOT expression file and and saves as data.frame data.treenames is a vector of tree names to attach; for treenames="*" all trees from slot treenames will be exported and expression levels attached as data.frame data.

removeExpr removes expression levels from ExprTreeSet and replaces data.frame data with an empty data.frame of dim(0,0).

Value

A ExprTreeSet object.

Note

Do not use attachExpr unless you know that your computer has sufficient RAM, especially when using exon arrays. It may be advisible to use a subset of treenames only.

Author(s)

Christian Stratowa

See Also

attachCall, removeCall

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
data.rma <- rma(data.test3,"tmp_Test3RMA0",tmpdir="",background="pmonly",normalize=TRUE,a
## attach data
data.rma <- attachExpr(data.rma)</pre>
```

attachInten-methods

```
## get data.frame
expr.rma <- validData(data.rma)
head(expr.rma)
## remove data
data.rma <- removeExpr(data.rma)
rm(scheme.test3, data.test3)
gc()</pre>
```

attachInten-methods

Attach/Remove Intensities

Description

Attach/remove raw CEL intensities to/from DataTreeSet.

Usage

```
attachInten(object, treenames = "*")
removeInten(object)
```

Arguments

| object | Object of class "DataTreeSet". |
|-----------|---|
| treenames | Object of class "list" representing the names of the ROOT data trees. |

Details

When CEL files will be imported using function import.data, the raw intensities will be stored in ROOT data trees. However, the intensities will not be saved in class DataTreeSet as slot data, thus avoiding memory problems. Function attachInten allows to fill slot data on demand.

attachInten exports intensities from data trees from ROOT data file and and saves as data.frame data. treenames is a vector of tree names to attach; for treenames="*" all trees from slot treenames will be exported and intensities attached as data.frame data.

removeInten removes intensities from DataTreeSet and replaces data.frame data with an empty data.frame of dim(0,0).

Value

A DataTreeSet object.

Note

Do not use attachInten unless you know that your computer has sufficient RAM, especially when using exon arrays. It may be advisible to use a subset of treenames only.

Author(s)

Christian Stratowa

See Also

attachBgrd, removeBgrd

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
dim(intensity(data.test3))
data.test3 <- attachInten(data.test3)
dim(intensity(data.test3))
head(intensity(data.test3))
data.test3 <- removeInten(data.test3)</pre>
```

```
dim(intensity(data.test3))
```

attachMask-methods Attach/Remove Scheme Mask

Description

Attach/remove scheme mask to/from SchemeTreeSet or to slot scheme of DataTreeSet.

Usage

```
attachMask(object)
```

```
removeMask(object)
```

Arguments

object Object of class "SchemeTreeSet" or "DataTreeSet".

Details

attachMask exports mask from scheme tree from ROOT scheme file and and saves mask as data.frame mask of slot scheme.

removeMask removes mask from SchemeTreeSet or from slot scheme of DataTreeSet and replaces data.frame mask with an empty data.frame of dim(0,0).

Value

A DataTreeSet object or SchemeTreeSet.

Note

Do not use attachMask unless you know that your computer has sufficient RAM, especially for exon array schemes.

Author(s)

Christian Stratowa

bgcorrect

See Also

import.expr.scheme, import.exon.scheme

Examples

```
## load existing ROOT scheme file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
dim(chipMask(scheme.test3))
scheme.test3 <- attachMask(scheme.test3)
dim(chipMask(scheme.test3))
head(chipMask(scheme.test3))
scheme.test3 <- removeMask(scheme.test3)</pre>
```

```
dim(chipMask(scheme.test3))
```

bgcorrect

Background Correction

Description

Background corrects probe intensities in an object of class DataTreeSet.

Usage

```
bgcorrect(xps.data, filename = character(0), filedir = getwd(), tmpdir = "", upd
bgcorrect.gc(xps.data, filename = character(0), filedir = getwd(), tmpdir = "",
bgcorrect.mas4(xps.data, filename = character(0), filedir = getwd(), tmpdir = ""
bgcorrect.mas5(xps.data, filename = character(0), filedir = getwd(), tmpdir = ""
bgcorrect.rma(xps.data, filename = character(0), filedir = getwd(), tmpdir = "",
xpsBgCorrect(object, ...)
```

Arguments

| xps.data | object of class DataTreeSet. |
|-----------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| update | logical. If TRUE the existing ROOT data file filename will be updated. |
| select | type of probes to select for background correction. |
| method | background method to use. |
| option | type of background correction to use. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |

bgcorrect

| params | vector of parameters for background method. |
|---------|---|
| verbose | logical, if TRUE print status information. |
| object | object of class DataSet. |
| | the arguments described above. |

Details

Background corrects probe intensities in an object of class DataTreeSet.

xpsBgCorrect is the DataSet method called by function bgcorrect, containing the same parameters.

Value

An DataTreeSet

Author(s)

Christian Stratowa

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.</pre>
## MAS4 sector background
data.bg.mas4 <- bgcorrect.mas4(data.test3,"tmp_Test3MAS4Bgrd",filedir=getwd(),tmpdir="",</pre>
## need to attach background intensities
data.bg.mas4 <- attachBgrd(data.bg.mas4)</pre>
## get data.frame
bg.mas4 <- validBgrd(data.bg.mas4)</pre>
head(bg.mas4)
## plot images
if (interactive()) {
image.dev(data.bg.mas4,bg=TRUE,col=rainbow(32))
image(matrix(bg.mas4[,1], ncol=ncols(schemeSet(data.bg.mas4)), nrow=nrows(schemeSet(data.
}
## Not run:
## examples using Affymetrix human tissue dataset (see also xps/examples/script4exon.R)
## example - exon array, e.g. HuEx-1_0-st-v2:
scmdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Schemes"</pre>
datdir <- "/Volumes/GigaDrive/CRAN/Workspaces/ROOTData"</pre>
scheme.exon <- root.scheme(paste(scmdir,"Scheme_HuEx10stv2r2_na25.root",sep="/"))</pre>
data.exon
           <- root.data(scheme.exon, paste(datdir,"HuTissuesExon_cel.root", sep="/"))
```

compute rma background

boxplot.dev

boxplot.dev Box Plots for Device

Description

Produce box-and-whisker plot(s) of the samples for the selected device.

Usage

boxplot.dev(x, which = "", size = 0, transfo = log2, range = 0, names = "namepar

Arguments

| Х | object of class DataTreeSet or ExprTreeSet. |
|---------|---|
| which | type of probes to be used, for details see validData. |
| size | length of sequence to be generated as subset. |
| transfo | a valid function to transform the data, usually $log2$, or 0. |
| range | determines how far the plot whiskers extend out from the box. |
| names | optional vector of sample names. |
| mar | plot margin. |
| las | style of axis labels. |
| dev | graphics device to plot to, i.e. one of "screen", "jpeg", "png", "pdf" or "ps". |
| outfile | the name of the output file. |
| W | the width of the device in pixels. |
| h | the height of the device in pixels. |
| | optional arguments to be passed to boxplot. |

Details

Produces a boxplot for slot data for an object of class DataTreeSet or ExprTreeSet for the selected graphics device.

Note

For a DataTreeSet object, data must first be attached using method attachInten.

Author(s)

Christian Stratowa

See Also

boxplot

boxplot-methods Box Plots

Description

Produce box-and-whisker plot(s) of the samples.

Usage

```
boxplot(x, which = "", size = 0, transfo = log2, range = 0, names
= "namepart", ...)
```

Arguments

| х | object of class DataTreeSet or ExprTreeSet. |
|---------|---|
| which | type of probes to be used, for details see validData. |
| size | length of sequence to be generated as subset. |
| transfo | a valid function to transform the data, usually "log2", or "0". |
| range | determines how far the plot whiskers extend out from the box. |
| names | optional vector of sample names. |
| | optional arguments to be passed to boxplot. |

Details

Creates a boxplot for slot data for an object of class DataTreeSet or ExprTreeSet.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as boxplot.

Note

For a DataTreeSet object, data must first be attached using method attachInten.

Author(s)

Christian Stratowa

See Also

boxplot.dev, boxplot

callFilter-methods

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
## need to attach scheme mask and probe intensities
data.test3 <- attachMask(data.test3)
data.test3 <- attachInten(data.test3)
if (interactive()) {
    boxplot(data.test3)
    }
## optionally remove mask and data to free memory
data.test3 <- removeInten(data.test3)
attactest3 <- removeMask(data.test3)
rm(scheme.test3, data.test3)
</pre>
```

callFilter-methods *Detection Call Filter*

Description

Detection Call Filter.

The cutoff value defines the upper threshold for allowed detection call p-values. If e.g. the number of samples exceeding this cutoff value is greater than samples then the corresponding expression dataframe row is flagged, i.e. flag = 0.

The Detection Call Filter flags all rows with: flag = (sum(call[i] >= cutoff) >= samples)

Usage

callFilter(object)
callFilter(object, value)<-</pre>

Arguments

| object | object of class PreFilter or | rUniFilte | r. |
|--------|------------------------------|-----------|-------------|
| value | character vector c (cutoff, | samples, | condition). |

Details

The method callFilter initializes the following parameters:

| cutoff: | the cutoff value for the filter: |
|------------|--|
| | cutoff = 1.0: present/absent call is used. |
| | cutoff < 1.0: detection p-value is used as cutoff. |
| samples: | this value depends on the condition used: |
| condition: | condition="samples": number of samples (default): |
| | condition="percent": percent of samples. |

Value

An initialized PreFilter or UniFilter object.

Author(s)

Christian Stratowa

Examples

```
## initialize PreFilter
prefltr <- PreFilter()
callFilter(prefltr) <- c(0.02,80.0,"percent")
str(prefltr)
## initialize UniFilter
unifltr <- UniFilter()
callFilter(unifltr) <- c(0.02,80.0,"percent")
str(unifltr)</pre>
```

callplot-methods Barplot of Percent Present and Absent Calls.

Description

Creates a barplot of percent Present/Marginal/Absent calls.

Usage

```
callplot(x, beside = TRUE, names = "namepart", col = c("red","green","blue"),
legend = c("P","M","A"), ...)
```

Arguments

| object of class CallTreeSet. |
|---|
| logical. If FALSE, the columns of height are portrayed as stacked bars, and if TRUE the columns are portrayed as juxtaposed bars. |
| optional vector of sample names. |
| color for P/M/A bars |
| legend for the plot, defaults to P/M/A. |
| optional arguments to be passed to barplot. |
| |

Details

Creates a barplot of percent Present/Marginal/Absent calls.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as callplot.

Author(s)

Christian Stratowa

CallTreeSet-class

See Also

pmplot

CallTreeSet-class Class CallTreeSet

Description

This class provides the link to the ROOT call file and the ROOT trees contained therein. It extends class ProcesSet.

Objects from the Class

Objects are created using functions mas5.call or dabg.call, respectively.

Slots

- calltype: Object of class "character" representing the call type, i.e. 'mas5' or 'dabg'.
- detcall: Object of class "data.frame". The data.frame can contain the detection calls stored in ROOT call trees.
- scheme: Object of class "SchemeTreeSet" providing access to ROOT scheme file.
- data: Object of class "data.frame". The data.frame can contain the data (i.e. p-values) stored in ROOT call trees.
- params: Object of class "list" representing relevant parameters.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT call trees are stored, usually 'CallTreeSet'.
- settype: Object of class "character" describing the type of treeset stored in setname, usually 'preprocess'.
- rootfile: Object of class "character" representing the name of the ROOT call file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

Extends

Class "ProcesSet", directly. Class "TreeSet", by class "ProcesSet", distance 2.

Methods

- attachCall signature(object = "CallTreeSet"): exports detection call data from ROOT
 call file and and saves as data.frame detcall.
- **attachPVal** signature(object = "CallTreeSet"): exports call p-values from ROOT call file and and saves as data.frame data.
- callplot signature(x = "CallTreeSet"): creates a barplot of percent present and absent calls.
- presCall signature (object = "CallTreeSet"): extracts the detection call data.frame.
- presCall<- signature(object = "CallTreeSet", value = "data.frame"):replaces the detection call data.frame.
- pvalData signature(object = "CallTreeSet"): extracts the detection p-value data.frame.
- pvalData<- signature(object = "CallTreeSet", value = "data.frame"):replaces the detection p-value data.frame.
- removePVal signature(object = "CallTreeSet"): replaces data.frame data with an empty data.frame of dim(0,0).

Author(s)

Christian Stratowa

See Also

related classes DataTreeSet, ExprTreeSet.

Examples

showClass("CallTreeSet")

cvFilter-methods Coefficient of Variation Filter

Description

This method initializes the Coefficient of Variation Filter.

The coefficient of variation is the standard deviation divided by the absolute value of the mean. The CV Filter flags all rows with: $flag = (cv \ge cutoff)$

Usage

```
cvFilter(object)
cvFilter(object, value)<-</pre>
```

dabg.call

Arguments

| object | object of class PreFilter. | | |
|--------|--------------------------------------|-------|-----------|
| value | <pre>numeric vector c (cutoff,</pre> | trim, | epsilon). |

Details

The method cvFilter initializes the following parameters:

| the cutoff level for the filter. |
|--|
| the trim value for trimmed mean (default is trim=0). |
| value to replace mean (default is epsilon=0.01): |
| epsilon > 0: replace mean=0 with epsilon. |
| epsilon = 0: always set mean=1. |
| |

Note, that for epsilon = 0 the filter flags all rows with: stdev >= cutoff

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
cvFilter(prefltr) <- c(0.3,0.0,0.01)
str(prefltr)</pre>
```

dabg.call Detection Above Background Call

Description

Computes the Detection Above Background Call first implemented for the Exon arrays.

Usage

Arguments

| xps.data | object of class DataTreeSet. |
|------------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| alpha1 | a significance threshold in (0,alpha2). |
| alpha2 | a significance threshold in (alpha1,0.5). |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE call data will be added to slots data and detcall. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | the arguments described above. |

Details

This function generates a detection p-value based on comparing the perfect match probe intensity to the intensity distribution provided by background probes sharing the same GC-content as the PM probe under consideration. For exon/genome arrays special 'antigenomic' background probes of defined GC-content are used, while for expression arrays the Mismatch probes will be grouped by their GC-content.

For exon/genome arrays it is necessary to supply option and exonlevel.

Following options are valid for exon arrays only:

| transcript: | expression levels are computed for transcript clusters, i.e. probe sets containing the same 'transcript_clu |
|-------------|--|
| exon: | expression levels are computed for exon clusters, i.e. probe sets containing the same 'exon_id', where e |
| probeset: | expression levels are computed for individual probe sets, i.e. for each 'probeset_id'. |

Following exonlevel annotations are valid for exon arrays:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |
| metafull: | full meta-probesets. |
| ambiguous: | ambiguous probesets only. |
| affx: | standard AFFX controls. |
| all: | combination of above. |

Following exonlevel annotations are valid for whole genome arrays:

| core: | probesets with category 'unique' and 'mixed'. |
|-----------|---|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |

dabg.call

all: combination of above.

Exon levels can also be combined, with following combinations being most useful:

```
exonlevel="metacore+affx":core meta-probesets plus AFFX controlsexonlevel="core+extended":probesets with cDNA supportexonlevel="core+extended+full":supported plus predicted probesets
```

Exon level annotations are described in the Affymetrix whitepaper 'exon_probeset_trans_clust_whitepaper.pdf'.

In order to use an alternative SchemeTreeSet set the corresponding SchemeTreeSet xps.scheme.

xpsDABGCall is the DataTreeSet method called by function dabg.call, containing the same parameters.

Value

 $A\,\texttt{CallTreeSet}$

Note

Yes, it is possible to compute DABG detection call for expression arrays, but it is very slow and thus not recommended.

Author(s)

Christian Stratowa

References

Affymetrix (2005) Exon Probeset Annotations and Transcript Cluster Groupings, Affymetrix Inc., Santa Clara, CA, exon_probeset_trans_clust_whitepaper.pdf.

See Also

mas5.call

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
## DABG detection call
call.dabg <- dabg.call(data.test3,"tmp_Test3DABG",verbose=FALSE)</pre>
```

```
## get data.frames
pval.dabg <- pvalData(call.dabg)
pres.dabg <- presCall(call.dabg)
head(pval.dabg)
head(pres.dabg)</pre>
```

plot results
if (interactive()) {
callplot(call.dabg)

```
}
rm(scheme.test3, data.test3)
gc()
```

DataTreeSet-class Class DataTreeSet

Description

This class provides the link to the ROOT data file and the ROOT trees contained therein. It extends class ProcesSet.

Objects from the Class

Objects can be created using the functions import.data or root.data.

Slots

- bgtreenames: Object of class "list" representing the names of optional ROOT background trees.
- bgrd: Object of class "data.frame". The data.frame can contain background intensities stored in ROOT background trees.
- projectinfo: Object of class "ProjectInfo" containing information about the project.
- scheme: Object of class "SchemeTreeSet" providing access to ROOT scheme file.
- data: Object of class "data.frame". The data.frame can contain the data (e.g. intensities) stored in ROOT data trees.
- params: Object of class "list" representing relevant parameters.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT data trees are stored, usually 'DataTreeSet'.
- settype: Object of class "character" describing the type of treeset stored in setname, usually 'rawdata'.
- rootfile: Object of class "character" representing the name of the ROOT data file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

Extends

Class "ProcesSet", directly. Class "TreeSet", by class "ProcesSet", distance 2.

DataTreeSet-class

Methods

- addData signature(object = "DataTreeSet"): import additional CEL-files and update ROOT data file rootfile.

- attachMask signature(object = "DataTreeSet"): exports scheme tree from ROOT
 scheme file and and saves as data.frame mask of slot scheme.
- background signature(object = "DataTreeSet"): extracts slot bgrd.
- background<- signature(object = "DataTreeSet", value = "data.frame"):
 replaces slot bgrd.</pre>
- **bgtreeNames** signature(object = "DataTreeSet"): **extracts slot** bgtreenames.
- intensity signature(object = "DataTreeSet"): extracts slot data.
- intensity<- signature(object = "DataTreeSet", value = "data.frame"):replaces slot data.</pre>
- mm signature(object = "DataTreeSet"): extracts the mismatch intensities.
- ncols signature(object = "DataTreeSet"): extracts the physical number of array columns
 from slot scheme.
- nrows signature(object = "DataTreeSet"): extracts the physical number of array
 rows from slot scheme.
- pm signature(object = "DataTreeSet"): extracts the perfect match intensities.
- pmplot signature(x = "DataTreeSet"): creates a barplot of mean perfect match and mismatch intensities.
- projectInfo signature(object = "DataTreeSet"): extracts slot projectinfo.
- projectInfo<- signature(object = "DataTreeSet", value = "ProjectInfo"):
 replaces slot projectinfo.</pre>
- rawCELName signature(object = "DataTreeSet"): returns the name(s) of the imported raw CEL-files.
- **removeBgrd** signature(object = "DataTreeSet"): replaces data.frame bgrd with an empty data.frame of dim(0,0).
- **removeInten** signature(object = "DataTreeSet"): replaces data.frame data with an empty data.frame of dim(0,0).
- removeMask signature(object = "DataTreeSet"): replaces data.frame mask from slot scheme with an empty data.frame of dim(0,0).
- validBgrd signature(object = "DataTreeSet"): extracts the valid data from data.frame
 bgrd.
- xpsBgCorrect signature(object = "DataTreeSet"): applies background correction methods. See bgcorrect.
- xpsDABGCall signature(object = "DataTreeSet"): computes DABG call.

xpsINICall signature(object = "DataTreeSet"): computes I/NI call. xpsMAS4 signature(object = "DataTreeSet"): computes MAS4 expression levels. xpsMAS5 signature(object = "DataTreeSet"): computes MAS5 expression levels. xpsMAS5Call signature(object = "DataTreeSet"): computes MAS5 detection call. xpsNormalize signature(object = "DataTreeSet"): applies normalization methods. xpsPreprocess signature(object = "DataTreeSet"): applies normalization methods. xpsRMA signature(object = "DataTreeSet"): computes RMA expression levels. xpsRMA signature(object = "DataTreeSet"): computes RMA expression levels.

Author(s)

Christian Stratowa

See Also

related classes ExprTreeSet, CallTreeSet.

Examples

showClass("DataTreeSet")

dfw

Distribution Free Weighted Expression Measure

Description

This function converts a DataTreeSet into an ExprTreeSet using the Distribution Free Weighted Fold Change (DFW) method.

Usage

```
dfw(xps.data,
   filename
              = character(0),
   filedir = getwd(),
tmpdir = "",
              = "",
   tmpdir
   normalize = TRUE,
              = 3,
   m
              = 1,
   n
              = 0.01,
   С
   option = "transcript",
   exonlevel = "",
   xps.scheme = NULL,
   add.data = TRUE,
   verbose = TRUE)
```

dfw

Arguments

| xps.data | object of class DataTreeSet. |
|------------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| normalize | logical. If TRUE normalize data using quantile normalization. |
| m | positive number as exponent of the weighted range WR. |
| n | positive number as exponent of the weighted standard deviation WSD. |
| С | scaling parameter. |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |

Details

This function computes the DFW (Distribution Free Weighted Fold Change) expression measure described in Chen et al. for both expression arrays and exon arrays. For exon arrays it is necessary to supply the requested option and exonlevel.

Following options are valid for exon arrays:

transcript:expression levels are computed for transcript clusters, i.e. probe sets containing the same 'transcript_clu
expression levels are computed for exon clusters, i.e. probe sets containing the same 'exon_id', where e
expression levels are computed for individual probe sets, i.e. for each 'probeset_id'.

Following exonlevel annotations are valid for exon arrays:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |
| metafull: | full meta-probesets. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Following exonlevel annotations are valid for whole genome arrays:

| core: | probesets with category 'unique', 'similar' and 'mixed'. |
|-----------|--|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Exon levels can also be combined, with following combinations being most useful:

```
exonlevel="metacore+affy": core meta-probesets plus AFFX controls
exonlevel="core+extended": probesets with cDNA support
exonlevel="core+extended+full": supported plus predicted probesets
```

Exon level annotations are described in the Affymetrix whitepaper exon_probeset_trans_clust_whitepaper.pdf: "Exon Probeset Annotations and Transcript Cluster Groupings".

In order to use an alternative SchemeTreeSet set the corresponding SchemeSet xps.scheme.

Value

An ExprTreeSet

Note

The expression measure obtained with DFW is given in linear scale, analogously to the expression measures computed with mas5 and rma.

For the analysis of many exon arrays it may be better to define a tmpdir, since this will store only the results in the main file and not e.g. background and normalized intensities, and thus will reduce the file size of the main file. For quantile normalization memory should not be an issue, however DFW depends on RAM unless you are using a temporary file.

Author(s)

Christian Stratowa

References

Chen, Z., McGee M., Liu Q., and Scheuermann, R.H. (2007), A distribution free summarization method for Affymetrix GeneChip arrays. Bioinformatics 23(3):321-327

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
data.dfw <- dfw(data.test3,"tmp_Test3DFW",verbose=FALSE)</pre>
```

```
## get data.frame
expr.dfw <- validData(data.dfw)
head(expr.dfw)</pre>
```

diffFilter-methods *Difference Filter*

Description

This method initializes the Difference Filter.

The difference is the maximum value minus minimum value for each row of the expression dataframe divided by the mean value of each row.

The Difference Filter flags all rows with: flag = ((max - min)/mean >= cutoff)

Usage

```
diffFilter(object)
diffFilter(object, value)<-</pre>
```

Arguments

| object | object of class PreFilter. | | |
|--------|----------------------------|-------|-----------|
| value | numeric vector c(cutoff, | trim, | epsilon). |

Details

The method diffFilter initializes the following parameters:

| cutoff: | the cutoff level for the filter. |
|----------|--|
| trim: | the trim value for trimmed mean (default is trim=0). |
| epsilon: | value to replace mean (default is epsilon=0.01): |
| | epsilon > 0: replace mean=0 with epsilon. |
| | epsilon = 0: always set mean=1. |

Note, that for epsilon = 0 the filter flags all rows with: (max - min) >= cutoff

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
diffFilter(prefltr) <- c(2.2,0.0,0.01)
str(prefltr)</pre>
```

existsROOTFile Test for Existing ROOT File

Description

Test if a ROOT file does already exist.

Usage

existsROOTFile(filename, tmp.rm = TRUE)

Arguments

| filename | name of ROOT file, including full path. |
|----------|--|
| tmp.rm | logical, if ${\tt TRUE}$ then exlude filenames beginning with dQuote(tmp_). |

Value

Return TRUE if file filename is an already existing ROOT file.

Note

It is possible to create temporary ROOT files called "tmp" or with filename starting with "tmp_" which can be overwritten. Thus by default temporary files will not be recognized by existsROOTFile. If you want to recognize temporary files, set tmp.rm = TRUE.

Author(s)

Christian Stratowa

See Also

isROOTFile

Examples

```
existsROOTFile(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"))
```

| exonLevel | Conversion of Parameter exonlevel to Integer |
|-----------|--|
|-----------|--|

Description

Conversion of parameter exonlevel to an integer vector.

Usage

```
exonLevel(exonlevel = "", chiptype = "GeneChip", as.sum = TRUE)
```

exonLevel

Arguments

| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
|-----------|---|
| chiptype | chip tpye, one of 'GeneChip', 'GenomeChip', 'ExonChip'. |
| as.sum | logical, if TRUE an integer vector of size three will be returned, if FALSE then the levels will be split into the basic integer representations. |

Details

Conversion of parameter exonlevel to an integer; this function is a utility function, which is usually only used internally.

Following exonlevel annotations are valid for exon arrays:

| core: | (=8192+1024) probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|---|
| metacore: | (=8192) core meta-probesets. |
| extended: | (=4096+512) probesets with other cDNA support. |
| metaextended: | (=4096) extended meta-probesets. |
| full: | (=2048+256) probesets supported by gene predictions only. |
| metafull: | (=2048) full meta-probesets. |
| ambiguous: | (=128) probesets that fall within multiple genes. |
| affx: | (=60) standard AFFX controls. |
| all: | (=16316) combination of above (including affx). |

Following exonlevel annotations are valid for whole genome arrays:

```
core:(=8192+1024) probesets with category 'unique', 'similar' and 'mixed'.metacore:(=8192) probesets with category 'unique' only.affx:(=60) standard AFFX controls.all:(=9276) combination of above (including affx).
```

Exon levels can also be combined, with following combinations being most useful:

| exonlevel="metacore+affx": | core meta-probesets plus AFFX controls |
|---------------------------------|--|
| exonlevel="core+extended": | probesets with cDNA support |
| exonlevel="core+extended+full": | supported plus predicted probesets |

Exon level annotations are described in the Affymetrix whitepaper exon_probeset_trans_clust_whitepaper.pdf: "Exon Probeset Annotations and Transcript Cluster Groupings".

Parameter exonlevel determines not only which probes are used for medianpolish, but also the probes used for background calculation and for quantile normalization. If you want to use seperate probes for background calculation, quantile normalization and medianpolish summarization, you can pass a numeric vector containing three integer values corresponding to the respective exonlevel. These integers must be the sum of the integers shown above, e.g. you can use exonlevel=c (16316, 8252, 8252), where 8252=8192+60 for "metacore+affx".

Value

an integer vector.

export.filter

Note

The following exonlevels are unsupported:

| control->bgp->genomic: | (=32768) genomic background probes. |
|----------------------------|---|
| control->bgp->antigenomic: | (=65536) antigenomic background probes. |
| normgene->intron: | (=131072) intronic controls. |
| normgene->exon: | (=262144) exronic controls. |
| rescue->FLmRNA->unmapped: | (=524288) unmapped mRNAs. |
| | |

For whole genome arrays it is possible (but not recommended) to use all probesets by using exonlevel=c(992316, 9 For exon arrays it is possible to use e.g. exonlevel=c(1032124, 1032124, 631868). However, please note that these settings are not recommended and not supported.

Author(s)

Christian Stratowa

See Also

rma, mas5

Examples

```
exonLevel("core", "GenomeChip")
exonLevel("all", "GenomeChip")
exonLevel("core+extended+full", "ExonChip")
exonLevel("core+extended+full", "ExonChip", as.sum=FALSE)
exonLevel(c(16316,8252,8252), "ExonChip")
```

export.filter Export filter data as text files

Description

Export data from classes FilterTreeSet or AnalysisTreeSet to outfile.

Usage

```
export.filter(xps.fltr, treename = "*", treetype = character(0), varlist = "*",
```

Arguments

| xps.fltr | an object of type FilterTreeSet or AnalysisTreeSet. |
|---|---|
| treename | tree name to export. |
| treetype | type of tree(s) to export, 'pfr', 'ufr' or 'stt'. |
| varlist | names of tree leaves to export. |
| outfile | name of output file. |
| sep | column separator |
| as.dataframe | if TRUE a data.frame will be returned. |
| verbose | logical, if TRUE print status information. |
| varlist outfile sep as.dataframe | names of tree leaves to export. name of output file. column separator if TRUE a data.frame will be returned. |

export.filter

Details

Export data from classes FilterTreeSet, or AnalysisTreeSet to outfile.

Parameter varlist lists the parameters to export:

```
- parameters are separated by ":", e.g. varlist="fUnitName:fFlag".
```

- for varlist="*" all valid parameters will be exported.

For class FilterTreeSet the following varlist parameters are valid:

| fUnitName: | unit name (probeset ID). |
|------------|--------------------------|
| fFlag: | mask. |

For class AnalysisTreeSet varlist can contain annotation parameters and parameters of the resulting data.

Following varlist annotation parameters are valid:

| fUnitName: | unit name (probeset ID). |
|----------------|-----------------------------------|
| fTranscriptID: | transcript_id (probeset ID). |
| fName: | gene name. |
| fSymbol: | gene symbol. |
| fAccession: | mRNA accession such as Refseq ID. |
| fEntrezID: | entrez ID. |
| fChromosome: | chromosome. |
| fStart: | start position. |
| fStop: | stop position. |
| fStrand: | strand on chromosome. |
| fCytoBand: | cytoband. |

For class AnalysisTreeSet the following varlist parameters are valid:

| mn1: | mean of group 1. |
|-------|---|
| mn2: | mean of group 2. |
| fc: | fold-change fc=mn2/mn1. |
| se: | standard error. |
| df: | degree of freedom. |
| stat: | t-statistic. |
| pval: | p-value. |
| nper: | number of permutations. |
| pcha: | p-chance. |
| padj: | adjusted p-value. |
| flag: | flag. |
| mask: | only rows with flag=1 will be exported. |

Value

If as .dataframe is TRUE, the data will be imported into the current R session as data.frame. Otherwise, NULL will be returned.

Author(s)

Christian Stratowa

export

See Also

export-methods

export

Export data as text files

Description

Export data from classes SchemeTreeSet, DataTreeSet, ExprTreeSet, or CallTreeSet
to outfile.

Usage

```
export.scheme(xps.scheme, treetype = character(0), varlist = "*", outfile = char
export.data(xps.data, treename = "*", treetype = "cel", varlist = "*", outfile =
export.expr(xps.expr, treename = "*", treetype = character(0), varlist = "*", ou
export.call(xps.call, treename = "*", treetype = character(0), varlist = "*", ou
export(object, ...)
```

Arguments

| xps.scheme | an object of type SchemeTreeSet. |
|--------------|--|
| xps.data | an object of type DataTreeSet. |
| xps.expr | an object of type ExprTreeSet. |
| xps.call | an object of type CallTreeSet. |
| treename | vector of tree names to export. |
| treetype | <pre>type of tree(s) to export, see validTreetype</pre> |
| varlist | names of tree leaves to export |
| outfile | name of output file. |
| sep | column separator |
| as.dataframe | if TRUE a data.frame will be returned. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| ••• | arguments treenames, treetype, varlist, outfile, sep, as. dataframe. |

Details

Export data from classes SchemeTreeSet, DataTreeSet, ExprTreeSet, or CallTreeSet
to outfile.

Parameter varlist lists the parameters to export:

- parameters are separated by ":", e.g. varlist="fInten:fStdev".

- for varlist=" * " all valid parameters will be exported.

For class DataTreeSet the following varlist parameters are valid:

export

| fInten: | intensities from e.g. tree.cel. |
|-----------|--|
| fStdev: | standard deviation from e.g. tree.cel. |
| fNPixels: | number of pixels from e.g. tree.cel. |
| fBg: | background values (background trees only). |

For classes <code>ExprTreeSet</code> and <code>CallTreeSet</code> varlist can contain annotation parameters and parameters of the resulting data.

Following varlist annotation parameters are valid:

| fUnitName: | unit name (probeset ID). |
|----------------|-----------------------------------|
| fTranscriptID: | transcript_id (probeset ID). |
| fName: | gene name. |
| fSymbol: | gene symbol. |
| fAccession: | mRNA accession such as Refseq ID. |
| fEntrezID: | entrez ID. |
| fChromosome: | chromosome. |
| fStart: | start position. |
| fStop: | stop position. |
| fStrand: | strand on chromosome. |
| fCytoBand: | cytoband. |

Following varlist parameters are valid for ExprTreeSet:

| fLevel: | expression level. |
|----------|---------------------|
| fStdev: | standard deviation. |
| fNPairs: | number of pairs. |

Following varlist parameters are valid for CallTreeSet:

| fCall: | detection call. |
|----------|--------------------|
| fPValue: | detection p-value. |

An example: varlist="fUnitName:fName:fSymbol:fLevel:fStdev:fEntrezID" export is a generic method to export data from ROOT trees as text file.

Value

If as .dataframe is TRUE, the data will be imported into the current R session as <code>data.frame</code>. Otherwise, <code>NULL</code> will be returned.

Author(s)

Christian Stratowa

See Also

export-methods

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"</pre>
```

export.root

data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.
export as table only
export(scheme.test3, treetype="idx", outfile="Test3_idx.txt",verbose=FALSE)
export as table and import as data.frame
ann <- export.scheme(scheme.test3, treetype="ann", outfile="Test3_ann.txt",as.dataframe=T
head(ann)
data <- export.data(data.test3, outfile="Test3_cel.txt",as.dataframe=TRUE,verbose=FALSE)
head(data)</pre>

export.root Export data from ROOT file

Description

Export data as text files directly from a ROOT file.

Usage

export.root(datafile = character(0), schemefile = character(0), treeset = character

Arguments

| datafile | name of ROOT data file including full path |
|--------------|--|
| schemefile | name of ROOT scheme file including full path |
| treeset | name of subdirectory in ROOT file where trees are stored |
| treename | name of ROOT tree to export. |
| treetype | type of tree(s) to export, see validTreetype. |
| varlist | names of tree leaves to export. |
| outfile | name of output file. |
| sep | column separator |
| as.dataframe | if TRUE a data.frame will be returned. |
| verbose | logical, if TRUE print status information. |

Details

Export data as text files directly from a ROOT file.

Value

If as .dataframe is TRUE, the data will be imported into the current R session as data.frame. Otherwise, NULL will be returned.

Author(s)

Christian Stratowa

See Also

export, export-methods

express

Examples

```
## export data directly from root file
schemefile <- paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/")
datafile <- paste(.path.package("xps"),"rootdata/DataTest3_cel.root",sep="/")
data <- export.root(datafile, schemefile, "DataSet", "*", "cel", "*", "DataOutFile.txt",
head(data)</pre>
```

express

Compute expression levels from raw data

Description

This function allows to combine different algorithms to compute expression levels, or to return the result for different algorithms only.

Usage

```
express(xps.data,
        filename = character(),
        filedir = getwd(),
        tmpdir = "",
        update = FALSE,
      # background correction
        bgcorrect.method = NULL,
        bgcorrect.select = character(),
        bgcorrect.option = character(),
        bgcorrect.params = list(),
      # normalization
        normalize.method = NULL,
        normalize.select = character(),
        normalize.option = character(),
        normalize.logbase = character(),
        normalize.params = list(),
      # expression values
        summarize.method = NULL,
        summarize.select = character(),
        summarize.option = character(),
        summarize.logbase = character(),
        summarize.params = list(),
      # reference values
        reference.index = 0,
        reference.method = "mean",
        reference.params = list(0),
      # misc.
        exonlevel = "",
        xps.scheme = NULL,
        add.data = TRUE,
        bufsize = 32000,
verbose = TRUE)
xpsPreprocess(object, ...)
```

Arguments

| xps.data | object of class DataTreeSet. |
|---------------|---|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| update | logical. If TRUE the existing ROOT data file filename will be updated. |
| bgcorrect.met | • • • |
| | background method to use. |
| bgcorrect.se | |
| | type of probes to select for background correction. |
| bgcorrect.opt | |
| 1 | type of background correction to use. |
| bgcorrect.pai | |
| normalize.met | vector of parameters for background method. |
| normarize.met | normalization method to use. |
| normalize.sel | |
| | type of probes to select for normalization. |
| normalize.opt | tion |
| | normalization option. |
| normalize.log | - |
| | logarithm base as character, one of '0', 'log', 'log2', 'log10'. |
| normalize.pa | |
| | vector of parameters for normalization method. |
| summarize.met | |
| summarize.sel | summarization method to use. |
| | type of probes to select for summarization. |
| summarize.opt | |
| | option determining the grouping of probes for summarization, one of 'tran- |
| | script', 'exon', 'probeset'; exon arrays only. |
| summarize.log | |
| | logarithm base as character, one of '0', 'log', 'log2', 'log10'. |
| summarize.pa | |
| | vector of parameters for summarization method. |
| reference.ind | |
| reference.met | index of reference tree to use, or 0. |
| rererence.met | for refindex=0, either trimmed mean or median of trees. |
| reference.pai | |
| Ť | vector of parameters for reference method. |
| exonlevel | exon annotation level determining which probes should be used for summariza- |
| | tion; exon/genome arrays only. |
| xps.scheme | optional alternative SchemeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| bufsize | integer which sets the buffer size of the tree branch baskets (default is 32000). |
| verbose | logical, if TRUE print status information. |
| object | object of class DataSet. |
| - | • |
| ••• | the arguments described above. |

express

Details

This function allows to combine different algorithms to compute expression levels, or to return the result for different algorithms only.

Please have a look at vignette "xpsPreprocess.pdf" for details on how to use function express.

xpsPreprocess is the DataSet method called by function express, containing the same parameters.

Value

An object of type DataTreeSet or ExprTreeSet.

Author(s)

Christian Stratowa

See Also

bgcorrect, normalize, summarize

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.</pre>
## compute rma with a single call to express()
expr.rma <- express(data.test3,"tmp_Test3Exprs",filedir=getwd(),tmpdir="",update=FALSE,</pre>
            bgcorrect.method="rma",bgcorrect.select="none",bgcorrect.option="pmonly:epane
            normalize.method="quantile", normalize.select="pmonly", normalize.option="trans
            summarize.method="medianpolish", summarize.select="pmonly", summarize.option="t
            verbose=FALSE)
## get expression data.frame
expr <- exprs(expr.rma)</pre>
head(expr)
## plot expression levels
if (interactive()) {
boxplot(expr.rma)
boxplot(log2(expr[,3:6]))
## Not run:
## examples using Affymetrix human tissue dataset (see also xps/examples/script4exon.R)
## example - exon array, e.g. HuEx-1_0-st-v2:
scmdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Schemes"</pre>
datdir <- "/Volumes/GigaDrive/CRAN/Workspaces/ROOTData"</pre>
scheme.exon <- root.scheme(paste(scmdir,"Scheme_HuEx10stv2r2_na25.root",sep="/"))</pre>
            <- root.data(scheme.exon, paste(datdir,"HuTissuesExon_cel.root", sep="/"))
data.exon
workdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Exon/hutissues/exon"
expr.rma <- express(data.exon, "HuExonExprs", filedir=workdir, tmpdir="", update=F,</pre>
            bgcorrect.method="rma",bgcorrect.select="antigenomic",bgcorrect.option="pmonl
            normalize.method="quantile", normalize.select="pmonly", normalize.option="trans
```

exprs-methods

summarize.method="medianpolish", summarize.select="pmonly", summarize.option="t exonlevel="metacore+affx")

End(Not run)

exprs-methods Get/Set Expression Values

Description

Get/set expression values from/for class ExprTreeSet.

Usage

```
exprs(object)
exprs(object, treenames = NULL) <- value</pre>
```

Arguments

| object | object of class ExprTreeSet. |
|-----------|---|
| treenames | character vector containing optional tree names to be used as subset. |
| value | data.frame containing expression values. |

Details

Get the expression values from slot data or set slot data to value.

Method exprs returns the expression values from slot data as data.frame, while replacement method exprs<- allows to replace slot data with a data.frame.

In order to create an ExprTreeSet containing only a subset of slot data, first export slot data using method exprs, create a character vector containing only treenames to be used in the subset, and then use replacement method exprs<- to replace slot data with the subset. Slots treenames and numtrees will be updated automatically.

Note: When creating character vector treenames it is sufficient to use the name part of the tree name w/o the extension.

Note: If you do not want to replace your current object, create first a copy of type ExprTreeSet by simply writing newobj <- oldobj, and use newobj for replacement. This is important since exprs<- does also update slots treenames and numtrees as already mentioned.

Author(s)

Christian Stratowa

See Also

pvalData, presCall
ExprTreeSet-class

Examples

```
## Not run:
## load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/'</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.</pre>
## create an ExprTreeSet
data.rma <- rma(data.test3,"tmp_TestRMA",tmpdir="",background="pmonly",normalize=TRUE,ver</pre>
## get expression values
value <- exprs(data.rma)</pre>
## selected treenames only
treenames <- c("TestA2", "TestB1")</pre>
## make a copy of your object if you do not want to replace it
subset.rma <- data.rma
## replace slot data with subset
exprs(subset.rma, treenames) <- value</pre>
str(subset.rma)
## End(Not run)
```

ExprTreeSet-class Class ExprTreeSet

Description

This class provides the link to the ROOT expression file and the ROOT trees contained therein. It extends class ProcesSet.

Objects from the Class

Objects are created using functions express, summarize or normalize, or the specialized functions rma, mas5 or mas4.

Slots

- exprtype: Object of class "character" representing the exression type, i.e. 'rma', 'mas5', 'mas4' or 'custom'.
- normtype: Object of class "character" representing the normalization type, i.e. 'mean', 'median', 'lowess', 'supsmu'.
- scheme: Object of class "SchemeTreeSet" providing access to ROOT scheme file.
- data: Object of class "data.frame". The data.frame can contain the data (e.g. expression levels) stored in ROOT data trees.
- params: Object of class "list" representing relevant parameters.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT data trees are stored, usually 'PreprocesSet'.
- settype: Object of class "character" describing the type of treeset stored in setname, usually 'preprocess'.

- rootfile: Object of class "character" representing the name of the ROOT data file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

Extends

Class "ProcesSet", directly. Class "TreeSet", by class "ProcesSet", distance 2.

Methods

attachExpr signature(object = "ExprTreeSet"): exports expression trees from ROOT
 expression file and and saves as data.frame data.

exprType signature(object = "ExprTreeSet"): extracts slot exprtype.

- exprType<- signature(object = "ExprTreeSet", value = "character"):replaces slot exprtype.
- exprs signature(object = "ExprTreeSet"): extracts the expression data.frame.
- exprs<- signature(object = "ExprTreeSet", value = "data.frame"):replaces
 the expression data.frame.</pre>
- mvaplot signature(x = "ExprTreeSet"): creates an MvA-plot.
- normType signature(object = "ExprTreeSet"): extracts slot normtype.
- normType<- signature(object = "ExprTreeSet", value = "character"):replaces slot normtype.
- **removeExpr** signature(object = "ExprTreeSet"): replaces data.frame data with an empty data.frame of dim(0,0).
- **se.exprs** signature (object = "ExprTreeSet"): extracts the standard deviation data.frame.

xpsNormalize signature(object = "ExprTreeSet"): applies normalization methods.

- **xpsPreFilter** signature(object = "ExprTreeSet"): applies prefiltering methods.
- xpsUniFilter signature(object = "ExprTreeSet"): applies unifiltering methods.

Author(s)

Christian Stratowa

See Also

related classes DataTreeSet, CallTreeSet.

Examples

showClass("ExprTreeSet")

extenPart

Description

Get the extension(s) of (tree) names.

Usage

extenPart(names, as.unique=TRUE)

Arguments

| names | vector of names. |
|-----------|--|
| as.unique | if TRUE return only unique extensions. |

Details

Extracts the extension part of names, e.g.of tree names of treename.treetype stored in a ROOT file.

Value

A vector of (unique) extensions.

Author(s)

Christian Stratowa

See Also

namePart

Examples

```
names <- c("TestA1.int", "TestA2.int")
extenPart(names)
extenPart(names, as.unique=FALSE)</pre>
```

farms

Description

This function converts a DataTreeSet into an ExprTreeSet using the Factor Analysis for Robust Microarray Summarization (FARMS) method.

Usage

```
farms(xps.data,
      filename = character(0),
      filedir = getwd(),
tmpdir = "",
      normalize = TRUE,
      weight
                 = 0.5,
      mu
                 = 0.0,
      mu = 0.0,
scale = 1.0,
      tol
                = 0.00001,
                 = 100,
      сус
      weighted = TRUE,
      version = "1.3.1",
option = "transcr
                 = "transcript",
      exonlevel = "",
      xps.scheme = NULL,
      add.data = TRUE,
      verbose = TRUE)
```

Arguments

| xps.data | object of class DataTreeSet. |
|-----------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| normalize | logical. If TRUE normalize data using quantile normalization. |
| weight | hyperparameter, usually set to 0.5 for version="1.3.1" and to 8.0 for version="1.3.0". |
| mu | hyperparameter allowing to correct for potential bias. |
| scale | scaling parameter, usually set to 1.0 for version="1.3.1" and to 2.0 for version="1.3.0". |
| tol | termination tolerance for EM algorithm. |
| сус | maximum number of cycles of EM algorithm. |
| weighted | logical, used only with version="1.3.1". Default is TRUE. |
| version | version of original farms package. Currently, version="1.3.1" and version="1.3.0" are implemented. Default is version="1.3.1". |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |

farms

| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
|------------|--|
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |

Details

This function computes the FARMS (Factor Analysis for Robust Microarray Summarization) expression measure described in Hochreiter et al. for both expression arrays and exon arrays.

Parameter version currently allows the user to choose between the original implementation of FARMS as implemented in package 'farms_1.3.0' or enhanced FARMS as implemented in package 'farms_1.3.1'. By default version="1.3.1" is used.

Parameter weight is a hyperparameter which determines the influence of the prior. For version="1.3.1" the value in the range of [0,1].

Parameter mu is a hyperparameter which allows to quantify different aspects of potential prior knowledge. Values near zero assume that most genes do not contain a signal and introduce a bias for loading matrix elements near zero.

Parameter weighted is a logical and indicates whether a weighted mean or a least square fit is used to summarize the loading matrix. It is applicable only to version="1.3.1".

For exon arrays it is necessary to supply the requested option and exonlevel.

Following options are valid for exon arrays:

transcript:expression levels are computed for transcript clusters, i.e. probe sets containing the same 'transcript_cheexon:expression levels are computed for exon clusters, i.e. probe sets containing the same 'exon_id', where expression levels are computed for individual probe sets, i.e. for each 'probeset_id'.

Following exonlevel annotations are valid for exon arrays:

| core: | probesets supported by RefSeq and full-length GenBank transcripts |
|---------------|---|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |
| metafull: | full meta-probesets. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Following exonlevel annotations are valid for whole genome arrays:

| core: | probesets with category 'unique', 'similar' and 'mixed'. |
|-----------|--|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Exon levels can also be combined, with following combinations being most useful:

exonlevel="metacore+affy":

core meta-probesets plus AFFX controls

```
exonlevel="core+extended": probesets with cDNA support
exonlevel="core+extended+full": supported plus predicted probesets
```

Exon level annotations are described in the Affymetrix whitepaper exon_probeset_trans_clust_whitepaper.pdf: "Exon Probeset Annotations and Transcript Cluster Groupings".

In order to use an alternative SchemeTreeSet set the corresponding SchemeSet xps.scheme.

Value

An ExprTreeSet

Note

The expression measure obtained with FARMS is given in linear scale, analogously to the expression measures computed with mas5 and rma.

For the analysis of many exon arrays it may be better to define a tmpdir, since this will store only the results in the main file and not e.g. background and normalized intensities, and thus will reduce the file size of the main file. For quantile normalization memory should not be an issue, however DFW depends on RAM unless you are using a temporary file.

Author(s)

Christian Stratowa

References

Hochreiter, S., Clevert D.-A., and Obermayer, K. (2006), A new summarization method for Affymetrix probe level data. Bioinformatics 22(8):943-949

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
data.farms <- farms(data.test3,"tmp_Test3FARMS",verbose=FALSE)
## get data.frame
expr.farms <- validData(data.farms)
head(expr.farms)</pre>
```

fcFilter-methods Fold-Change Filter

Description

This method initializes the Fold-Change Filter. The fold-change is determined by the mean value of group 2 divided by the mean value of group 1. The Fold-Change Filter flags all rows with: $flag = (fc \ge cutoff)$

Usage

```
fcFilter(object)
fcFilter(object, value)<-</pre>
```

Arguments

| object | object of class UniFilter. | |
|--------|--------------------------------------|------------|
| value | <pre>numeric vector c (cutoff,</pre> | direction) |

Details

The method fcFilter initializes the following parameters:

| cutoff: | the cutoff level for the filter. |
|------------|--|
| direction: | direction="both" (default): select up and downregulated genes. |
| | direction="up": select upregulated genes only. |
| | direction="down": select downregulated genes only. |

Value

An initialized UniFilter object.

Author(s)

Christian Stratowa

Examples

```
unifltr <- UniFilter()
fcFilter(unifltr) <- c(1.5,"both")
str(unifltr)</pre>
```

Filter-class Base Class Filter

Description

Base class for classes PreFilter and UniFilter.

Slots

numfilters: Object of class "numeric" giving the number of filters applied.

Methods

numberFilters signature(object = "Filter"): number of filters applied.

Author(s)

Christian Stratowa

See Also

related classes PreFilter, UniFilter.

Examples

showClass("Filter")

```
FilterTreeSet-class
```

Class FilterTreeSet

Description

This class provides the link to the ROOT filter file and the ROOT trees contained therein. It extends class ProcesSet.

Objects from the Class

Objects are currently created using function prefilter.

Slots

- filter: Object of class "Filter" currently providing access to the PreFilter settings.
- exprset: Object of class "ExprTreeSet" providing direct access to the ExprTreeSet used for filtering.
- callset: Object of class "CallTreeSet" providing direct access to the optional CallTreeSet used for filtering.
- scheme: Object of class "SchemeTreeSet" providing access to ROOT scheme file.
- data: Object of class "data.frame". The data.frame contains the data of the filter stored in ROOT filter trees.
- params: Object of class "list" representing relevant parameters.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT trees are stored, currently 'PreFilterSet'.
- settype: Object of class "character" describing the type of treeset stored in setname, currently 'prefilter'.
- rootfile: Object of class "character" representing the name of the ROOT file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

gapFilter-methods

Extends

Class "ProcesSet", directly. Class "TreeSet", by class "ProcesSet", distance 2.

Methods

```
validData signature(object = "FilterTreeSet"): extracts data.frame data.
```

Author(s)

Christian Stratowa

See Also

related classes AnalysisTreeSet.

Examples

```
showClass("FilterTreeSet")
```

gapFilter-methods Gap Filter

Description

This method initializes the Gap Filter.

The gapFilter looks for genes that might usefully discriminate between two groups. To do this we look for a gap in the ordered expression values. The gap should come in the central portion, thus a parameter window is defined to exclude jumps in the initial window values and the final window values.

```
The Gap Filter flags all rows with: flag = ((gap[i+1] - gap[i])/mean >= cutoff)
gapFilter(object)
gapFilter(object, value)<-</pre>
```

Arguments

| object | object of class PreFilter. | | | |
|--------|----------------------------|---------|-------|-----------|
| value | numeric vector c(cutoff, | window, | trim, | epsilon). |

Details

The method gapFilter initializes the following parameters:

| cutoff: | the cutoff level for the filter. | |
|----------|--|--|
| window: | trim value for the ordered expression levels (default is window=0.05). | |
| trim: | the trim value for trimmed mean (default is trim=0). | |
| epsilon: | value to replace mean (default is epsilon=0.01): | |
| | epsilon > 0: replace mean=0 with epsilon. | |
| | epsilon = 0: always set mean=1. | |

Note, that for epsilon = 0 the filter flags all rows with: (gap[i+1] - gap[i]) >= cutoff

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
gapFilter(prefltr) <- c(0.3,0.05,0.0,0.01)
str(prefltr)</pre>
```

getChipName Get Chip Name

Description

Get chip name from ROOT scheme file.

Usage

```
getChipName(rootfile)
```

Arguments

rootfile name of ROOT scheme file, including full path.

Details

Extracts the chip name directly from ROOT scheme file rootfile.

Value

 \boldsymbol{a} character with the chip name.

Author(s)

Christian Stratowa

See Also

getChipType,getNameType

Examples

```
## correct usage
getChipName(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"))
## incorrect usage
getChipName(paste(.path.package("xps"),"rootdata/DataTest3_cel.root",sep="/"))
```

getChipType Get Chip Type

Description

Get chip type from ROOT scheme file.

Usage

```
getChipType(rootfile)
```

Arguments

rootfile name of ROOT scheme file, including full path.

Details

Extracts the chip type directly from ROOT scheme file rootfile.

Value

a character with the chip type, either 'GeneChip' or 'ExonChip'.

Author(s)

Christian Stratowa

See Also

getChipName, getNameType

Examples

```
## correct usage
getChipType(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"))
## incorrect usage
getChipType(paste(.path.package("xps"), "rootdata/DataTest3_cel.root", sep="/"))
```

getDatatype Get Data Type

Description

Get data type corresponding to tree type.

Usage

getDatatype(treetype)

Arguments

treetype tree type.

Details

Get data type corresponding to tree type. Valid tree types are described in validTreetype.

Value

a character with the correct data type, i.e. 'rawdata', 'preprocess' or 'normation'.

Author(s)

Christian Stratowa

See Also

type2Exten,validTreetype

Examples

getDatatype("cel")
getDatatype("tbw")

getNameType Get Chip Name and Type

Description

Get chip name and type from ROOT scheme file.

Usage

```
getNameType(rootfile)
```

Arguments

rootfile name of ROOT scheme file, including full path.

Details

Extracts the chip name and type directly from ROOT scheme file rootfile.

Value

a list with parameters:

| chipname | chip name. |
|----------|---|
| chiptype | chip type, either 'GeneChip' or 'ExonChip'. |

Author(s)

Christian Stratowa

getNumberTrees

See Also

getChipName, getChipType

Examples

```
## correct usage
getNameType(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"))
## incorrect usage
getNameType(paste(.path.package("xps"),"rootdata/DataTest3_cel.root",sep="/"))
```

getNumberTrees Get Number of Trees

Description

Get number of trees stored in a ROOT file.

Usage

```
getNumberTrees(rootfile, treetype = "*", setname = NULL)
```

Arguments

| rootfile | name of ROOT file, including full path. |
|----------|---|
| treetype | tree type. |
| setname | name of ROOT subdirectory containing trees. |

Details

Extracts the number of trees of treetype stored in ROOT file rootfile. Valid tree types are listed in validTreetype. For treetype="*" the total number of trees in rootfile are returned.

If setname is provided, only trees in subdirectory setname are counted.

Value

Number of trees.

Author(s)

Christian Stratowa

Examples

```
getNumberTrees(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"))
getNumberTrees(paste(.path.package("xps"), "rootdata/DataTest3_cel.root", sep="/"))
```

getProbeInfo

Description

Get GeneChip probe information from root scheme file.

Usage

getProbeInfo(rootfile)

Arguments

rootfile name of ROOT scheme file, including full path.

Details

Extracts GeneChip probe information directly from ROOT scheme file rootfile.

Value

a list with parameters:

| nrows | physical number of rows in the array. |
|------------|--|
| ncols | physical number of columns in the array. |
| nprobes | number of probes on the array. |
| ncontrols | number of controls on the array. |
| ngenes | number of genes on the array. |
| nunits | number of units on the array. |
| nprobesets | umber of probesets on the array. |
| naffx | number of AFFX controls on the array. |

Author(s)

Christian Stratowa

Examples

```
getProbeInfo(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"))
```

getTreeData-methods

Export Tree Data

Description

Exports tree data from ROOT data file and and saves as data.frame.

Usage

```
getTreeData(object, treetype = "cel", varlist = "fInten")
```

Arguments

| object | Object of class "ProcesSet". |
|----------|---|
| treetype | type of tree to export, see validTreetype |
| varlist | names of tree leaves to export. |

Details

Exports tree leaves from ROOT data file and and saves as data.frame.

Value

A data.frame.

Author(s)

Christian Stratowa

See Also

export

getTreeNames Get Tree Names

Description

Get tree names stored in a ROOT file.

Usage

```
getTreeNames(rootfile, treetype = "*", setname = NULL, gettitle = FALSE)
```

Arguments

| rootfile | name of ROOT file, including full path. |
|----------|---|
| treetype | tree type. |
| setname | name of ROOT subdirectory containing trees. |
| gettitle | If TRUE the titles of the trees will be returned. |

Details

Extracts the tree names of treetype stored in ROOT file rootfile. Valid tree types are listed in validTreetype. For treetype="*" names for all trees in rootfile are returned.

If setname is provided, only tree names in subdirectory setname are returned.

Value

A vector of tree names. For gettitle=TRUE a vector of tree titles.

Author(s)

Christian Stratowa

Examples

```
getTreeNames(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"))
getTreeNames(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"), "scm")
getTreeNames(paste(.path.package("xps"), "rootdata/DataTest3_cel.root", sep="/"))
```

highFilter-methods Upper Threshold Filter

Description

This method initializes the Upper Threshold Filter. The cutoff value defines the upper threshold for allowed expression levels. If e.g. the number of samples exceeding this cutoff value is greater than parameter then the corresponding dataframe row is flagged, i.e. flag = 0. The Upper Threshold Filter flags all rows with: flag = (sum(expression[i] <= cutoff) >= parameter)

Usage

```
highFilter(object)
highFilter(object, value)<-</pre>
```

Arguments

| object | object of class PreFilter. | | |
|--------|-----------------------------|------------|-------------|
| value | character vector c (cutoff, | parameter, | condition). |

Details

The method highFilter initializes the following parameters:

| cutoff: | the upper threshold level for the filter. |
|------------|---|
| parameter: | this value depends on the condition used: |
| condition: | condition="samples": number of samples (default): |
| | condition="percent": percent of samples. |
| | condition="mean": mean value of samples. |
| | condition="percentile": percentile of samples. |
| | |

hist-methods

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
highFilter(prefltr) <- c(14.5,75.0,"percent")
str(prefltr)</pre>
```

hist-methods Density Estimate

Description

Plot the density estimates for each sample.

Usage

```
hist(x, which = "", size = 0, transfo = log2, ylab = "density", xlab
= "log intensity", type = "l", col = 1:6, ...)
```

Arguments

| х | object of class DataTreeSet or ExprTreeSet. |
|---------|---|
| which | type of probes to be used, for details see validData. |
| size | length of sequence to be generated as subset. |
| transfo | a valid function to transform the data, usually "log2", or "0". |
| ylab | a title for the y axis. |
| xlab | a title for the x axis. |
| type | type for the plot. |
| col | colors to use for the different arrays. |
| | optional arguments to be passed to plot. |

Details

 $Plots \ the \ non-parametric \ density \ estimates \ using \ values \ contained \ in \ the \ columns \ of \ slot \ data.$

For a DataTreeSet object, data must first be attached using method attachInten.

Note

For exon array raw data only a limited number of samples can be displayed as density plot due to memory limitations. To display all samples it is proposed to use function root.density instead.

Author(s)

Christian Stratowa

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
## need to attach scheme mask and probe intensities
data.test3 <- attachMask(data.test3)
data.test3 <- attachInten(data.test3)
if (interactive()) {
hist(data.test3)
}
## optionally remove mask and data to free memory
data.test3 <- removeInten(data.test3)
data.test3 <- removeMask(data.test3)</pre>
```

```
image.dev
```

Image for Device

Description

Creates an image for each sample for the selected device.

Usage

```
image.dev(x, bg = FALSE, transfo = log2, col = gray((0:64)/64), names = "namepar
```

Arguments

| Х | object of class DataTreeSet. |
|---------|--|
| bg | logical. If FALSE, intensities from slot data will be used; if TRUE, background intensities from slot bgrd will be used. |
| transfo | a valid function to transform the data, usually "log2", or "0". |
| col | color range for intensities. |
| names | optional vector of sample names. |
| xlab | a title for the x axis. |
| ylab | a title for the y axis. |
| mar | plot margins. |
| dev | graphics device to plot to, i.e. one of "screen", "jpeg", "png", "pdf" or "ps". |
| outfile | the name of the output file. |
| W | the width of the device in pixels. |
| h | the height of the device in pixels. |
| | optional arguments to be passed to image. |

image-methods

Details

Creates an image for each array for the selected graphics device.

For bgrd=TRUE the distribution of the computed background intensities will be shown; this can be useful to see potential density gradients caused by hybridization conditions. For the computation of background intensities see function bgcorrect; it is suggested to use bgcorrect.mas4 to identify density gradients.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as image.

Note

Data must first be attached to class DataTreeSet using method attachInten.

Author(s)

Christian Stratowa

See Also

image-methods, image

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
## need to attach scheme mask and data
data.test3 <- attachMask(data.test3)
data.test3 <- attachInten(data.test3)
if (interactive()) {
image.dev(data.test3)
}
## to avoid memory comsumption of R remove data:
data.test3 <- removeInten(data.test3)
data.test3 <- removeMask(data.test3)</pre>
```

image-methods Display an Image

Description

Creates an image for each sample.

Usage

```
image(x, bg = FALSE, transfo = log2, col = gray((0:64)/64), names
= "namepart", xlab = "", ylab = "", ...)
```

Arguments

| Х | object of class DataTreeSet. |
|---------|--|
| bg | logical. If FALSE, intensities from slot data will be used; if TRUE, background intensities from slot bgrd will be used. |
| transfo | a valid function to transform the data, usually "log2", or "0". |
| col | color range for intensities. |
| names | optional vector of sample names. |
| | optional arguments to be passed to image. |
| | |

Details

Creates an image for each array. For bgrd=TRUE the distribution of the computed background intensities will be shown; this can be useful to see potential density gradients caused by hybridization conditions. For the computation of background intensities see function bgcorrect; it is suggested to use bgcorrect.mas4 to identify density gradients.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as image.

Note

Data must first be attached to class DataTreeSet using method attachInten.

Author(s)

Christian Stratowa

See Also

image.dev, image

import.data Import CEL files into a DataTreeSet

Description

Import the Affymetrix CEL files into a ROOT file and create S4 class DataTreeSet

Usage

```
import.data(xps.scheme,
    filename = character(0),
    filedir = getwd(),
    celdir = NULL,
    celfiles = "*",
    celnames = NULL,
    project = NULL,
    verbose = TRUE)
```

import.data

Arguments

| xps.scheme | a SchemeTreeSet containing the correct scheme for the CEL-files |
|------------|---|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| celdir | system directory containing the CEL-files for corresponding scheme. |
| celfiles | optional vector of CEL-files to be imported. |
| celnames | optional vector of names which should replace the CEL-file names. |
| project | optional class ProjectInfo. |
| verbose | logical, if TRUE print status information. |

Details

import.data is used to import CEL-files from directory celdir into a ROOT data file. To import only a subset of CEL-files, list these CEL-files as vector celfiles.

To import CEL-files from different directories, vector celfiles must contain the full path for each CEL-file and celdir must be celdir=NULL.

Currently, the following types of Affymetrix CEL-files can be imported: text (version 3), xml, binary (xda), generic (agcc,calvin)

An S4 class DataTreeSet will be created, serving as R wrapper to the ROOT data file filename.

Use function root.data to access the ROOT data file from new R sessions to avoid creating a new ROOT data file for every session.

Value

A DataTreeSet object.

Note

As mentioned above, use function root.data to access the ROOT data file from new R sessions to avoid creating a new ROOT data file for every R session.

Do not separate filename of ROOT files with dots, use underscores, e.g. do not use filename="Data.Test3" but use filename="Data_Test3" or filename="DataTest3" instead.

To every ROOT data file the extension "_cel" is attached to filename to easily recognize ROOT data files containing the raw CEL data, e.g. for filename="Data_Test3" the final name is "Data_Test3_cel.root". Extension "root" is added automatically, so that ROOT is able to recognize the file as ROOT file.

Once a ROOT file is created it can not be overwritten, it must be deleted manually first. Only ROOT files called "tmp" or with filename starting with "tmp_" will be re-created automatically.

If CEL-file names contain dots and/or colons as characters, these characters will be replaced by underscores. It is recommended to use parameter celnames to create shorter CEL names and to replace special characters.

Author(s)

Christian Stratowa

See Also

root.data,DataTreeSet

Examples

import.exon.scheme Import CLF, PGF and annotation files into a SchemeTreeSet

Description

Import the Affymetrix CLF, PGF, and probeset and transcript annotation files into a ROOT file and create S4 class SchemeTreeSet

Usage

```
import.exon.scheme(filename = character(0),
    filedir = getwd(),
    layoutfile = character(0),
    schemefile = character(0),
    probeset = character(0),
    transcript = character(0),
    control = "",
    add.mask = FALSE,
    verbose = TRUE)
```

Arguments

| filename | file name of ROOT scheme file. |
|------------|---|
| filedir | system directory where ROOT scheme file should be stored. |
| layoutfile | name of CLF-file, including full path. |
| schemefile | name of PGF-file, including full path. |
| probeset | name of probeset annotation-file, including full path. |
| transcript | name of transcript annotation-file, including full path. |
| control | optional name of controls.ps-file, including full path. |
| add.mask | logical. If ${\tt TRUE}\xspace$ mask information will be included as slot <code>mask</code> . |
| verbose | logical, if TRUE print status information. |

import.exon.scheme

Details

import.exon.scheme is used to import all information for an Affymetrix exon array into a ROOT scheme file, including CLF and PGF-files, and the current Afymetrix probeset and transcript annotation files.

An S4 class SchemeTreeSet will be created, serving as R wrapper to the ROOT scheme file filename.

Since a new ROOT scheme file needs only to be created when new annotation files are available from the Affymetrix website, it is recommended to store all ROOT scheme files in a commonly accessible system directory filedir.

Use function root.scheme to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every session.

Value

A SchemeTreeSet object.

Warning

The current version of 'xps' should be able to import all Affymetrix exon array annotation files up to November 2008. However, since Affymetrix is still changing the headers and/or columns of the annotation files, future annotation files may require adaptation of the source code, thus the current version of 'xps' may not be able to read these files.

Note

As mentioned above, use function root.scheme to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every R session.

Do not separate filename of ROOT files with dots, use underscores, e.g. do not use filename="Scheme_HuEx10s but use filename="Scheme_HuEx10stv2r2_na27" instead. Extension "root" is added automatically, so that ROOT is able to recognize the file as ROOT file.

Do not set add.mask=TRUE unless you know that your computer has sufficient RAM.

Do not add item control unless you want to use one of the old annotation files where the probeset annotation file does not contain the AFFX controls.

Author(s)

Christian Stratowa

See Also

import.expr.scheme, root.scheme, SchemeTreeSet

Examples

```
## Not run:
## define paths
scmdir <- "/common/path/schemes"
libdir <- "/my/path/Affy/libraryfiles"
anndir <- "/my/path/Affy/Annotation"
## create scheme for HuEx-1_0-st-v2.r2 Exon array
scheme.huex10stv2r2.na27 <- import.exon.scheme("Scheme_HuEx10stv2r2_na27",filedir=scmdir,</pre>
```

```
layoutfile=paste(libdir,"HuEx-1_0-st-v2_libraryfile/HuEx-1_0-
                             schemefile=paste(libdir,"HuEx-1_0-st-v2_libraryfile/HuEx-1_0-
                             probeset=paste(anndir,"HuEx-1_0-st-v2.na27.hg18.probeset.csv'
                             transcript=paste(anndir,"HuEx-1_0-st-v2.na27.hg18.transcript.
## access ROOT scheme file from new R session
scheme.huex10stv2r2 <- root.scheme(paste(scmdir,"Scheme_HuEx10stv2r2_na27.root",sep="/"))</pre>
## create scheme for HuGene-1_0-st-v1.r4 as exon array
scheme.hugene10stv1r4.na27 <- import.exon.scheme("Scheme_HuGene10stv1r4_na27_2",filedir=s</pre>
                               layoutfile=paste(libdir,"HuGene-1_0-st-v1.r4.analysis-lib-f
                               schemefile=paste(libdir,"HuGene-1_0-st-v1.r4.analysis-lib-f
                              probeset=paste(anndir,"HuGene-1_0-st-v1.na27.2.hg18.probeset
                               transcript=paste(anndir,"HuGene-1_0-st-v1.na27.hg18.transcr
## access ROOT scheme file from new R session
scheme.hugenel0stvlr4 <- root.scheme(paste(scmdir,"Scheme_HuGenel0stvlr4_na27_2.root",sep</pre>
## create scheme for HuEx-1_0-st-v2.r2 Exon array with the old annotation file
scheme.huex10stv2r2.old <- import.exon.scheme("Scheme_HuEx10stv2r2_old",filedir=scmdir,</pre>
                            layoutfile=paste(libdir,"HuEx-1_0-st-v2_libraryfile/HuEx-1_0-s
                            schemefile=paste(libdir,"HuEx-1_0-st-v2_libraryfile/HuEx-1_0-st
                           probeset=paste(anndir,"HuEx-1_0-st-probeset-annot.csv",sep="/'
                            transcript=paste(anndir, "HuEx-1_0-st-transcript-annot.csv", sep
                            control=paste(libdir,"HuEx-1_0-st-v2_libraryfile/HuEx-1_0-st-w
## End(Not run)
```

import.expr.scheme Import CDF, probe and annotation files into a SchemeTreeSet

Description

Import the Affymetrix CDF, probe and annotation files into a ROOT file and create S4 class Scheme-TreeSet

Usage

```
import.expr.scheme(filename = character(0),
    filedir = getwd(),
    schemefile = character(0),
    probefile = character(0),
    annotfile = character(0),
    chipname = NULL,
    add.mask = FALSE,
    verbose = TRUE)
```

Arguments

| filename | file name of ROOT scheme file. |
|------------|---|
| filedir | system directory where ROOT scheme file should be stored. |
| schemefile | name of CDF-file, including full path. |
| probefile | name of probe-file, including full path. |

| annotfile | name of annotation-file, including full path. |
|-----------|--|
| chipname | optional chip name when using an alternative CDF-file. |
| add.mask | logical. If TRUE mask information will be included as slot mask. |
| verbose | logical, if TRUE print status information. |

Details

import.expr.scheme is used to import all information for an Affymetrix expression array into a ROOT scheme file, including CDF-file, the corresponding probe file, and the current Afymetrix annotation file.

Usually, chipname is extracted from the name of the CDF-file, however, when using an alternative CDF-file, e.g. from BrainArray or AffyProbeMiner, a chipname must be supplied which starts with (or contains) the exact Affymetrix chip name.

An S4 class <code>SchemeTreeSet</code> will be created, serving as R wrapper to the <code>ROOT</code> scheme file filename.

Since a new ROOT scheme file needs only to be created when a new annotation file is available from the Affymetrix website, it is recommended to store all ROOT scheme files in a commonly accessible system directory filedir.

Use function root.scheme to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every session.

Value

A SchemeTreeSet object.

Note

As mentioned above, use function root.scheme to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every R session.

Do not separate filename of ROOT files with dots, use underscores, e.g. do not use filename="Scheme.Test3.mbut use filename="Scheme_Test3_na27" or simply filename="SchemeTest3na27" instead. Extension "root" is added automatically, so that ROOT is able to recognize the file as ROOT file.

For a few probesets, parsing the Affymetrix annotation files will provide ambiguous results. Setting verbose=11 will list these probesets.

Author(s)

Christian Stratowa

See Also

import.exon.scheme, import.genome.scheme, root.scheme, SchemeTreeSet

Examples

```
## Not run:
## define paths
scmdir <- "/common/path/schemes"
libdir <- "/my/path/Affy/libraryfiles"
anndir <- "/my/path/Affy/Annotation"</pre>
```

import.genome.scheme

Import CLF, PGF and annotation files into a SchemeTreeSet

Description

Import the Affymetrix CLF, PGF and transcript annotation files into a ROOT file and create S4 class SchemeTreeSet

Usage

```
import.genome.scheme(filename = character(0),
    filedir = getwd(),
    layoutfile = character(0),
    schemefile = character(0),
    transcript = character(0),
    add.mask = FALSE,
    verbose = TRUE)
```

Arguments

| filename | file name of ROOT scheme file. |
|------------|---|
| filedir | system directory where ROOT scheme file should be stored. |
| layoutfile | name of CLF-file, including full path. |
| schemefile | name of PGF-file, including full path. |
| transcript | name of transcript annotation-file, including full path. |
| add.mask | logical. If ${\tt TRUE}\xspace$ mask information will be included as slot <code>mask</code> . |
| verbose | logical, if TRUE print status information. |

import.genome.scheme

Details

import.genome.scheme is used to import all information for an Affymetrix whole genome array into a ROOT scheme file, including CLF and PGF-files, and the current Afymetrix transcript annotation files.

An S4 class SchemeTreeSet will be created, serving as R wrapper to the ROOT scheme file filename.

Since a new ROOT scheme file needs only to be created when new annotation files are available from the Affymetrix website, it is recommended to store all ROOT scheme files in a commonly accessible system directory filedir.

Use function root.scheme to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every session.

Value

A SchemeTreeSet object.

Warning

The current version of 'xps' is able to import all Affymetrix genome array annotation files up to November 2008, i.e. all files of release 3 (r3) and earlier. However, in January 2009 Affymetrix has updated all CLF, PGF and annotation files to release 4 (r4) and added a new probeset annotation file, thus in effect changing the whole genome arrays to exon arrays!

Thus, for release 4 (r4) files, function import.genome.scheme can no longer be used, but you must use function import.exon.scheme instead (see examples).

Note

As mentioned above, use function root.scheme to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every R session.

Do not separate filename of ROOT files with dots, use underscores, e.g. do not use filename="Scheme_HuGene10stv1_na27" instead. Extension "root" is added automatically, so that ROOT is able to recognize the file as ROOT file.

Do not set add.mask=TRUE unless you know that your computer has sufficient RAM.

Do not add item control unless you want to use one of the old annotation files where the probeset annotation file does not contain the AFFX controls.

Author(s)

Christian Stratowa

See Also

import.exon.scheme, root.scheme, SchemeTreeSet

Examples

```
## Not run:
## define paths
scmdir <- "/common/path/schemes"
libdir <- "/my/path/Affy/libraryfiles"
anndir <- "/my/path/Affy/Annotation"</pre>
```

ini.call

Informative/Non-Informative Call

Description

Computes the Informative/Non-Informative Call for the exclusion of non-informative probe sets.

Usage

| ini.call(xps.data, | | |
|--------------------|---|---------------|
| filename | = | character(0), |
| filedir | = | getwd(), |
| tmpdir | = | "", |
| weight | = | 0.5, |
| mu | = | 0.0, |
| scale | = | 1.0, |
| tol | = | 0.00001, |
| сус | = | 100, |
| alphal | = | 0.4, |
| alpha2 | = | 0.6, |
| version | = | "1.3.1", |
| option | = | "transcript", |
| exonlevel | = | "", |
| xps.scheme | = | NULL, |
| add.data | = | TRUE, |
| verbose | = | TRUE) |

xpsINICall(object, ...)

Arguments

| xps.data | object of class DataTreeSet. |
|----------|---|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| weight | hyperparameter, usually set to 0.5 for version="1.3.1" and to 8.0 for version="1.3.0". |
| mu | hyperparameter allowing to correct for potential bias. |
| scale | scaling parameter, usually set to 1.0 for version="1.3.1" and to 2.0 for version="1.3.0". |

ini.call

| tol | termination tolerance for EM algorithm. |
|------------|--|
| сус | maximum number of cycles of EM algorithm. |
| alphal | a significance threshold in (0,alpha2). |
| alpha2 | a significance threshold in (alpha1,1.0). |
| version | version of original farms package. Currently, version="1.3.1" and version="1.3.0" are implemented. Default is version="1.3.1". |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE call data will be added to slots data and detcall. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | the arguments described above. |

Details

In contrast to mas5.call this function quantifies the signal-to-noise ratio for each probe set, as described in Talloen et al. Thus, the returned p-values and detection calls have a different meaning:

The p-value that is returned estimates the signal-to-noise ratio (SNR):

P-values (SNR) of less than 0.5 indicate that there is more signal than noise and the corresponding genes are considered to be 'informative' for further analysis. In contrast, values greater than 0.5 indicate 'non-informative' genes.

The informative call is computed by thresholding the p-value as in:

call "P" if p-value < alpha1 call "M" if alpha1 <= p-value < alpha2 call "A" if alpha2 <= p-value</pre>

Here "P" should be considered as informative "I", "M" as marginally informative, and "A" as non-informative "NI".

The defaults for alpha1=0.4 and alpha2=0.6 are set to allow "M" calls. In order to get the same results as package 'farms_1.3.1', you need to set alpha1=0.5 and alpha2=0.5.

For exon/genome arrays it is necessary to supply option and exonlevel.

Following options are valid for exon arrays only:

transcript:expression levels are computed for transcript clusters, i.e. probe sets containing the same 'transcript_cheexon:expression levels are computed for exon clusters, i.e. probe sets containing the same 'exon_id', where expression levels are computed for individual probe sets, i.e. for each 'probeset_id'.

Following exonlevel annotations are valid for exon arrays:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |

ini.call

| full meta-probesets. |
|---------------------------|
| ambiguous probesets only. |
| standard AFFX controls. |
| combination of above. |
| |

Following exonlevel annotations are valid for whole genome arrays:

| core: | probesets with category 'unique' and 'mixed'. |
|-----------|---|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |
| all: | combination of above. |

Exon levels can also be combined, with following combinations being most useful:

| exonlevel="metacore+affy": | core meta-probesets plus AFFX controls |
|---------------------------------|--|
| exonlevel="core+extended": | probesets with cDNA support |
| exonlevel="core+extended+full": | supported plus predicted probesets |

Exon level annotations are described in the Affymetrix whitepaper 'exon_probeset_trans_clust_whitepaper.pdf'.

 $In order \ to \ use \ an \ alternative \ {\tt SchemeTreeSet} \ set \ the \ corresponding \ SchemeTreeSet \ {\tt xps.scheme.schemeTreeSet} \ set \ the \ corresponding \ SchemeTreeSet \ {\tt xps.schemeTreeSet} \ set \ the \ schemeTreeSet \ set \ set \ set \ schemeTreeSet \ set \ se$

xpsINICall is the DataTreeSet method called by function ini.call, containing the same parameters.

Value

A CallTreeSet

Note

Since I/NI-calls distinguish only between informative and non-informative genes, the calls are identical for all samples.

Author(s)

Christian Stratowa

References

Talloen, W., Clevert D.-A., Hochreiter, S., Amaratunga, D., Bijnens, J., Kass, S., and Gohlmann, H.W.H. (2006), I/NI-calls for the exclusion of non-informative genes: a highly effective filtering tool for microarray data. Bioinformatics 23(21):2897-2902

See Also

farms, mas5.call

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

initialize-methods

```
## I/NI call
call.ini <- ini.call(data.test3, "tmp_Test3INI", verbose=FALSE)
## get data.frames
snr.ini <- pvalData(call.ini)
inf.ini <- presCall(call.ini)
head(snr.ini)
head(inf.ini)
## plot results
if (interactive()) {
callplot(call.ini)
}
rm(scheme.test3, data.test3)
gc()</pre>
```

initialize-methods Initialize Classes

Description

Initialize S4 classes.

Methods

Internal method to initialize S4 classes.

intensity-methods Get/Set Data Values

Description

Get/set data values from/for class DataTreeSet.

Usage

```
intensity(object)
intensity(object, filename = NULL, verbose = FALSE) <- value</pre>
```

Arguments

| object | object of class DataTreeSet. |
|----------|---|
| filename | $\label{eq:character} \mbox{ character vector containing optional ROOT file name.}$ |
| verbose | logical, if TRUE print status information. |
| value | data.frame containing expression values. |

Details

Get the intensity values from slot data or set slot data to value.

Method intensity returns the data values from slot data as data.frame, while replacement method intensity <- allows to replace slot data with a data.frame.

Using replacement method intensity<- with default settings will not change the data stored in the ROOT data file, and thus will not have any effect on subsequent processing methods. If you really want to use the replacement data for further processing you must supply a new ROOT filename. This will export each intensity column of value as CEL-file (version 3), which will then be imported into the new ROOT data file filename.

Warning: Do not use replacement method intensity <- until you really know what you are doing!

Note: The first two columns of replacement data.frame value must be the (X,Y) coordinates, followed by the intensities whereby the number of intensity columns must be identical to the columns to be replaced.

Note: If you do not want to replace your current object, create first a copy of type DataTreeSet by simply writing newobj <- oldobj, and use newobj for replacement. This is important since intensity<- does also update slots rootfile, filedir and treenames when a new filename was chosen.

Note: The CEL-files created are fully functional CEL-files (version 3), however some header rows such as GridCornerUL, AlgorithmParameters, and some of the data in DatHeader are placeholders only.

Warning: The CEL-files created WILL REPLACE THE ORIGINAL CEL-files, if they have identical names to the original CEL-files and the original CEL-files are located in the working directory. Thus the original CEL-files should preferable be located in directory celdir of function import.data.

Author(s)

Christian Stratowa

See Also

validData

Examples

```
## Not run:
## load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
## get intensity values
value <- intensity(data.test3)</pre>
```

make a copy of your object if you do not want to replace it newdata.test3 <- data.test3</pre>

```
## replace slot data with value
intensity(newdata.test3, "ReplacementData", FALSE) <- value
str(newdata.test3)</pre>
```

now you can create an ExprTreeSet using the new intensity data

isROOTFile

data.rma <- rma(newdata.test3, "ReplacementRMA", tmpdir="", background="none", normalize=TRUE
End(Not run)</pre>

isROOTFile Test for ROOT File

Description

Test if a file is a valid ROOT file.

Usage

isROOTFile(filename)

Arguments

filename name of ROOT file, including full path.

Value

Return TRUE if file filename is a valid ROOT file.

Author(s)

Christian Stratowa

See Also

existsROOTFile

Examples

isROOTFile(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/"))

lowFilter-methods Lower Threshold Filter

Description

This method initializes the Lower Threshold Filter. The cutoff value defines the lower threshold for allowed expression levels. If e.g. the number of samples lower than this cutoff value is greater than parameter then the corresponding dataframe row is flagged, i.e. flag = 0.

The Lower Threshold Filter flags all rows with: flag = (sum(expression[i] >= cutoff)
>= parameter)

Usage

lowFilter(object)
lowFilter(object, value)<-</pre>

Arguments

| object | object of class PreFilter. | | |
|--------|-----------------------------|------------|-------------|
| value | character vector c (cutoff, | parameter, | condition). |

Details

The method lowFilter initializes the following parameters:

```
cutoff:the lower threshold level for the filter.parameter:this value depends on the condition used:condition:condition="samples": number of samples (default):condition="percent": percent of samples.condition="mean": mean value of samples.condition="percentile": percentile of samples.
```

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
lowFilter(prefltr) <- c(4.0,3,"samples")
str(prefltr)</pre>
```

madFilter-methods Median Absolute Deviation Filter

Description

```
This method initializes the Median Absolute Deviation Filter.
The MAD Filter flags all rows with: flag = (mad >= cutoff)
Usage
madFilter(object)
madFilter(object, value)<-</pre>
```

Arguments

| object | object of class PreFilter. | |
|--------|-------------------------------------|-----------|
| value | <pre>numeric vector c(cutoff,</pre> | epsilon). |

Details

The method madFilter initializes the following parameters:

| cutoff: | the cutoff level for the filter. |
|----------|--|
| epsilon: | value to replace mean (default is epsilon=0.01). |

mas4

Note, that epsilon has no effect on mad.

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
madFilter(prefltr) <- c(0.5,0.01)
str(prefltr)</pre>
```

mas4

MAS 4.0 Expression Measure

Description

This function converts a DataTreeSet into an ExprTreeSet using the XPS implementation of Affymetrix's MAS 4.0 expression measure.

Usage

```
mas4(xps.data,
filename = character(0),
filedir = getwd(),
tmpdir = "",
normalize = FALSE,
sc = 500,
option = "transcript",
exonlevel = "",
update = FALSE,
xps.scheme = NULL,
add.data = TRUE,
verbose = TRUE)
```

xpsMAS4(object, ...)

Arguments

| xps.data | object of class DataTreeSet. |
|-----------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| normalize | logical. If ${\tt TRUE}$ scale normalization is used after an ${\tt ExprTreeSet}$ is obtained. |
| SC | value at which all arrays will be scaled to. |

| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
|------------|--|
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| update | logical. If TRUE the existing ROOT data file filename will be updated. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | arguments filename, filedir, tmpdir, option, exonlevel, xps. scheme. |

Details

This function computes the Affymetrix MAS 4.0 expression measure, i.e. the 'Average Difference' expression level, as implemented in XPS.

If normalize=TRUE then the expression levels will be scaled to sc. For sc=0 the expression levels will be scaled to the mean expression level.

xpsMAS4 is the DataTreeSet method called by function mas4, however, expression levels will not be scaled to a common mean expression level.

For further details see mas5.

Value

An ExprTreeSet

Note

In contrast to function mas4, expression levels computed with xpsMAS4 will not be scaled to a common mean expression level.

Author(s)

Christian Stratowa

References

Affymetrix (1999) GeneChip Expression Analysis Algorithm Tutorial, Affymetrix Inc., Santa Clara, CA.

See Also

xpsMAS4, express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

data.mas4 <- mas4(data.test3,"tmp_Test3MAS4",tmpdir="",normalize=TRUE,sc=500, update=TRUE
mas5.call

```
## get data.frame
expr.mas4 <- validData(data.mas4)
head(expr.mas4)
## plot results (negative expression values!)
if (interactive()) {
boxplot(expr.mas4)
}
rm(scheme.test3, data.test3)
gc()</pre>
```

mas5.call MAS 5.0 Absolute Detection Call

Description

Performs the Wilcoxon signed rank-based gene expression presence/absence detection algorithm first implemented in the Affymetrix Microarray Suite version 5.

Usage

```
mas5.call(xps.data,
    filename = character(0), filedir = getwd(), tmpdir = "",
    tau = 0.015, alpha1 = 0.04, alpha2 = 0.06, ignore.saturated = TRUE, bg
    option = "transcript", exonlevel = "", xps.scheme = NULL, add.data = T
```

```
xpsMAS5Call(object, ...)
```

Arguments

| xps.data | object of class DataTreeSet. | |
|------------------|--|--|
| filename | file name of ROOT data file. | |
| filedir | system directory where ROOT data file should be stored. | |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. | |
| tau | a small positive constant. | |
| alpha1 | a significance threshold in (0,alpha2). | |
| alpha2 | a significance threshold in (alpha1,0.5). | |
| ignore.saturated | | |
| | logical. If TRUE do the saturation correction described in the paper, with a saturation level of 46000. | |
| bgcorrect.option | | |
| | bgcorrect option determining wether to subtract background first, one of 'none' or 'correctbg'. | |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. | |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. | |
| xps.scheme | optional alternative SchemeTreeSet. | |

mas5.call

| add.data | logical. If TRUE call data will be added to slots data and detcall. |
|----------|---|
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | the arguments described above. |

Details

This function performs the hypothesis test:

H0: median(Ri) = tau, corresponding to absence of transcript H1: median(Ri) > tau, corresponding to presence of transcript

where Ri = (PMi - MMi) / (PMi + MMi) for each i a probe-pair in the probe-set represented by data. The p-value that is returned estimates the usual quantity:

Pr(observing a more "present looking" probe-set than data | data is absent)

Small p-values imply presence while large ones imply absence of transcript. The detection call is computed by thresholding the p-value as in:

call "P" if p-value < alpha1 call "M" if alpha1 <= p-value < alpha2 call "A" if alpha2 <= p-value

The defaults for tau, alpha1 and alpha2 correspond to those in MAS5.0 for expression arrays. However, when using this function for exon or whole genome arrays, new values for alpha1 and alpha2 must be determined. The recommended function for exon/genome arrays is dabg.call.

In order to use an alternative SchemeTreeSet set the corresponding SchemeTreeSet xps.scheme.

xpsMAS5Call is the DataTreeSet method called by function mas5.call, containing the same parameters.

Value

A CallTreeSet

Author(s)

Christian Stratowa

References

Liu, W. M. and Mei, R. and Di, X. and Ryder, T. B. and Hubbell, E. and Dee, S. and Webster, T. A. and Harrington, C. A. and Ho, M. H. and Baid, J. and Smeekens, S. P. (2002) Analysis of high density expression microarrays with signed-rank call algorithms, Bioinformatics, 18(12), pp. 1593-1599.

Liu, W. and Mei, R. and Bartell, D. M. and Di, X. and Webster, T. A. and Ryder, T. (2001) Rankbased algorithms for analysis of microarrays, Proceedings of SPIE, Microarrays: Optical Technologies and Informatics, 4266.

Affymetrix (2002) Statistical Algorithms Description Document, Affymetrix Inc., Santa Clara, CA, whitepaper.http://www.affymetrix.com/support/technical/whitepapers/sadd_whitepaper.pdf

See Also

dabg.call

mas5

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/'</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.</pre>
## MAS5 detection call
call.mas5 <- mas5.call(data.test3,"tmp_Test3Call",tmpdir="",verbose=FALSE)</pre>
## get data.frames
pval.mas5 <- pvalData(call.mas5)</pre>
pres.mas5 <- presCall(call.mas5)</pre>
head(pval.mas5)
head(pres.mas5)
## plot results
if (interactive()) {
callplot(call.mas5, beside=FALSE, ylim=c(0,125))
}
rm(scheme.test3, data.test3)
gc()
```

```
mas5
```

MAS 5.0 Expression Measure

Description

This function converts a DataTreeSet into an ExprTreeSet using the XPS implementation of Affymetrix's MAS 5.0 expression measure.

Usage

```
mas5(xps.data,
filename = character(0),
filedir = getwd(),
tmpdir = "",
normalize = FALSE,
sc = 500,
option = "transcript",
exonlevel = "",
update = FALSE,
xps.scheme = NULL,
add.data = TRUE,
verbose = TRUE)
```

```
xpsMAS5(object, ...)
```

Arguments

| xps.data | object of class DataTreeSet. |
|----------|------------------------------|
| filename | file name of ROOT data file. |

| filedir | system directory where ROOT data file should be stored. |
|------------|--|
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| normalize | logical. If TRUE scale normalization is used after an ExprTreeSet is obtained. |
| SC | value at which all arrays will be scaled to. |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| update | logical. If TRUE the existing ROOT data file filename will be updated. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | arguments filename, filedir, tmpdir, option, exonlevel, xps. scheme. |

Details

This function computes the Affymetrix MAS 5.0 expression measure as implemented in XPS. Although this implementation is based on the Affymetrix 'sadd_whitepaper.pdf', it can be used to compute an expression level for both expression arrays and exon arrays. For exon arrays it is necessary to supply the requested option and exonlevel.

Following options are valid for exon arrays:

transcript:expression levels are computed for transcript clusters, i.e. probe sets containing the same 'transcript_clu
expression levels are computed for exon clusters, i.e. probe sets containing the same 'exon_id', where e
probeset:probeset:expression levels are computed for individual probe sets, i.e. for each 'probeset_id'.

Following exonlevel annotations are valid for exon arrays:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |
| metafull: | full meta-probesets. |
| ambiguous: | ambiguous probesets only. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Following exonlevel annotations are valid for whole genome arrays:

| core: | probesets with category 'unique', 'similar' and 'mixed'. |
|-----------|--|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

mas5

Exon levels can also be combined, with following combinations being most useful:

```
exonlevel="metacore+affx": core meta-probesets plus AFFX controls
exonlevel="core+extended": probesets with cDNA support
exonlevel="core+extended+full": supported plus predicted probesets
```

Exon level annotations are described in the Affymetrix whitepaper 'exon_probeset_trans_clust_whitepaper.pdf'.

If normalize=TRUE then the expression levels will be scaled to sc. For sc=0 the expression levels will be scaled to the mean expression level.

If update=TRUE then the existing ROOT file filename will be updated, however, this is usually only recommended as option for function express.

In order to use an alternative SchemeTreeSet set the corresponding SchemeTreeSet xps.scheme.

xpsMAS5 is the DataTreeSet method called by function mas5, however, expression levels will not be scaled to a common mean expression level.

Value

An ExprTreeSet

Note

In contrast to function mas5, expression levels computed with xpsMAS5 will not be scaled to a common mean expression level.

Author(s)

Christian Stratowa

References

Affymetrix (2002) Statistical Algorithms Description Document, Affymetrix Inc., Santa Clara, CA, whitepaper.http://www.affymetrix.com/support/technical/whitepapers/sadd_whitepaper.pdf

Affymetrix (2005) Exon Probeset Annotations and Transcript Cluster Groupings, Affymetrix Inc., Santa Clara, CA, exon_probeset_trans_clust_whitepaper.pdf.

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
data.mas5 <- mas5(data.test3,"tmp_Test3MAS5",tmpdir="",normalize=TRUE,sc=500,update=TRUE,
## get data.frame
expr.mas5 <- validData(data.mas5)
head(expr.mas5)</pre>
```

```
## plot results
if (interactive()) {
boxplot(data.mas5)
boxplot(log2(expr.mas5))
}
rm(scheme.test3, data.test3)
gc()
```

mboxplot-methods Box Plots of Relative M Values

Description

Produce boxplots of relative M values for the set of arrays.

Usage

```
mboxplot(x, which = "", size = 0, transfo = \log_2, method = "mean", range = 0, ylim = c(-1, 1), outline = FALSE, names = "namepart", ...)
```

Arguments

| Х | object of class DataTreeSet or ExprTreeSet. |
|---------|---|
| which | type of probes to be used, for details see validData. |
| size | length of sequence to be generated as subset. |
| transfo | a valid function to transform the data, usually "log2", or "0". |
| method | method to create the reference data, "mean" or "median". |
| range | determines how far the plot whiskers extend out from the box. |
| ylim | range for the plotted y values. |
| outline | if outline is not true, the outliers are not drawn. |
| names | optional vector of sample names. |
| | optional arguments to be passed to boxplot. |

Details

Create boxplots of M plots, where M is determined relative to a pseudo-mean reference chip.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as boxplot.

Note

For a DataTreeSet object, data must first be attached using method attachInten.

Author(s)

Christian Stratowa

metaProbesets

See Also

boxplot.dev, boxplot

Examples

```
# load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
# need to attach scheme mask and probe intensities
data.test3 <- attachMask(data.test3)
data.test3 <- attachInten(data.test3)
if (interactive()) {
mboxplot(data.test3, ylim=c(-6,6))
}
# optionally remove mask and data to free memory
data.test3 <- removeInten(data.test3)
data.test3 <- removeMask(data.test3)</pre>
```

metaProbesets Create MetaProbeset File for APT

Description

Create MetaProbeset File for APT function "apt-probeset-summarize".

Usage

```
metaProbesets(xps.scheme, infile = character(0), outfile = character(0), exonlev
```

Arguments

| xps.scheme | exon SchemeTreeSet. |
|------------|--|
| infile | Name of file containing exon transcript_cluster_ids. |
| outfile | Name of resulting file containing meta probeset definitions. |
| exonlevel | exon annotation level determining which probes should be used. |

Details

This function allows to create a metaprobeset file for APT function "apt-probeset-summarize" to be used with option "-m". The infile must contain exon transcript_cluster_ids, one per line, e.g. one can export the rownames (data.rma)

The resulting file may be useful if you want to compare results created with xps to results created with APT function "apt-probeset-summarize".

Value

None.

Author(s)

Christian Stratowa

Examples

```
## Not run:
## first, load ROOT exon scheme file:
scmdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Schemes"
scheme.exon <- root.scheme(paste(scmdir,"Scheme_HuEx10stv2r2_na25.root",sep="/"))
metaProbesets(scheme.exon,"metacore.txt","metacoreList.mps","metacore")
## End(Not run)
```

mvaplot.dev MvA Scatter Plot for Device

Description

Produce scatter plots of M values vs A values of the samples for the selected device.

Usage

mvaplot.dev(x, transfo = log2, method = "median", names = "namepart", ylim = c(-

Arguments

| Х | object of class ExprTreeSet. |
|---------|--|
| transfo | a valid function to transform the data, usually "log2", or "0". |
| method | method to compute M, "mean" or "median". |
| names | optional vector of sample names. |
| ylim | range for the plotted M values. |
| xlab | a title for the x axis. |
| ylab | a title for the y axis. |
| pch | either an integer specifying a symbol or a single character to be used in plotting points. |
| | 1 |
| mar | plot margin. |
| dev | graphics device to plot to, i.e. one of "screen", "jpeg", "png", "pdf" or "ps". |
| outfile | the name of the output file. |
| W | the width of the device in pixels. |
| h | the height of the device in pixels. |
| | optional arguments to be passed to plot. |

Details

Produces mvaplots for slot data for an object of class ExprTreeSet for the selected graphics device.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as myaplot.

mvaplot-methods

Author(s)

Christian Stratowa

See Also

mvaplot

mvaplot-methods M vs A Plot

Description

Produce scatter plots of M values vs A values of the samples.

Usage

mvaplot(x, transfo = log2, method = "median", names = "namepart", ylim = c(-6,6), ...)

Arguments

| х | object of class ExprTreeSet. |
|---------|---|
| transfo | a valid function to transform the data, usually "log2", or "0". |
| method | method to compute M, "mean" or "median". |
| names | optional vector of sample names. |
| ylim | range for the plotted M values. |
| | optional arguments to be passed to plot. |

Details

Produces myaplots for slot data for an object of class ExprTreeSet.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as myaplot.

Author(s)

Christian Stratowa

See Also

mvaplot.dev

namePart

Description

Get (tree) names w/o their extension.

Usage

```
namePart(names)
```

Arguments

names vector of names.

Details

Extracts the name part of names, e.g.of tree names of treename.treetype stored in a ROOT file.

Value

A vector of tree names w/o its extension.

Author(s)

Christian Stratowa

See Also

extenPart

Examples

```
names <- c("TestA1.int", "TestA2.int")
namePart(names)</pre>
```

| normalize | Normalization on Affymetrix Probe Level Data or on Expression Lev- |
|-----------|--|
| | els |

Description

Functions that allow to normalize Affymetrix arrays both at the probe level ("low-level normalization") and/or at the expression level ("high-level normalization").

normalize

Usage

```
normalize(xps.data, filename = character(0), filedir = getwd(), tmpdir = "", upd
normalize.constant(xps.data, filename = character(0), filedir = getwd(), tmpdir
normalize.lowess(xps.data, filename = character(0), filedir = getwd(), tmpdir =
normalize.quantiles(xps.data, filename = character(0), filedir = getwd(), tmpdir
normalize.supsmu(xps.data, filename = character(0), filedir = getwd(), tmpdir
xpsNormalize(object, ...)
```

Arguments

| xps.data | object of class DataTreeSet or ExprTreeSet. |
|-----------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| update | logical. If TRUE the existing ROOT data file filename will be updated. |
| select | type of probes to select for normalization. |
| method | normalization method to use. |
| option | option determining the grouping of probes for normalization, and the selection of the probes. |
| logbase | logarithm base as character, one of '0', 'log', 'log2', 'log10'. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| refindex | index of reference tree to use, or 0. |
| refmethod | for refindex=0, either trimmed mean or median of trees. |
| params | vector of parameters for normalization method. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet or ExprTreeSet. |
| | the arguments described above. |

Details

Functions that allow to normalize Affymetrix arrays both at the probe level ("low-level normalization") and/or at the expression level ("high-level normalization").

Please have a look at vignette "xpsPreprocess.pdf" for details on how to use function normalize.

xpsNormalize are the DataTreeSet or ExprTreeSet methods, respectively, called by function normalize, containing the same parameters.

Value

An object of type DataTreeSet or ExprTreeSet.

pm-methods

Warning

Functions normalize.lowess and normalize.supsmu have only be tested for objects of type ExprTreeSet but not for objects of type DataTreeSet, i.e. for probe level intensities.

Author(s)

Christian Stratowa

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
## RMA background
data.bg.rma <- bgcorrect.rma(data.test3, "tmp_Test3NormRMA", filedir=getwd(), tmpdir="", verb
## normalize quantiles
data.qu.rma <- normalize.quantiles(data.bg.rma, "tmp_Test3NormRMA", filedir=getwd(), tmpdir=
## summarize medianpolish
data.mp.rma <- summarize.rma(data.qu.rma, "tmp_Test3NormRMA", filedir=getwd(), tmpdir="", upd
</pre>
```

pm-methods *Methods for accessing perfect matches and mismatches*

Description

Methods for accessing perfect match (PM) and mismatch (MM) probes.

Usage

pm(object, which = "pm")
mm(object, which = "mm")

Arguments

| object | object of class DataTreeSet. |
|--------|--|
| which | type of perfect match or mismatch probes to be returned. |

Details

For expression arrays all the perfect match (pm) or mismatch (mm) probes on the arrays the object represents are returned as data.frame.

For exon arrays, pm returns the probes of the different exon levels as data.frame, i.e. which can have one of the following values:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |

pmplot-methods

| probesets supported by gene predictions only. |
|---|
| full meta-probesets. |
| standard AFFX controls. |
| |

For whole genome arrays, pm returns the probes of the different exon levels as data.frame, i.e. which can have one of the following values:

| core: | probesets with category 'unique' and 'mixed'. |
|-----------|---|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |

For exon/genome arrays, mm returns the background probes as data.frame, i.e. which is either "genomic" or "antigenomic".

Value

A data.frame.

Author(s)

Christian Stratowa

See Also

validData

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
## need to attach scheme mask and probe intensities
data.test3 <- attachMask(data.test3)
data.test3 <- attachInten(data.test3)
pm <- pm(data.test3)
nm <- mm(data.test3)
head(pm)
head(mm)
## optionally remove mask and data to free memory
data.test3 <- removeInten(data.test3)
data.test3 <- removeMask(data.test3)</pre>
```

pmplot-methods Barplot of PM and MM Intensities.

Description

Creates a barplot of mean perfect match and mismatch intensities.

Usage

```
pmplot(x, which = "", size = 0, transfo = NULL, method = "mean", names
= "namepart", beside = TRUE, col = c("red", "blue"), legend = c("PM", "MM"),
...)
```

Arguments

| Х | object of class DataTreeSet. |
|---------|--|
| which | type of probes to be used, for details see validData. |
| size | length of sequence to be generated as subset. |
| transfo | a valid function to transform the data, usually "log2", or "0". |
| method | method to compute average intensities, "mean" or "median". |
| names | optional vector of sample names. |
| beside | logical. If FALSE, mean intensities are portrayed as stacked bars, and if TRUE the columns are portrayed as juxtaposed bars. |
| col | color of PM, MM bars. |
| legend | a vector of text used to construct a legend for the plot, or a logical indicating whether a legend should be included. |
| | optional arguments to be passed to barplot. |

Details

Produces barplots of mean perfect match and mismatch intensities for slot data for an object of class ExprTreeSet.

For names=NULL full column names of slot data will be displayed while for names="namepart" column names will be displayed without name extension. If names is a vector of column names, only these columns will displayed as pmplot.

Note

Data must first be attached to class DataTreeSet using method attachInten.

Author(s)

Christian Stratowa

See Also

boxplot.dev, boxplot, barplot

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
## need to attach scheme mask and probe intensities
data.test3 <- attachMask(data.test3)</pre>
```

PreFilter-class

```
data.test3 <- attachInten(data.test3)
if (interactive()) {
pmplot(data.test3)
}
## optionally remove mask and data to free memory
data.test3 <- removeInten(data.test3)
data.test3 <- removeMask(data.test3)</pre>
```

PreFilter-class Class PreFilter

Description

Class PreFilter allows to apply different filters to class ExprTreeSet, i.e. to the expression level data.frame data.

Objects from the Class

Objects can be created by calls of the form new("PreFilter", ...). Alternatively, the contructor PreFilter can be used.

Slots

```
mad: Object of class "list" describing parameters for madFilter.
cv: Object of class "list" describing parameters for cvFilter.
variance: Object of class "list" describing parameters for varFilter.
difference: Object of class "list" describing parameters for diffFilter.
ratio: Object of class "list" describing parameters for ratioFilter.
gap: Object of class "list" describing parameters for gapFilter.
hithreshold: Object of class "list" describing parameters for highFilter.
lothreshold: Object of class "list" describing parameters for lowFilter.
quantile: Object of class "list" describing parameters for quantileFilter.
numfilters: Object of class "list" describing parameters for quantileFilter.
```

Extends

Class "Filter", directly.

Methods

```
callFilter signature(object = "PreFilter"): extracts slot prescall.
```

callFilter<- signature(object = "PreFilter", value = "character"): replaces
 slot prescall with character vector c(cutoff, samples, condition).</pre>

cvFilter signature(object = "PreFilter"): extracts slot cv.

diffFilter signature(object = "PreFilter"): extracts slot difference.

- diffFilter<- signature(object = "PreFilter", value = "numeric"): replaces
 slot difference with numeric vector c(cutoff, trim, epsilon).</pre>
- gapFilter signature(object = "PreFilter"): extracts slot gap.
- gapFilter<- signature(object = "PreFilter", value = "numeric"): replaces
 slot gap with numeric vector c(cutoff, window, trim, epsilon).</pre>
- **highFilter** signature (object = "PreFilter"): extracts slot hithreshold.
- highFilter<- signature(object = "PreFilter", value = "character"): replaces
 slot hithreshold with character vector c(cutoff, parameter, condition).</pre>
- **lowFilter** signature(object = "PreFilter"): extracts slot lothreshold.
- lowFilter<- signature(object = "PreFilter", value = "character"): replaces
 slot lothreshold with character vector c(cutoff, parameter, condition).</pre>
- madFilter signature(object = "PreFilter"): extracts slot mad.
- madFilter<- signature(object = "PreFilter", value = "numeric"): replaces
 slot mad with numeric vector c(cutoff, epsilon).</pre>
- quantileFilter signature(object = "PreFilter"): extracts slot quantile.
- quantileFilter<- signature(object = "PreFilter", value = "numeric"): replaces
 slot quantile with numeric vector c(cutoff, loquantile, hiquantile).</pre>
- ratioFilter signature(object = "PreFilter"): extracts slot ratio.
- ratioFilter<- signature(object = "PreFilter", value = "numeric"):replaces
 slot ratio with numeric vector c(cutoff).</pre>
- varFilter signature(object = "PreFilter"): extracts slot variance.
- varFilter<- signature(object = "PreFilter", value = "numeric"): replaces slot variance with numeric vector c(cutoff, trim, epsilon).

Author(s)

Christian Stratowa

See Also

related classes Filter, UniFilter.

Examples

```
## for demonstration purposes only: initialize all pre-filters
prefltr <- new("PreFilter")
madFilter(prefltr) <- c(0.5,0.01)
cvFilter(prefltr) <- c(0.3,0.0,0.01)
varFilter(prefltr) <- c(0.6,0.02,0.01)
diffFilter(prefltr) <- c(2.2,0.0,0.01)
ratioFilter(prefltr) <- c(1.5)
gapFilter(prefltr) <- c(0.3,0.05,0.0,0.01)
lowFilter(prefltr) <- c(4.0,3,"samples")
highFilter(prefltr) <- c(14.5,75.0,"percent")
quantileFilter(prefltr) <- c(0.02,80.0,"percent")
str(prefltr)
```

PreFilter-constructor

Constructor for Class PreFilter

Description

Constructor for class PreFilter allows to apply different filters to class ExprTreeSet, i.e. to the expression level data.frame data.

Usage

| PreFilter(mad | | = | character(), |
|---------------|-------------|---|--------------|
| (| CV | = | character(), |
| , | variance | = | character(), |
| (| difference | = | character(), |
| : | ratio | = | character(), |
| (| gap | = | character(), |
| - | lothreshold | = | character(), |
|] | hithreshold | = | character(), |
| (| quantile | = | character(), |
|] | prescall | = | character()) |

Arguments

| mad | "character" vector describing parameters for madFilter. |
|-------------|--|
| CV | "character" vector describing parameters for cvFilter. |
| variance | "character" vector describing parameters for varFilter. |
| difference | "character" vector describing parameters for diffFilter. |
| ratio | "character" vector describing parameters for ratioFilter. |
| gap | "character" vector describing parameters for gapFilter. |
| lothreshold | "character" vector describing parameters for lowFilter. |
| hithreshold | "character" vector describing parameters for highFilter. |
| quantile | "character" vector describing parameters for quantileFilter. |
| prescall | "character" vector describing parameters for callFilter. |

Details

The PreFilter constructor allows to apply the following filters to class ExpTreeSet:

| character vector c(cutoff,epsilon). |
|---|
| character vector c(cutoff,trim,epsilon). |
| character vector c(cutoff,trim,epsilon). |
| character vector c(cutoff,trim,epsilon). |
| character vector c(cutoff). |
| character vector c(cutoff,window,trim,epsilon). |
| character vector c(cutoff,parameter,condition). |
| character vector c(cutoff,parameter,condition). |
| character vector c(cutoff,loquantile,hiquantile). |
| character vector c(cutoff,samples,condition). |
| |

prefilter

Value

An object of type "PreFilter"

Note

Function PreFilter is used as constructor for class PreFilter so that the user need not know details for creating S4 classes.

Author(s)

Christian Stratowa

See Also

Filter, UniFilter

Examples

```
str(prefltr)
```

```
## alternatively add character vectors as methods after creation of constructor
prefltr <- PreFilter()
madFilter(prefltr) <- c(0.5,0.01)
gapFilter(prefltr) <- c(0.3,0.05,0.0,0.01)
lowFilter(prefltr) <- c(4.0,3,"samples")
highFilter(prefltr) <- c(14.5,75.0,"percent")
str(prefltr)</pre>
```

```
prefilter
```

Function for Applying a PreFilter to an ExprTreeSet

Description

This function applies a PreFilter to an ExprTreeSet.

Usage

```
prefilter(xps.expr,
    filename = character(0),
    filedir = getwd(),
    filter = NULL,
    minfilters = 999,
    logbase = "log2",
    treename = "PreFilter",
    xps.call = NULL,
    verbose = TRUE)
```

xpsPreFilter(object, ...)

```
90
```

prefilter

Arguments

| xps.expr | object of class ExprTreeSet. |
|------------|---|
| filename | file name of ROOT filter file. |
| filedir | system directory where ROOT filter file should be stored. |
| filter | object of class PreFilter. |
| minfilters | minimum number of initialized filter methods to satisfy (default is all filters). |
| logbase | convert data to logarithm of base: "0", "log", "log2" (default), "log10" |
| treename | tree name to be used in ROOT filter file. |
| xps.call | optional object of class CallTreeSet. |
| verbose | logical, if TRUE print status information. |
| object | object of class ExprTreeSet. |
| | same arguments as function prefilter. |

Details

This function applies the different filters initialized with constructor PreFilter to the ExprTreeSet xps.expr.

Slot minfilters determines the minimum number of initialized filters, which must be satisfied so that the mask is set to flag=1. For minfilters=1 at least one filter must be satisfied, equivalent to logical 'OR'; for minfilters=999 all filters must be satisfied, equivalent to logical 'AND'.

If method callFilter was initialized with constructor PreFilter then CallTreeSet xps.call must be supplied, usually created with function mas5.call.

Value

A FilterTreeSet

Author(s)

Christian Stratowa

See Also

PreFilter, unifilter

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
## second, create an ExprTreeSet
data.rma <- rma(data.test3,"tmp_TestRMA",tmpdir="",background="pmonly",normalize=TRUE,ver
## note: do not copy/paste this code, it is necessary only because R CMD check fails sind
data.rma@rootfile <- paste(.path.package("xps"),"rootdata/tmp_Test3RMA.root",sep="/")
data.rma@filedir <- paste(.path.package("xps"),"rootdata",sep="/")</pre>
```

```
## third, construct a PreFilter
prefltr <- PreFilter(mad=c(0.5,0.01),lothreshold=c(6.0,0.02,"mean"),hithreshold=c(10.5,80)</pre>
```

presCall-methods

```
## finally, create a FilterTreeSet
rma.pfr <- prefilter(data.rma,"tmp_Test3Prefilter",getwd(),prefltr,2,verbose=FALSE)
str(rma.pfr)
## End(Not run)</pre>
```

presCall-methods Get/Set Present Call Values

Description

Get/set present call values from/for class CallTreeSet.

```
Usage
presCall(object)
presCall(object, treenames = NULL) <- value
pvalData(object)
pvalData(object, treenames = NULL) <- value</pre>
```

Arguments

| object | object of class CallTreeSet. |
|-----------|---|
| treenames | character vector containing optional tree names to be used as subset. |
| value | data.frame containing present call values. |

Details

Get the p-values from slot data or present calls from slot detcall, or set slot data or detcall, respectively, to value.

Method presCall returns the present calls from slot detcall as data.frame, while replacement method presCall<- allows to replace slot detcall with a data.frame.

Method pvalData returns the p-values from slot data as data.frame, while replacement method pvalData<- allows to replace slot data with a data.frame.

In order to create an CallTreeSet containing only a subset of e.g. slot data, first export slot data using method pvalData, create a character vector containing only treenames to be used in the subset, and then use replacement method pvalData<- to replace slot data with the subset. Slots treenames and numtrees will be updated automatically for pvalData<- but not for presCall<-.

Note: When creating character vector treenames it is sufficient to use the name part of the tree name w/o the extension.

Note: If you do not want to replace your current object, create first a copy of type CallTreeSet by simply writing newobj <- oldobj, and use newobj for replacement.

Author(s)

Christian Stratowa

ProcesSet-class

See Also

exprs

Examples

```
## Not run:
## load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"), "schemes/SchemeTest3.root", sep="/'</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"), "rootdata/DataTest3_cel.</pre>
## create an CallTreeSet
call.mas5 <- mas5.call(data.test3,"tmp_TestMAS5Call",tmpdir="",verbose=FALSE)</pre>
## get p-values
value <- pvalData(call.mas5)</pre>
## selected treenames only
treenames <- c("TestA2", "TestB1")</pre>
## make a copy of your object if you do not want to replace it
subset.call <- call.mas5</pre>
## replace slot data with subset
exprs(subset.call, treenames) <- value</pre>
str(subset.call)
## End(Not run)
```

ProcesSet-class Class ProcesSet

Description

This class provides access to class SchemeTreeSet for the derived classes DataTreeSet, ExprTreeSet and CallTreeSet. It extends class TreeSet.

Objects from the Class

Usually, no objects are created from it.

Slots

scheme: Object of class "SchemeTreeSet" providing access to ROOT scheme file.

- data: Object of class "data.frame". The data.frame can contain the data stored in ROOT data trees.
- params: Object of class "list" representing relevant parameters.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT trees are stored, usually one of 'DataTreeSet', 'PreprocesSet', 'CallTreeSet'.
- settype: Object of class "character" describing the type of treeset stored in setname, usually one of 'rawdata', 'preprocess'.
- rootfile: Object of class "character" representing the name of the ROOT file, including full path.

- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

Extends

Class "TreeSet", directly.

Methods

- **chipName** signature(object = "ProcesSet"): **extracts** slot chipname from slot scheme.
- **chipType** signature(object = "ProcesSet"): **extracts slot** chiptype from **slot** scheme.
- export signature(object = "ProcesSet"): exports ROOT trees as text file, see exportmethods.
- getTreeData signature(object = "ProcesSet"): exports tree data from ROOT file rootfile, and saves as data.frame data.
- hist signature(x = "ProcesSet"): creates a plot showing the histograms for data.frame
 data.
- schemeFile signature(object = "ProcesSet"): extracts the ROOT scheme file from
 slot scheme.
- schemeFile<- signature(object = "ProcesSet"), value = "character"):replaces the ROOT scheme file from slot scheme.</pre>
- schemeSet signature(object = "ProcesSet"): extracts slot scheme.
- schemeSet<- signature(object = "ProcesSet"), value = "SchemeTreeSet"):
 replaces slot scheme with a different SchemeTreeSet.</pre>

Author(s)

Christian Stratowa

See Also

derived classes DataTreeSet, ExprTreeSet, CallTreeSet.

Examples

showClass("ProcesSet")

ProjectInfo-class Class ProjectInfo

Description

This class allows to save the relevant project information in the ROOT data file and in class DataTreeSet.

Objects from the Class

```
Objects can be created by calls of the form
```

```
new("ProjectInfo", submitter=[character], laboratory=[character], contact=[chara
...).
```

Alternatively, the constructor ProjectInfo can be used.

Slots

submitter: Object of class "character" representing the name of the submitter.

laboratory: Object of class "character" representing the laboratory of the submitter.

contact: Object of class "character" representing the contact address of the submitter.

project: Object of class "list" representing the project information.

author: Object of class "list" representing the author information.

dataset: Object of class "list" representing the dataset information.

source: Object of class "list" representing the sample source information.

sample: Object of class "list" representing the sample information.

celline: Object of class "list" representing the sample information for cell lines.

primarycell: Object of class "list" representing the sample information for primary cells.

tissue: Object of class "list" representing the sample information for tissues.

biopsy: Object of class "list" representing the sample information for biopsies.

arraytype: Object of class "list" representing the array information.

- hybridizations: Object of class "data.frame" representing the hybridization information for each hybridization.
- treatments: Object of class "data.frame" representing the treatment information for each hybridization.

Methods

projectInfo signature(object = "ProjectInfo"): extracts slot project.

- projectInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot project with character vector c(name,date,type,description,comments).
- authorInfo signature(object = "ProjectInfo"): extracts slot author.

authorInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot author with character vector c(lastname,firstname,type,company,department,email, phone,comments).

datasetInfo signature(object = "ProjectInfo"): extracts slot dataset.

datasetInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot dataset with character vector c(name,type,sample,submitter,date,description,comments).</pre>

sourceInfo signature(object = "ProjectInfo"): extracts slot source.

- sourceInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot source with character vector c(name,type,species,subspecies,description,comments).</pre>
- sampleInfo signature(object = "ProjectInfo"): extracts slot sample.
- sampleInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot sample with character vector c(name,type,sex,phenotype,genotype,extraction, isxenograft,xenostrain,xenosex,xenoage,xenoageunit,comments).
- cellineInfo signature(object = "ProjectInfo"): extracts slot celline.
- **cellineInfo<-** signature(object = "ProjectInfo", value = "character"): replaces slot celline with character vector c(name,type,parent,atcc,modification,sex,phenotype, genotype,extraction,isxenograft,xenostrain,xenosex,xenoage,xenoageunit,comments).
- primcellInfo signature(object = "ProjectInfo"): extracts slot primarycell.
- primcellInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot primarycell with character vector c(name,type,date,description,sex,phenotype, genotype,extraction,isxenograft,xenostrain,xenosex,xenoage,xenoageunit,comments).
- tissueInfo signature(object = "ProjectInfo"): extracts slot tissue.
- biopsyInfo signature(object = "ProjectInfo"): extracts slot biopsy.
- arrayInfo signature(object = "ProjectInfo"): extracts slot arraytype.
- arrayInfo<- signature(object = "ProjectInfo", value = "character"):replaces slot arraytype with character vector c(chipname,chiptype,description,comments).
- **hybridizInfo** signature(object = "ProjectInfo"): extracts slot hybridizations.
- hybridizInfo<- signature(object = "ProjectInfo", value = "character"): replaces slot hybridizations with vector of character vectors with each containing c(name,type,inputname,date,
- treatmentInfo signature(object = "ProjectInfo"): extracts slot treatments.
- treatmentInfo<- signature(object = "ProjectInfo", value = "character"):
 replaces slottreatments with vector of character vectors with each containing c(name,type,concentration,concentration)</pre>
- **show** signature (object = "ProjectInfo"): shows the content of ProjectInfo.

Author(s)

Christian Stratowa

Examples

```
project <- new("ProjectInfo", submitter="Christian", laboratory="home", contact="email")
projectInfo(project) <- c("TestProject", "20060106", "Project Type", "use Test3 data for
authorInfo(project) <- c("Test3Set", "MC", "Tissue", "Stratowa", "20060106", "description"
sourceInfo(project) <- c("Unknown", "source type", "Homo sapiens", "caucasian", "descript
primcellInfo(project) <- c("Mel31", "primary cell", 20071123, "extracted from patient", "ma
arrayInfo(project) <- c(c("TestA1", "hyb type", "TestA1.CEL", 20071117, "my prep1", "star</pre>
```

ProjectInfo-constructor

```
c ("TestA2", "hyb type", "TestA2.CEL", 20071117, "my prep2", "star
c ("TestB1", "hyb type", "TestB1.CEL", 20071117, "my prep1", "star
c ("TestB2", "hyb type", "TestB2.CEL", 20071117, "my prep2", "star
treatmentInfo(project) <- c(c("TestA1", "DMSO", 4.3, "mM", 1.0, "hours", "intravenous", "my con
c ("TestA2", "DMSO", 4.3, "mM", 8.0, "hours", "intravenous", "my con
c ("TestB1", "DrugA2", 4.3, "mM", 1.0, "hours", "intravenous", "my con
c ("TestB2", "DrugA2", 4.3, "mM", 8.0, "hours", "intravenous", "my con
show(project)
```

ProjectInfo-constructor Constructor for Class ProjectInfo

Description

Constructor for class ProjectInfo class allows to save the relevant project information in the ROOT data file and in class DataTreeSet.

Usage

```
ProjectInfo(submitter = character(),
    laboratory = character(),
    contact = character(),
    project = character(),
    author = character(),
    dataset = character(),
    source = character(),
    source = character(),
    sample = character(),
    celline = character(),
    primarycell = character(),
    tissue = character(),
    biopsy = character(),
    arraytype = character(),
    hybridizations = character(),
    treatments = character())
```

Arguments

| submitter | "character" representing the name of the submitter. |
|-------------|---|
| laboratory | "character" representing the laboratory of the submitter. |
| contact | "character" representing the contact address of the submitter. |
| project | "character" vector representing the project information. |
| author | "character" vector representing the author information. |
| dataset | "character" vector representing the dataset information. |
| source | "character" vector representing the sample source information. |
| sample | "character" vector representing the sample information. |
| celline | "character" vector representing the sample information for cell lines. |
| primarycell | "character" vector representing the sample information for primary cells. |
| tissue | "character" vector representing the sample information for tissues. |

| biopsy | "character" vector representing the sample information for biopsies. |
|--------------|--|
| arraytype | "character" vector representing the array information. |
| hybridizatio | ns |
| | "character" vector representing the hybridization information for each hybridization. |
| treatments | "character" vector representing the treatment information for each hybridiza- tion. |

Details

The ProjectInfo constructor allows to save the following project information in the ROOT data file and in class DataTreeSet:

| submitter: | name of the submitter. |
|-----------------|---|
| laboratory: | laboratory of the submitter. |
| contact: | contact address of the submitter. |
| project: | character vector c(name,date,type,description,comments). |
| author: | character vector c(lastname,firstname,type,company,department,email, phone,comments) |
| dataset: | character vector c(name,type,sample,submitter,date,description,comments). |
| source: | character vector c(name,type,species,subspecies,description,comments). |
| sample: | character vector c(name,type,sex,phenotype,genotype,extraction, isxenograft,xenostrain,xenose |
| celline: | character vector c(name,type,parent,atcc,modification,sex,phenotype, genotype,extraction,isxer |
| primarycell: | character vector c(name,type,date,description,sex,phenotype, genotype,extraction,isxenograft,x |
| tissue: | character vector c(name,type,development,morphology,disease,stage, donorage,ageunit,status,s |
| biopsy: | character vector c(name,type,morphology,disease,stage,donorage,ageunit, status,sex,phenotype |
| arraytype: | character vector c(chipname, chiptype, description, comments). |
| hybridizations: | vector of character vectors with each containing c(name,type,inputname,date,preparation,proto |
| treatments: | vector of character vectors with each containing c(name,type,concentration,concentrationunit,ti |

Value

An object of type "ProjectInfo"

Note

Function ProjectInfo is used as constructor for class ProjectInfo so that the user need not know details for creating S4 classes.

Author(s)

Christian Stratowa

See Also

ProjectInfo

Examples

quantileFilter-methods

```
c("TestB1","hyb type","TestB1.CEL",20071117,"my p
c("TestB2","hyb type","TestB2.CEL",20071117,"my p
str(project)
## alternatively add character vectors as methods after creation of constructor
authorInfo(project) <- c("Stratowa","Christian","Project Leader","Company","Dept","cst
datasetInfo(project) <- c("Test3Set","MC","Tissue","Stratowa","20060106","description",
treatmentInfo(project) <- c(c("TestA1","DMSO",4.3,"mM",1.0,"hours","intravenous","my comm
c("TestB1","DrugA2",4.3,"mM",8.0,"hours","intravenous","my com
str(project)
```

quantileFilter-methods

Quantile Filter

Description

This method initializes the Quantile Filter. The Quantile Filter flags all rows with: flag = (quantile[high]/quantile[low] >= cutoff)

Usage

quantileFilter(object)
quantileFilter(object, value)<-</pre>

Arguments

| object | object of class PreFilter. | | |
|--------|-------------------------------------|-------------|--------------|
| value | <pre>numeric vector c(cutoff,</pre> | loquantile, | hiquantile). |

Details

The method quantileFilter initializes the following parameters:

| cutoff: | the cutoff level for the filter. |
|-------------|---|
| loquantile: | value for low quantile (default is loquantile=0.05). |
| hiquantile: | value for high quantile (default is hiquantile=0.95). |

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
quantileFilter(prefltr) <- c(3.0, 0.05, 0.95)
str(prefltr)</pre>
```

ratioFilter-methods

Ratio Filter

Description

This method initializes the Ratio Filter. The ratio is the maximum value divided by minimum value for each row of the expression dataframe.

The Ratio Filter flags all rows with: flag = (max/min >= cutoff)

Usage

```
ratioFilter(object)
ratioFilter(object, value)<-</pre>
```

Arguments

| object | object of class PreFilter. |
|--------|-------------------------------------|
| value | <pre>numeric value c(cutoff).</pre> |

Details

The method ratioFilter initializes the following parameters:

cutoff: the cutoff level for the filter.

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
ratioFilter(prefltr) <- c(1.5)
str(prefltr)</pre>
```

rawCELName-methods Method for getting names of the raw CEL-files

Description

Method for getting names (and full path) of the original CEL-files.

Usage

rawCELName(object, treename = "*", fullpath = TRUE)

rma

Arguments

| object | object of class DataTreeSet. |
|----------|---|
| treename | treename, for which the name of the original CEL-file should be returned. |
| fullpath | logical, if TRUE return full path. |

Details

Since CEL-files can be imported with import.data using alternative celnames, method rawCELName allows to return the original name and optionally the full path for each CEL-file.

Value

A character vector.

Author(s)

Christian Stratowa

See Also

import.data

Examples

```
## load existing ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
rawCELName(data.test3)
rawCELName(data.test3, treename = "TestA2.cel", fullpath = FALSE)
```

rma

Robust Multi-Array Average Expression Measure

Description

This function converts a DataTreeSet into an ExprTreeSet using the robust multi-array average (RMA) expression measure.

Usage

```
rma(xps.data,
    filename = character(0),
    filedir = getwd(),
    tmpdir = "",
    background = "pmonly",
    normalize = TRUE,
    option = "transcript",
    exonlevel = "",
    params = list(16384, 0.0, 1.0, 10, 0.01, 1.0),
    xps.scheme = NULL,
```

```
add.data = TRUE,
verbose = TRUE)
```

.. ..

xpsRMA(object, ...)

Arguments

| xps.data | object of class DataTreeSet. |
|------------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| background | probes used to compute background, one of 'pmonly', 'mmonly', 'both'; for genome/exon arrays one of 'genomic', 'antigenomic' |
| normalize | logical. If TRUE normalize data using quantile normalization. |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| params | list of (default) parameters for rma. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | the arguments described above. |

Details

This function computes the RMA (Robust Multichip Average) expression measure described in Irizarry et al. for both expression arrays and exon arrays. For exon arrays it is necessary to supply the requested option and exonlevel.

Following options are valid for exon arrays:

transcript:expression levels are computed for transcript clusters, i.e. probe sets containing the same 'transcript_clu
expression levels are computed for exon clusters, i.e. probe sets containing the same 'exon_id', where e
probeset:probeset:expression levels are computed for individual probe sets, i.e. for each 'probeset_id'.

Following exonlevel annotations are valid for exon arrays:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |
| metafull: | full meta-probesets. |
| ambiguous: | ambiguous probesets only. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Following exonlevel annotations are valid for whole genome arrays:

| core: | probesets with category 'unique', 'similar' and 'mixed'. |
|-----------|--|
| metacore: | probesets with category 'unique' only. |
| affx: | standard AFFX controls. |
| all: | combination of above (including affx). |

Exon levels can also be combined, with following combinations being most useful:

| exonlevel="metacore+affx": | core meta-probesets plus AFFX controls |
|--|--|
| exonlevel="core+extended": | probesets with cDNA support |
| <pre>exonlevel="core+extended+full":</pre> | supported plus predicted probesets |

Exon level annotations are described in the Affymetrix whitepaper exon_probeset_trans_clust_whitepaper.pdf: "Exon Probeset Annotations and Transcript Cluster Groupings".

In order to use an alternative SchemeTreeSet set the corresponding SchemeSet xps.scheme.

xpsRMA is the DataSet method called by function rma, containing the same parameters.

Value

An ExprTreeSet

Note

In contrary to other implementations of RMA the expression measure is given to you in linear scale, analogously to the expression measures computed with mas5 and mas4.

Please note that the default settings of params gives results which are identical to the results obtained with APT (Affymetrix Power Tools) and with package $affy_1.14.2$ or earlier. If you want to obtain results which are identical to the results obtained with $affy_1.16.0$ or later then you need to set params = list (16384, 0.0, 0.4, 10, 0.01, 1.0).

By setting parameter background="none" it is possible to skip background correction .

For the analysis of many exon arrays it may be better to define a tmpdir, since this will store only the results in the main file and not e.g. background and normalized intensities, and thus will reduce the file size of the main file. For quantile normalization memory should not be an issue, however medianpolish depends on RAM unless you are using a temporary file.

Parameter exonlevel determines not only which probes are used for medianpolish, but also the probes used for background calculation and for quantile normalization. If you want to use seperate probes for background calculation, quantile normalization and medianpolish summarization, you can pass a numeric vector containing three integer values corresponding to the respective exonlevel, e.g. you can use exonlevel=c(16316, 8252, 8252), see function exonLevel for more details.

Author(s)

Christian Stratowa

References

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Rafael. A. Irizarry, Benjamin M. Bolstad, Francois Collin, Leslie M. Cope, Bridget Hobbs and Terence P. Speed (2003), Summaries of Affymetrix GeneChip probe level data Nucleic Acids Research 31(4):e15

Bolstad, B.M., Irizarry R. A., Astrand M., and Speed, T.P. (2003), A Comparison of Normalization Methods for High Density Oligonucleotide Array Data Based on Bias and Variance. Bioinformatics 19(2):185-193

Irizarry, RA, Hobbs, B, Collin, F, Beazer-Barclay, YD, Antonellis, KJ, Scherf, U, Speed, TP (2003) Exploration, Normalization, and Summaries of High Density Oligonucleotide Array Probe Level Data. Biostatistics .Vol. 4, Number 2: 249-264

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root", sep="/"</pre>
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
data.rma <- rma(data.test3,"tmp_Test3RMA",tmpdir="",background="pmonly",normalize=TRUE,ve</pre>
## get data.frame
expr.rma <- validData(data.rma)</pre>
head(expr.rma)
## plot results
if (interactive()) {
boxplot(data.rma)
boxplot(log2(expr.rma))
}
rm(scheme.test3, data.test3)
gc()
## Not run:
## examples using Affymetrix human tissue dataset (see also xps/examples/script4exon.R)
## first, load ROOT scheme file and ROOT data file from e.g.:
scmdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Schemes"</pre>
datdir <- "/Volumes/GigaDrive/CRAN/Workspaces/ROOTData"</pre>
## 1. example - expression array, e.g. HG-U133_Plus_2:
scheme.u133p2 <- root.scheme(paste(scmdir,"Scheme_HGU133p2_na25.root",sep="/"))</pre>
              <- root.data(scheme.u133p2, paste(datdir,"HuTissuesU133P2_cel.root",sep="/"
data.u133p2
workdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Exon/hutissues/u133p2"
data.rma <- rma(data.u133p2,"MixU133P2RMA",filedir=workdir,tmpdir="",</pre>
                background="pmonly", normalize=TRUE)
## 2. example - whole genome array, e.g. HuGene-1_0-st-v1:
scheme.genome <- root.scheme(paste(scmdir,"Scheme_HuGene10stv1r3_na25.root",sep="/"))</pre>
              <- root.data(scheme.genome, paste(datdir,"HuTissuesGenome_cel.root",sep="/'
data.genome
workdir <- "/Volumes/GigaDrive/CRAN/Workspaces/Exon/hutissues/hugene"
```

root.browser-methods

root.browser-methods

Open the ROOT object browser

Description

Open the ROOT object browser to see all objects stored in a ROOT file including ROOT trees.

Usage

```
root.browser(object)
```

Arguments

object an object of type SchemeTreeSet, DataTreeSet, ExprTreeSet, or CallTreeSet

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT browser, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

root.call

Create class CallTreeSet accessing ROOT detection call file

Description

Create class CallTreeSet accessing ROOT detection call file.

Usage

```
root.call(xps.scheme, rootfile = character(0), treetype = character(0), treename
```

Arguments

| xps.scheme | A SchemeTreeSet containing the correct scheme for the ROOT data file. |
|------------|---|
| rootfile | name of ROOT data file, including full path. |
| treetype | tree type. |
| treenames | optional character vector of tree names to get only subset of trees. |

Details

An S4 class CallTreeSet will be created, serving as R wrapper to the existing ROOT detection call file rootfile.

Parameter treetype must be supplied to identify the ROOT trees for slots data and detcall. Valid tree types are listed in validTreetype.

To get the names of all trees with their extensions treetype, which are stored in rootfile, you can call function getTreeNames first.

If the CallTreeSet should only handle a subset of the trees stored in rootfile, the tree names must be supplied as vector treenames.

Value

A CallTreeSet object.

Author(s)

Christian Stratowa

See Also

root.data,root.expr

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
## MAS5 detection call
detcall.mas5 <- mas5.call(data.test3,"tmp_Test3CallAll",tmpdir="",verbose=FALSE)
## use subset of trees
sub.call <- root.call(scheme.test3, "tmp_Test3CallAll.root", "dc5", c("TestA2", "TestB1")</pre>
```

```
root.data
```

Create class DataTreeSet accessing ROOT data file

Description

Create class DataTreeSet accessing ROOT data file.

Usage

```
root.data(xps.scheme, rootfile = character(0), celnames = "*")
```

root.density

Arguments

| xps.scheme | A SchemeTreeSet containing the correct scheme for the ROOT data file. |
|------------|---|
| rootfile | name of ROOT data file, including full path. |
| celnames | optional character vector of tree names to get only subset of trees. |

Details

An S4 class DataTreeSet will be created, serving as R wrapper to the existing ROOT data file rootfile.

If the DataTreeSet should only handle a subset of the trees stored in rootfile, the tree names must be supplied as vector celnames.

To get the names of all trees stored in rootfile you can call function getTreeNames first.

Value

A DataTreeSet object.

Note

Use root.data to access the ROOT data file from new R sessions to avoid creating a new ROOT data file for every R session.

Author(s)

Christian Stratowa

See Also

import.data,DataTreeSet

Examples

```
## get scheme and import CEL-files from package
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- import.data(scheme.test3,"tmp_datatest3",celdir=paste(.path.package("xps"),</pre>
```

```
## use subset of CEL-files
subdata.test3 <- root.data(scheme.test3,"tmp_datatest3_cel.root", celnames=c("TestA1.cel"</pre>
```

root.density ROOT Density Plot

Description

Creates a ROOT density plot for one or all ROOT tree(s).

Usage

```
root.density(x, treename = "*", logbase = "log2", canvasname = "DensityPlot", sa
```

Arguments

| х | object of class DataTreeSet or ExprTreeSet. |
|------------|---|
| treename | name of tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT density plot for one or all tree(s) present in rootfile.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

root.hist1D

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.density(data.test3, "*")
```

```
root.density(data.test3, "TestA1.cel")
root.density(data.test3, "TestA1.cel", save.as="png")
```

End(Not run)
root.expr

Description

Create class ExprTreeSet accessing ROOT expression file.

Usage

```
root.expr(xps.scheme, rootfile = character(0), treetype = character(0), treename
```

Arguments

| xps.scheme | A SchemeTreeSet containing the correct scheme for the ROOT data file. |
|------------|---|
| rootfile | name of ROOT data file, including full path. |
| treetype | tree type. |
| treenames | optional character vector of tree names to get only subset of trees. |

Details

An S4 class ExprTreeSet will be created, serving as R wrapper to the existing ROOT expression file rootfile.

Parameter treetype must be supplied to identify the ROOT trees for slot data. Valid tree types are listed in validTreetype.

To get the names of all trees with their extensions treetype, which are stored in rootfile, you can call function getTreeNames first.

If the ExprTreeSet should only handle a subset of the trees stored in rootfile, the tree names must be supplied as vector treenames.

Value

A ExprTreeSet object.

Author(s)

Christian Stratowa

See Also

root.data,root.call

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
# rma
all.rma <- rma(data.test3,"tmp_Test3RMAAll",tmpdir="",background="pmonly",normalize=TRUE,
## use subset of trees
sub.rma <- root.expr(scheme.test3, "tmp_Test3RMAAll.root", "mdp", c("TestA2.mdp", "TestB1</pre>
```

root.graph1D

root.graph1D ROOT1D-Graph

Description

Creates a ROOT 1D-graph for a ROOT tree.

Usage

```
root.graph1D(x, treename = character(0), logbase = "log2", option = "P", canvasn
```

Arguments

| х | object of class DataTreeSet or ExprTreeSet. |
|------------|---|
| treename | name of tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| option | ROOT TGraph::PaintGraph option, usually one of "P", "*", "L". |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT 1D-graph for tree treename present in rootfile.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

root.graph2D

root.graph2D

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.graph1D(data.test3, "TestA1.cel")
## End(Not run)
```

root.graph2D ROOT 2D-Graph

Description

Creates a ROOT 2D-graph for a ROOT tree.

Usage

```
root.graph2D(x, treename1 = character(0), treename2 = character(0), logbase = "lo
```

Arguments

| х | object of class DataTreeSet or ExprTreeSet. |
|------------|---|
| treename1 | name of first tree, must be present in rootfile of object x. |
| treename2 | name of second tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| option | ROOT TGraph::PaintGraph option, usually one of "P", "*", "L". |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT 2D-graph for trees treename1 and treename2 present in rootfile.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

root.hist1D

Author(s)

Christian Stratowa

See Also

root.graph1D, root.mvaplot

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/'
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.graph2D(data.test3, "TestA1.cel", "TestB1.cel")
## End(Not run)
```

root.hist1D ROOT1D-Histogram

Description

Creates a ROOT 1D-histogram for a ROOT tree.

Usage

```
root.hist1D(x, treename = character(0), logbase = "log2", type = "hist", option
```

Arguments

| х | object of class DataTreeSet or ExprTreeSet. |
|------------|---|
| treename | name of tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| type | ROOT 1D-hist or density, i.e. "hist" or "density". |
| option | ROOT 1D-hist option only, usually one of "HIST", "B", "C", "E". |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT 1D-histogram for tree treename present in rootfile.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

root.hist2D

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

root.hist2D,root.hist3D

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.hist1D(data.test3, "TestA1.cel")
root.hist1D(data.test3, "TestA1.cel", type="density")
## End(Not run)
```

root.hist2D ROOT 2D-Histogram

Description

Creates a ROOT 2D-histogram for a ROOT tree.

Usage

```
root.hist2D(x, treename1 = character(0), treename2 = character(0), logbase = "log
```

Arguments

| Х | object of class DataTreeSet or ExprTreeSet. |
|------------|---|
| treename1 | name of first tree, must be present in rootfile of object x. |
| treename2 | name of second tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| option | ROOT hist TH2 option, usually one of "SCAT", "COLZ", "BOX", "SURF2", "SURF3". |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT 2D-histogram for trees treename1 and treename2 present in rootfile.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

root.hist1D,root.hist3D

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.hist2D(data.test3, "TestA1.cel", "TestB1.cel", option="COLZ")
## End(Not run)
```

root.hist3D ROOT 3D-Histogram

Description

Creates a ROOT 3D-histogram for a ROOT tree.

Usage

```
root.hist3D(x, treename1 = character(0), treename2 = character(0), treename3 = c
```

Arguments

| Х | object of class DataTreeSet or ExprTreeSet. |
|-----------|---|
| treename1 | name of first tree, must be present in rootfile of object x. |
| treename2 | name of second tree, must be present in rootfile of object x. |
| treename3 | name of third tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| option | ROOT hist TH3 option, usually one of "HIST", "SCAT", "BOX". |

root.image

| canvasname | name of ROOT canvas |
|------------|---|
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT 3D-histogram for trees treename1, treename2 and treename3 present in rootfile. By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

By moving the mouse into the middle of the canvas, the cursor changes and you can rotate the 3Dhistogram. By selecting menu "View->View With->OpenGL" the OpenGL viewer opens, where you can rotate the 3D-histogram interactively.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

root.hist1D,root.hist2D

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.hist3D(data.test3, "TestA1.cel", "TestB2.cel", "TestB1.cel", option="BOX")
```

End(Not run)

root.image ROOT Image

Description

Creates a ROOT image for a ROOT tree.

Usage

```
root.image(x, treename = character(0), leafname = "fInten", logbase = "log2", op
```

Arguments

| х | object of class DataTreeSet. |
|------------|---|
| treename | name of tree, must be present in rootfile of object x. |
| leafname | leaf name of tree, usual "fInten" or "fBg". |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| option | ROOT graph option, usually. one of "COL", "COLZ". |
| zlim | size limits c(min,max) of leafname. |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the device in pixels. |
| h | the height of the device in pixels. |

Details

Creates a ROOT image for tree treename present in rootfile.

To zoom-in move the mouse cursor to the x-axis (y-axis) until it changes to a hand and click-drag to select an axis-range. To unzoom move the mouse cursor to the x-axis (y-axis) until it changes to a hand and right-click to select "Unzoom".

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

image-methods, image

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.image(data.test3, "TestA1.cel")
root.image(data.test3, "TestA1.cel", save.as="png")
```

End(Not run)

root.merge.data Create class DataTreeSet by merging ROOT data files

Description

Create class DataTreeSet by merging different ROOT data files.

Usage

```
root.merge.data(xps.scheme, rootfiles = list(), celnames = "*")
```

Arguments

| xps.scheme | A SchemeTreeSet containing the correct scheme for the ROOT data file. |
|------------|---|
| rootfiles | list of ROOT data file(s), including full path. |
| celnames | optional character vector of tree names to get only subset of trees. |

Details

This function allows to merge data trees from different existing ROOT data files.

An S4 class DataTreeSet will be created, serving as R wrapper to the existing ROOT data file(s) rootfiles.

If the DataTreeSet should only handle a subset of the trees stored in rootfiles, the tree names must be supplied as vector celnames.

To get the names of all trees stored in separate rootfiles you can call function getTreeNames first.

Value

A DataTreeSet object.

Author(s)

Christian Stratowa

See Also

root.data,DataTreeSet

Examples

```
## get scheme and import CEL-files from package
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- import.data(scheme.test3,"tmp_datatest3",celdir=paste(.path.package("xps"),
## get subset of CEL-files
subdataA <- root.data(scheme.test3,"tmp_datatest3_cel.root", celnames=c("TestA1.cel","Tes
subdataB <- root.data(scheme.test3,"tmp_datatest3_cel.root", celnames=c("TestB1.cel","Tes
## merge data</pre>
```

```
dataAB <- root.merge.data(scheme.test3,c(rootFile(subdataA),rootFile(subdataB)), celnames</pre>
```

root.mvaplot

root.mvaplot

Description

Creates a ROOT M vs A plot for a ROOT tree.

Usage

```
root.mvaplot(x, treename1 = character(0), treename2 = character(0), logbase = "lo
```

Arguments

| х | object of class ExprTreeSet or DataTreeSet. |
|------------|---|
| treename1 | name of first tree, must be present in rootfile of object x. |
| treename2 | name of second tree, must be present in rootfile of object x. |
| logbase | usually "log2", or "0", determines if leaf data should be converted to log. |
| option | ROOT TGraph::PaintGraph option, usually one of "P", "*". |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the canvas in pixels. |
| h | the height of the canvas in pixels. |

Details

Creates a ROOT M vs A plot for trees treename1 and treename2 present in rootfile.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT canvas, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

See Also

root.graph1D

root.profile

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
# compute RMA
data.rma <- rma(data.test3,"Test3RMA",tmpdir="",background="pmonly",normalize=TRUE)
root.mvaplot(data.rma, "TestA1.mdp", "TestB1.mdp")
## End(Not run)
```

root.profile ROOT Profile Plot

Description

Creates a ROOT profile plot, i.e. a plot of parallel coordinates

Usage

root.profile(x, treename = "*", varlist = NULL, as.log = TRUE, globalscale = TRU

Arguments

| х | S4 object, usually of class DataTreeSet or ExprTreeSet. |
|-------------|---|
| treename | name of tree, usually all trees present in rootfile of object x. |
| varlist | leaf name of tree, usual "fInten" or "fLevel". |
| as.log | logical indicating if varlist should be drawn as logarithmic data. |
| globalscale | logical indicating if all axes should be drawn at the same scale. |
| boxes | logical indicating if box-and-whisker plots should be drawn. |
| ylim | <pre>size limits c(min,max) of varlist.</pre> |
| canvasname | name of ROOT canvas |
| save.as | graphics type for saving canvas, one of "ps", "eps", "pdf", "jpg", "gif", "png", "tiff" |
| W | the width of the device in pixels. |
| h | the height of the device in pixels. |

Details

Creates a ROOT profile plot for all trees treename="*" present in rootfile. In this case varlist must be the name of one tree leaf only; for varlist=NULL leaf "fInten" will be used for class DataTreeSet and leaf "fLevel" will be used for class ExprTreeSet. If treename is the name of one tree only then varlist can contain up to all leaves of the tree, separated by colons, e.g. varlist="fLevel:fStdev".

For boxes=TRUE the profile plot draws box-and-whisker plots and can thus be considered the equivalent of the usual boxplot.

A ROOT profile plot, i.e. a plot of parallel coordinates, is drawn in a "TreeViewer", a graphic user interface designed to handle ROOT trees. You can activate context menus by right-clicking on items or inside the right panel.

The "TreeViewer" is explained in http://root.cern.ch/root/html/TTreeViewer. html.

By selecting menu "File->Save->canvasname.xxx" you can save the figure as e.g. *gif, *.jpg, *.pdf, *.ps or even as C++ macro.

Alternatively, you can save the plot by setting save.as. However, this will close the canvas immediately after opening it.

Note

Always select menu item "Quit ROOT" from menu "File" to close the ROOT tree viewer, otherwise you are in the CINT C/C++ interpreter from ROOT. To exit CINT, you need to type ".q".

Author(s)

Christian Stratowa

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.
root.profile(data.test3)
```

End(Not run)

ROOT

ROOT An Object-Oriented Data Analysis Framework

Description

ROOT system overview

Details

ROOT is a modular object-oriented framework aimed at solving the data analysis challenges of high-energy physics. The relevant features of ROOT are as follows:

Architecture: The ROOT architecture is a layered class hierarchy with over 500 classes divided into different categories. Most of the classes inherit from a common base class TObject, which provides the default behavior and protocol for all objects.

ROOT Files: Object input/output is handled by class TFile, which has a UNIX-like directory structure and provides a hierarchical sequential and direct access persistent object store. ROOT files store information in a machine independent format and support on-the-fly data compression. Furthermore, ROOT files are self-describing: for every object stored in TFile, a dictionary describing the corresponding class is written to the file. A dictionary generator, called ROOTCINT, parses the class header files and generates a dictionary. Note: TFile can be considered to be the ROOT analogon to an R environment.

root.scheme

Data Trees: Any object derived from TObject can be written to a file with an associated key TKey. However, each key has an overhead in the directory structure in memory. To reduce this overhead, a novel concept, called Trees (class TTree) has been developed. Trees are designed to support very large numbers of complex objects in a large number of files. A Tree consists of branches (TBranch) with each branch described by its leaves (TLeaf). Trees allow direct and random access to any entry of a selected subset of branches. Thus, Trees extend and replace the usual data tables. The concept of Tree friends allows the joining of many trees as one virtual tree. However, unlike table joins in an RDBMS, the processing time is independent of the number of tree friends. Note: TTree can be considered to be the ROOT analogon to an R data.frame.

CINT: CINT is an interactive C/C++ interpreter, which is aimed at processing C/C++ scripts, called macros. Currently, CINT covers 99% of ANSI C and 95% of ANSI C++. CINT offers a gdb-like debugger for interpreted programs and allows the automatic compilation of scripts using ACLiC, the automatic compiler of libraries for CINT. Although available as independent program, CINT is embedded in ROOT as command line interpreter and macro processor, as well as dictionary generator.

User interaction: The ROOT system can be accessed from the command line, by writing macros, or via a graphic user interface (e.g. RootBrowser). Furthermore, it is possible to write libraries and applications. The ROOT GUI classes allow the development of full-featured standalone applications. Note: A macro can be considered to be the ROOT analogon of an R script. The RootBrowser can be opened using function root.browser

Platform independence: The ROOT system is available for most platforms and operating systems, including Linux, MacOS X, and the major flavors of UNIX and Windows. ROOT and ROOT-derived applications can be compiled for any supported platform.

Author(s)

The ROOT team http://root.cern.ch/root/Authors.html

References

ROOT User Guide http://root.cern.ch/root/doc/RootDoc.html

ROOT publications http://root.cern.ch/root/Publications.html

Christian Stratowa (2003), Distributed Storage and Analysis of Microarray Data in the Terabyte Range: An Alternative to BioConductor http://www.ci.tuwien.ac.at/Conferences/ DSC-2003/Proceedings/Stratowa.pdf

root.scheme Create class SchemeTreeSet accessing ROOT scheme file

Description

Create class SchemeTreeSet accessing ROOT scheme file.

Usage

root.scheme(rootfile = character(0), add.mask = FALSE)

Arguments

| rootfile | name of ROOT scheme file, including full path. |
|----------|---|
| add.mask | if TRUE mask information will be included as slot mask. |

Details

An S4 class SchemeTreeSet will be created, serving as R wrapper to the ROOT scheme file rootfile.

Value

A SchemeTreeSet object.

Note

Use this function to access the ROOT scheme file from new R sessions to avoid creating a new ROOT scheme file for every R session.

Do not set add.mask=TRUE for exon arrays unless you know that your computer has sufficient RAM.

Author(s)

Christian Stratowa

See Also

import.expr.scheme, import.exon.scheme, SchemeTreeSet

Examples

```
## create class SchemeSet to access the ROOT scheme file for the Test3 GeneChip
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
str(scheme.test3)
## Not run:
## scheme set for existing human root exon scheme file
scheme.huex10stv2r2.na22 <- root.scheme("/my/path/schemes/Scheme_HuEx10stv2r2_na22.root")</pre>
```

End(Not run)

SchemeTreeSet-class

Class SchemeTreeSet

Description

This class provides the link to the ROOT scheme file and the ROOT trees contained therein. It extends class TreeSet.

Objects from the Class

Objects can be created using the functions import.expr.scheme, import.exon.scheme, import.genome.scheme or root.scheme.

SchemeTreeSet-class

Slots

chipname: Object of class "character" representing the Affymetrix chip name.

- chiptype: Object of class "character" representing the chip tpye, either 'GeneChip', 'GenomeChip' or 'ExonChip'.
- probeinfo: Object of class "list" representing chip information, including nrows, ncols, number of probes, etc.
- mask: Object of class "data.frame". The data.frame can contain the mask used to identify the probes as e.g. PM, MM or control probes.
- setname: Object of class "character" representing the name to the ROOT file subdirectoy where the ROOT scheme trees are stored; it is identical to chipname.
- settype: Object of class "character" describing the type of treeset stored in setname, i.e. 'scheme'.
- rootfile: Object of class "character" representing the name of the ROOT scheme file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

Extends

Class "TreeSet", directly.

Methods

- **chipMask** signature(object = "SchemeTreeSet"): **extracts** data.frame mask.
- chipMask<- signature(object = "SchemeTreeSet", value = "data.frame"):
 replaces data.frame mask.</pre>
- **chipName** signature(object = "SchemeTreeSet"): extracts slot chipname.
- chipType signature(object = "SchemeTreeSet"): extracts slot chiptype.
- chipType<- signature(object = "SchemeTreeSet", value = "character"):replaces slot chiptype.
- export signature(object = "SchemeTreeSet"): exports ROOT trees as text file, see
 export-methods.
- ncols signature(object = "SchemeTreeSet"): extracts the physical number of array columns from slot probeinfo.
- nrows signature(object = "SchemeTreeSet"): extracts the physical number of array
 rows from slot probeinfo.
- probeInfo signature(object = "SchemeTreeSet"): extracts slot probeinfo.
- removeMask signature(object = "SchemeTreeSet"): replaces data.frame mask with an empty data.frame of dim(0,0).

summarize

Author(s)

Christian Stratowa

Examples

showClass("SchemeTreeSet")

summarize

Probe Set Summarizing Functions

Description

Converts Affymetrix probe level data to expression levels by summarizing the probe set values into one expression measure and a standard error for this summary.

Usage

```
summarize(xps.data, filename = character(0), filedir = getwd(), tmpdir = "", upd
summarize.mas4(xps.data, filename = character(0), filedir = getwd(), tmpdir = ""
summarize.mas5(xps.data, filename = character(0), filedir = getwd(), tmpdir = ""
summarize.rma(xps.data, filename = character(0), filedir = getwd(), tmpdir = "",
xpsSummarize(object, ...)
```

Arguments

| xps.data | object of class DataTreeSet. |
|------------|--|
| filename | file name of ROOT data file. |
| filedir | system directory where ROOT data file should be stored. |
| tmpdir | optional temporary directory where temporary ROOT files should be stored. |
| update | logical. If TRUE the existing ROOT data file filename will be updated. |
| select | type of probes to select for summarization. |
| method | summarization method to use. |
| option | option determining the grouping of probes for summarization, one of 'tran- script', 'exon', 'probeset'; exon arrays only. |
| logbase | logarithm base as character, one of '0', 'log', 'log2', 'log10'. |
| exonlevel | exon annotation level determining which probes should be used for summariza- tion; exon/genome arrays only. |
| params | vector of parameters for summarization method. |
| xps.scheme | optional alternative SchemeTreeSet. |
| add.data | logical. If TRUE expression data will be included as slot data. |
| verbose | logical, if TRUE print status information. |
| object | object of class DataTreeSet. |
| | the arguments described above. |

TreeSet-class

Details

Converts Affymetrix probe level data to expression levels by summarizing the probe set values into one expression measure and a standard error for this summary.

xpsSummarize is the DataTreeSet method called by function summarize, containing the same parameters.

Value

An ExprTreeSet.

Author(s)

Christian Stratowa

See Also

express

Examples

```
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
## RMA background
data.bg.rma <- bgcorrect.rma(data.test3,"tmp_Test3RMA",filedir=getwd(),tmpdir="",verbose=
## normalize quantiles
data.qu.rma <- normalize.quantiles(data.bg.rma,"tmp_Test3RMA",filedir=getwd(),tmpdir="",v
## summarize medianpolish
data.mp.rma <- summarize.rma(data.qu.rma,"tmp_Test3RMA",filedir=getwd(),tmpdir="",update=</pre>
```

```
## get expression data.frame
expr.rma <- exprs(data.mp.rma)
head(expr.rma)</pre>
```

```
## plot expression levels
if (interactive()) {
    boxplot(data.mp.rma)
    boxplot(log2(expr.rma[,3:6]))
}
```

TreeSet-class Class TreeSet

Description

This is the virtual base class for all other classes providing the link to a ROOT file and the ROOT trees contained therein.

Objects from the Class

A virtual Class: No objects may be created from it.

Slots

- setname: Object of class "character" representing the name to the ROOT file subdirectory where the ROOT trees are stored, usually one of 'DataTreeSet', 'PreprocesSet', 'CallTreeSet'.
- settype: Object of class "character" describing the type of treeset stored in setname, usually one of 'scheme', 'rawdata', 'preprocess'.
- rootfile: Object of class "character" representing the name of the ROOT file, including full path.
- filedir: Object of class "character" describing the full path to the system directory where rootfile is stored.
- numtrees: Object of class "numeric" representing the number of ROOT trees stored in subdirectoy setname.
- treenames: Object of class "list" representing the names of the ROOT trees stored in subdirectoy setname.

Methods

- export signature(object = "TreeSet"): exports ROOT trees as text file, see exportmethods.
- fileDir signature(object = "TreeSet"): extracts slot filedir.
- fileDir<- signature(object = "TreeSet", value = "character"): replaces slot
 filedir.</pre>
- root.browser signature(object = "TreeSet"): opens the ROOT file browser.
- rootFile signature(object = "TreeSet"): extracts slot rootfile.
- rootFile<- signature(object = "TreeSet", value = "character"): replaces slot
 rootfile.</pre>
- setName signature(object = "TreeSet"): extracts slot setname.
- setName<- signature(object = "TreeSet", value = "character"): replaces slot
 setname.</pre>
- setType signature(object = "TreeSet"): extracts slot settype.
- setType<- signature(object = "TreeSet", value = "character"): replaces slot
 settype.</pre>
- treeNames signature(object = "TreeSet"): extracts slot treenames.

Author(s)

Christian Stratowa

See Also

derived classes SchemeTreeSet, DataTreeSet, ExprTreeSet, CallTreeSet.

Examples

showClass("TreeSet")

type2Exten

Description

Convert Method Type to Tree Extension.

Usage

```
type2Exten(type, datatype)
```

Arguments

| type | method type. |
|----------|--------------|
| datatype | data type. |

Details

For every datatype different methods, i.e. algorithms exist which can be applied. Valid datatypes are 'preprocess' and 'normation'.

For datatype 'preprocess' the following methods can be applied:

| mean: | trimmed mean |
|----------------|----------------|
| median: | median |
| quantile: | quantile |
| tukeybiweight: | tukey biweight |
| medianpolish: | median polish |

For datatype 'normation' the following methods can be applied:

| mean: | trimmed mean |
|-----------|--------------|
| median: | median |
| quantile: | quantile |
| lowess: | lowess |
| supsmu: | supsmu |

The tree extensions are described in validTreetype.

Value

A character with the correct tree extension.

Author(s)

Christian Stratowa

See Also

getDatatype,validTreetype

Examples

```
type2Exten("quantile", "preprocess")
type2Exten("medianpolish", "preprocess")
type2Exten("supsmu", "normation")
```

UniFilter-class Class UniFilter

Description

Class UniFilter allows to apply different unitest filters to class ExprTreeSet, i.e. to the expression level data.frame data.

Objects from the Class

```
Objects can be created by calls of the form new ("UniFilter", ...). Alternatively, the contructor UniFilter can be used.
```

Slots

foldchange: Object of class "list" describing parameters for fcFilter.
prescall: Object of class "list" describing parameters for callFilter.
unifilter: Object of class "list" describing parameters for unitestFilter.
unitest: Object of class "list" describing parameters for uniTest.
numfilters: Object of class "numeric" giving the number of filters applied.

Extends

Class "Filter", directly.

Methods

- **callFilter** signature(object = "UniFilter"): extracts slot prescall.
- callFilter<- signature(object = "UniFilter", value = "character"): replaces
 slot prescall with character vector c(cutoff, samples, condition).</pre>
- fcFilter signature(object = "UniFilter"): extracts slot foldchange.
- fcFilter<- signature(object = "UniFilter", value = "numeric"): replaces slot
 foldchange with numeric vector c(cutoff, direction).</pre>
- uniTest signature(object = "UniFilter"): extracts slot unitest.
- uniTest<- signature(object = "UniFilter", value = "character"):replaces
 slot unitest with character vector c(type, alternative, correction, numperm, mu, paired,
 conflevel, varequ).</pre>

```
unitestFilter signature(object = "UniFilter"): extracts slot unifilter.
```

unitestFilter<- signature(object = "UniFilter", value = "character"):replaces slot unifilter with character vector c(cutoff, variable).

Author(s)

Christian Stratowa

UniFilter-constructor

See Also

related classes Filter, PreFilter.

Examples

```
unifltr <- new("UniFilter", unitest=list("t.test"))
fcFilter(unifltr) <- c(1.5,"both")
unitestFilter(unifltr) <- c(0.01,"pval")
str(unifltr)</pre>
```

UniFilter-constructor

Constructor for Class UniFilter

Description

Constructor for class UniFilter allows to apply different unitest filters to class ExprTreeSet, i.e. to the expression level data.

Usage

```
UniFilter(unitest = "t.test",
    foldchange = character(),
    prescall = character(),
    unifilter = character())
```

Arguments

| unitest | "character" vector describing parameters for uniTest. |
|------------|---|
| foldchange | "character" vector describing parameters for fcFilter. |
| prescall | "character" vector describing parameters for callFilter. |
| unifilter | "character" vector describing parameters for unitestFilter. |

Details

The UniFilter constructor allows to apply the following unitest filters to class ExprTreeSet:

| unitest: | character vector c(type,alternative,correction.numperm,mu,paired,conflevel,varequ). |
|-------------|---|
| foldchange: | character vector c(cutoff,direction). |
| prescall: | character vector c(cutoff,samples,condition). |
| unifilter: | character vector c(cutoff,variable). |

Value

```
An object of type "UniFilter"
```

Note

Function UniFilter is used as constructor for class UniFilter so that the user need not know details for creating S4 classes.

unifilter

Author(s)

Christian Stratowa

See Also

UniFilter, PreFilter

Examples

```
fcFilter(unifltr) <- c(1.5, "both")
unitestFilter(unifltr) <- c(0.01, "pval")
str(unifltr)</pre>
```

unifilter

Function for Applying an UniFilter to an ExprTreeSet

Description

This function applies an UniFilter to an ExprTreeSet.

Usage

```
unifilter(xps.expr,
         filename = character(0),
         filedir = getwd(),
         filter
                 = NULL,
         minfilters = 999,
                  = "log2",
         logbase
                  = character(0),
         group
         treename
                   = "UniTest",
         xps.fltr
                   = NULL,
         xps.call = NULL,
         update = FALSE,
         verbose = TRUE)
```

xpsUniFilter(object, ...)

Arguments

| xps.expr | object of class ExprTreeSet. |
|----------|---|
| filename | file name of ROOT filter file. |
| filedir | system directory where ROOT filter file should be stored. |
| filter | object of class UniFilter. |

unifilter

| minfilters | minimum number of initialized filter methods to satisfy (default is all filters). |
|------------|---|
| logbase | convert data to logarithm of base: "0", "log", "log2" (default), "log10" |
| group | a character vector assigning the trees of xps.expr to one of two groups. |
| treename | tree name to be used in ROOT filter file. |
| xps.fltr | optional object of class FilterTreeSet. |
| xps.call | optional object of class CallTreeSet. |
| update | logical. If TRUE the existing ROOT filter file filename will be updated. |
| verbose | logical, if TRUE print status information. |
| object | object of class ExprTreeSet. |
| | same arguments as function unifilter. |

Details

This function applies the different filters initialized with constructor UniFilter to the ExprTreeSet xps.expr.

Slot minfilters determines the minimum number of initialized filters, which must be satisfied so that the mask is set to flag=1. For minfilters=1 at least one filter must be satisfied, equivalent to logical 'OR'; for minfilters=999 all filters must be satisfied, equivalent to logical 'AND'.

If pre-filtering should be done before applying function unifilter then a FilterTreeSet xps.fltr must be supplied, created with function prefilter.

If method callFilter was initialized with constructor UniFilter then CallTreeSet xps.call must be supplied, usually created with function mas5.call.

Value

An AnalysisTreeSet

Note

Internally, slot group will be converted to integer values using as.integer (as.factor(group)), thus group=c("GrpA", "GrpA", "GrpB", "GrpB") will result in a fold-change of fc=mean(GrpB)/mean(Gr

Author(s)

Christian Stratowa

See Also

UniFilter, prefilter

Examples

```
## Not run:
## first, load ROOT scheme file and ROOT data file
scheme.test3 <- root.scheme(paste(.path.package("xps"),"schemes/SchemeTest3.root",sep="/"
data.test3 <- root.data(scheme.test3, paste(.path.package("xps"),"rootdata/DataTest3_cel.</pre>
```

```
## second, create an ExprTreeSet
data.rma <- rma(data.test3,"tmp_Test3_RMA",tmpdir="",background="pmonly",normalize=TRUE,v
## note: do not copy/paste this code, it is necessary only because R CMD check fails sinc
data.rma@rootfile <- paste(.path.package("xps"),"rootdata/tmp_Test3RMA.root",sep="/")</pre>
```

```
data.rma@filedir <- paste(.path.package("xps"), "rootdata", sep="/")
## third, construct an UniFilter
unifiltr <- UniFilter(unitest=c("t.test", "two.sided", "none", 0, 0.0, FALSE, 0.95, TRUE), foldcha
## finally, create an AnalysisTreeSet
rma.ufr <- unifilter(data.rma, "tmp_Test3Unifilter", getwd(), unifiltr, group=c("GrpA", "GrpA",
str(rma.ufr)
## End(Not run)</pre>
```

unitestFilter-methods

Unitest Filter

Description

This method initializes the Unitest Filter.

Applying an unitest such as the t.test to two groups returns the p-value for the test and the value of the t-statistic. The Unitest Filter allows to select only rows satisfying e.g. a certain p-value as cutoff.

```
The Unitest Filter flags all rows with: flag = (variable <= cutoff)
Usage
unitestFilter(object)
unitestFilter(object, value)<-</pre>
```

Arguments

| object | object of class UniFilter. | |
|--------|-----------------------------|------------|
| value | character vector c (cutoff, | variable). |

Details

The method unitestFilter initializes the following parameters:

```
cutoff: the cutoff level for the filter.
variable: variable="pval" (default): p-value.
variable="stat": univariate statistic.
variable="padj": optional adjusted p-value.
variable="pcha": optional p-value obtained by permutations.
```

Value

An initialized UniFilter object.

Author(s)

Christian Stratowa

Examples

unifltr <- UniFilter()</pre>

uniTest-methods

```
unitestFilter(unifltr) <- c(0.01,"pval")
str(unifltr)</pre>
```

uniTest-methods A Two-Group Unitest

Description

Unitest performs a a two group uni-test such as the t.test on each row of the expression dataframe. The Unitest returns a dataframe containing the results of the test.

Usage

```
uniTest(object)
uniTest(object, value)<-</pre>
```

Arguments

| object | object of class UniFilter. | | | | |
|--------|----------------------------|--------------|-------------|----------|-----|
| value | character vector c (type, | alternative, | correction, | numperm, | mu, |
| | paired, conflevel, | varequ) | | | |

Details

The method uniTest initializes the following parameters:

| type: | a character string specifying the type of test: currently "t.test" (default) or "normal.test". | |
|--------------|---|--|
| alternative: | e: a character string specifying the alternative hypothesis, must be one of "two.sided" (default | |
| correction: | a correction to adjust p-values for multiple comparisons: | |
| | correction="none": no correction (default). | |
| | correction="bonferroni": Bonferroni correction. | |
| | correction="BH" or "fdr": correction for false discovery rate (Benjamini & Hochberg). | |
| | correction="BY": correction for false discovery rate (Benjamini & Yekutieli). | |
| | correction="hochberg": Hochberg correction. | |
| | correction="holm": Holm correction. | |
| | correction="wy": Westfall-Young step-down adjusted p-chance (E.Manduchi). | |
| numperm: | optional number of permutations used to determine p-chance (default is 0). | |
| mu: | a number indicating the true value of the difference in means for a two sample test (default is 0). | |
| paired: | a logical indicating whether you want a paired uni-test (default is FALSE). | |
| conflevel: | confidence level of the interval (default is 0.95). | |
| varequ: | a logical variable indicating whether to treat the two variances as being equal. If TRUE then the poo | |

Value

An initialized UniFilter object.

Author(s)

Christian Stratowa

References

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Dudoit S., Yang Y.H., Callow M.J., Speed T.P. (2000) Statistical methods for identifying differentially expressed genes in replicated cDNA microarray experiments. *Technical report* **578**; UC Berkeley.

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Examples

```
unifltr <- UniFilter()
uniTest(unifltr) <- c("t.test", "two.sided", "none", 0, 0.0, FALSE, 0.98, TRUE)
str(unifltr)</pre>
```

validCall-methods Get Valid Detection Call Values

Description

Extracts valid present call values with unit names as row names.

```
Usage
```

```
validCall(object, which = "UnitName")
validPVal(object, which = "UnitName")
```

Arguments

| object | object of class CallTreeSet. |
|--------|--------------------------------------|
| which | name of column containing unit name. |

Details

Method validCall returns the present calls from slot detcall as data.frame and uses column which as row names, usually the probeset IDs stored in column "UnitName".

Method validPVal returns the detection call p-values from slot data as data.frame and uses column which as row names, usually the probeset IDs stored in column "UnitName".

Value

A data.frame.

Author(s)

Christian Stratowa

validData-methods

See Also

validData,validExpr

validData-methods Extract Subset of Data

Description

Extracts a subset of valid data from data.frame data.

Usage

validData(object, which = "")

Arguments

| object | object of class DataTreeSet, ExprTreeSet or CallTreeSet. |
|--------|---|
| which | type of probes to be returned for DataTreeSet, otherwise name of column containing unit name. |

Details

For class DataTreeSet and expression arrays, validData returns all the perfect match or mismatch probes on the arrays the object represents as data.frame, i.e. which can have the following values:

| pm: | perfect match probes. |
|-------|---|
| mm: | mismatch probes. |
| both: | both perfect match and mismatch probes. |

For class DataTreeSet and exon arrays, validData returns the probes of the different exon levels as data.frame, i.e. which can have one of the following values:

| core: | probesets supported by RefSeq and full-length GenBank transcripts. |
|---------------|--|
| metacore: | core meta-probesets. |
| extended: | probesets with other cDNA support. |
| metaextended: | extended meta-probesets. |
| full: | probesets supported by gene predictions only. |
| metafull: | full meta-probesets. |
| affx: | standard AFFX controls. |
| all: | combination of above. |
| genomic: | genomic background probes. |
| antigenomic: | antigenomic background probes. |
| | |

For class ExprTreeSet validData returns the valid expression levels from slot data with unit names as row names, usually the probeset IDs stored in column which="UnitName".

For class CallTreeSet validData returns the valid detection call p-values from slot data with unit names as row names, usually the probeset IDs stored in column which="UnitName".

Value

A data.frame.

Author(s)

Christian Stratowa

See Also

pm, mm, validExpr, validCall

validExpr-methods Get Valid Expression Levels

Description

Extracts valid expression levels with unit names as row names from data.frame data.

Usage

validExpr(object, which = "UnitName")

Arguments

| object | object of class ExprTreeSet. |
|--------|--------------------------------------|
| which | name of column containing unit name. |

Details

Method validExpr returns the expression levels from slot data and uses column which as row names, usually the probeset IDs stored in column "UnitName".

Value

A data.frame.

Author(s)

Christian Stratowa

See Also

validData,validCall

validTreetype Validate Tree Type

Description

Validate tree type for corresponding data type.

Usage

validTreetype(treetype, datatype)

Arguments

| treetype | tree type. |
|----------|------------|
| datatype | data type. |

Details

Every ROOT tree has an extension, which describes the type of data stored in this tree. For example, 'TestA1.cel' is the tree name that stores the CEL-file data for 'TestA1.CEL'.

Trees with datatype="scheme" have the following extensions:

scm: scheme tree containing (x,y)-coordinates and mask for UNIT_ID.

idx: unit tree containing UnitName (i.e. probeset id), NumCells, NumAtoms, UnitType, for UNIT_ID.

prb: probe tree containing probe sequences.

ann: transcript annotation tree.

anx: exon annotation tree; exon arrays only.

anp: probeset annotation tree; exon arrays only.

cxy: coordinate tree containing CLF-file information; exon arrays only.

exn: exon tree; exon arrays only.

pbs: probeset tree; exon arrays only.

Trees with datatype="rawdata" have the following extensions:

cel: data tree containing CEL-file data.

Trees with datatype="preprocess" have the following extensions:

int: intensity tree containing background-corrected intensities.

sbg: background tree containing MAS4 sector background levels.

wbg: background tree containing MAS5 weighted sector background levels.

rbg: background tree containing RMA background levels.

gbg: background tree containing GC-content background levels.

cmn: cell tree containing preprocessed intensities using algorithm 'mean'.

 $\ensuremath{\mathsf{cmd}}$: cell tree containing preprocessed intensities using algorithm 'median'.

clw: cell tree containing preprocessed intensities using algorithm 'lowess'.

css: cell tree containing preprocessed intensities using algorithm 'supsmu'.

 ${\tt cqu:}\ cell\ tree\ containing\ preprocessed\ intensities\ using\ algorithm\ 'quantile'.$

dc5: detection tree containing MAS5 detection call and p-value.

dab: detection tree containing DABG detection call and p-value.

amn: expression tree containing expression levels computed with 'arithmetic mean'.

gmn: expression tree containing expression levels computed with 'geometric mean'.

 ${\tt wmn}:$ expression tree containing expression levels computed with 'weighted mean'.

wdf: expression tree containing expression levels computed with 'weighted difference'.

adf: expression tree containing expression levels computed with 'average difference'. tbw: expression tree containing expression levels computed with 'tukey biweight'. mdp: expression tree containing expression levels computed with 'median polish'.

Trees with datatype="normation" have the following extensions:

tmn: expression tree after normalization using algorithm 'trimmed mean'.

med: expression tree after normalization using algorithm 'median'.

ksm: expression tree after normalization using algorithm 'kernel smoother'.

low: expression tree after normalization using algorithm 'lowess'.

sup: expression tree after normalization using algorithm 'supsmu'.

qua: expression tree after normalization using algorithm 'quantile'.

mdp: expression tree after normalization using algorithm 'median polish'.

Value

Returns the valid treetype, otherwise an error message is returned.

Note

Not all tree types are used in the current package.

Author(s)

Christian Stratowa

See Also

getDatatype, type2Exten

Examples

```
validTreetype("prb", "scheme")
validTreetype("cel", "rawdata")
validTreetype("tbw", "preprocess")
```

varFilter-methods Variance Filter

Description

This method initializes the Variance Filter.
The Variance Filter flags all rows with: flag = (var/mean >= cutoff)
Usage
varFilter(object)
varFilter(object, value)<-</pre>

Arguments

| object | object of class PreFilter. | | |
|--------|-------------------------------------|-------|-----------|
| value | <pre>numeric vector c(cutoff,</pre> | trim, | epsilon). |

volcanoplot-methods

Details

The method varFilter initializes the following parameters:

```
cutoff: the cutoff level for the filter.
trim: the trim value for trimmed mean (default is trim=0).
epsilon: value to replace mean (default is epsilon=0.01):
epsilon > 0: replace mean=0 with epsilon.
epsilon = 0: always set mean=1.
```

Note, that for epsilon = 0 the filter flags all rows with: variance >= cutoff

Value

An initialized PreFilter object.

Author(s)

Christian Stratowa

Examples

```
prefltr <- PreFilter()
varFilter(prefltr) <- c(0.6,0.02,0.01)
str(prefltr)</pre>
```

volcanoplot-methods

Volcano Plot

Description

Produce a scatter plot of fold-change values vs p-values, called volcano plot.

Usage

```
volcanoplot(x, labels = "", p.value = "pval", mask = FALSE, show.cutoff
= TRUE, cex.text = 0.7, col.text = "blue", col.cutoff = "grey", xlim
= NULL, xlab = "Log2(Fold-Change)", ylab = "-Log10(P-Value)", pch =
'.', ...)
```

Arguments

| Х | object of class AnalysisTreeSet. | |
|-------------|---|--|
| labels | optional transcript labels to be drawn at plotting points. | |
| p.value | type of p-value, 'pval' for p-value, 'padj' for adjusted p-value, or 'pcha' for p-chance. | |
| mask | logical, if TRUE draw only points for transcripts satisfying the univariate test. | |
| show.cutoff | logical, if TRUE draw lines indicating cutoff. | |
| cex.text | magnification to be used for optional labels. | |
| col.text | color to be used for optional labels. | |

xpsOptions

| col.cutoff | color to be used for lines indicating cutoff, if show.cutoff=TRUE. |
|------------|---|
| xlim | optional range for the plotted fold-change values. |
| xlab | label of x-axis. |
| ylab | label of y-axis. |
| pch | either an integer specifying a symbol or a single character to be used as the default in plotting points. |
| | optional arguments to be passed to plot. |

Details

Produces a volcano plot for slot data for an object of class AnalysisTreeSet.

It is possible to label the points of the volcano plot, whereby the following labels parameters are valid:

| fUnitName: | unit name (probeset ID). |
|--------------|--------------------------|
| fName: | gene name. |
| fSymbol: | gene symbol. |
| fChromosome: | chromosome. |
| fCytoBand: | cytoband. |

Author(s)

Christian Stratowa

xpsOptions xps Options

Description

Options for xps

Usage

```
xpsOptions(debug=FALSE)
```

Arguments

debug logical, if TRUE, print debug information.

Details

Currently only used to set debug to FALSE or TRUE.

Value

A global variable debug.xps can be set to TRUE.

Author(s)

Christian Stratowa

xps-package xps Package Overview

Description

xps Package Overview

Details

Important data classes: SchemeTreeSet, DataTreeSet, ExprTreeSet, CallTreeSet, FilterTreeSet, AnalysisTreeSet. Full help on methods and associated functions is available from within class help pages.

Additional data classes: ProjectInfo, PreFilter, UniFilter.

The package handles pre-processing, normalization, filtering and analysis of Affymetrix GeneChip expression arrays, including exon array systems (Exon 1.0 ST: core, extended, full probesets), gene array systems (Gene 1.0 ST) and plate array systems on computers with 1 GB RAM only. It imports Affymetrix .CDF, .CLF, .PGF and .CEL as well as Affymetrix annotation files, and computes e.g. RMA, MAS5, FARMS, DFW, MAS5-calls, DABG-calls, I/NI-calls. It is an R wrapper to XPS (eXpression Profiling System), which is based on ROOT, an object-oriented framework developed at CERN. Thus, the prior installation of ROOT is a prerequisite for the usage of this package, see the README file. However, no knowledge of ROOT is required. ROOT is licensed under LGPL and can be downloaded from http://root.cern.ch.

Author(s)

Christian Stratowa <cstrato@aon.at>

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