# codelink

October 5, 2010

Codelink-class Codelink

# Description

This is the storage class for Codelink data

### **Objects from the Class**

Object are created after reading text codelink files with readCodelink()

### Description

Codelink objects contain a single "list", which contains the following elements:

sample Object of class "character" containing the sample names
file Object of class "character" containing the file names
name Object of class "character" containing the probe ID
method Object of class "list" containing log information
Smean Object of class "matrix" containing spot mean intensities
Bmedian Object of class "matrix" containing background median intensities
Ri Object of class "matrix" containing raw intensities
Ni Object of class "matrix" containing normalized intensities
snr Object of class "matrix" containing signal to noise ratio values
flag Object of class "character" containing assigned flags

#### Note

More details are in the package vignette

### Author(s)

Diego Diez

# Examples

## End(Not run)

CodelinkSet-class CodelinkSet

# Description

This is the storage class for Codelink data

### **Objects from the Class**

Object are created after reading text codelink files with readCodelink2()

### Description

CodelinkSet objects are derived from ExpressionSet and therefore inherits all the methods.

### Note

More details are in the package vignette

### Author(s)

Diego Diez

### Examples

```
## End(Not run)
```

CodelinkSetUnique-class
Class CodelinkSetUnique

### Description

This is the storage class for Codelink data- with unique probes

# **Objects from the Class**

Object are created after applying averageProbes() on an object of the class CodelinkSet

# Description

CodelinkSetUnique objects are derived from ExpressionSet and therefore inherits all the methods.

# Note

More details are in the package vignette

# SNR

# Author(s)

Diego Diez

# Examples

```
## Not run:
    data(codelink.exprs)
    foo <- averageProbes(codelink.exprs)</pre>
```

## End(Not run)

SNR

Calculate SNR

# Description

Compute SNR inside read.Codelink.

# Usage

SNR(Smean, Bmedian, Bstdev)

# Arguments

| Smean   | matrix of Smean intensities.             |
|---------|--|
| Bmedian | matrix of background median intensities. |
| Bstdev  | matrix of background standard deviation. |

# Author(s)

Diego Diez

| arrayNew | Create a new x11 device |
|----------|-------------------------|
|          |                         |

# Description

Create a new x11 device with dimensions suited to be used with imageCodelink().

# Usage

```
arrayNew(f=2, chip="rwgcod")
```

# Arguments

| chip | character; Codelink chip to be used in imageCodelink. |
|------|---|
| f    | numerical; scaling factor.                            |

### Author(s)

Diego Diez

### Examples

```
## Not run:
    data(codelink.example)
    arrayNew()
    imageCodelink(codelink.example)
```

## End(Not run)

arraySize

Determine the size of the array

# Description

When loading Codelink arrays in text format (as exported from the Codelink software) this function retrieves the correct size of the array. This is useful beacuse those files contain and indetermined number of empty lines at the end. Thus, reading the entire data matrix don't works.

#### Note

Not meant to be used directly.

#### Author(s)

Diego Diez

as.matrix.Codelink Return a matrix of intensity values

### Description

Takes a Codelink object and returns a matrix with the intensity values available.

# Usage

```
## S3 method for class 'Codelink':
as.matrix(x, ...)
```

### Arguments

| Х | an object of class "Codelink".                                |
|---|---|
|   | additional arguments added to generic as.matrix since R-2.5.x |

# Value

A matrix with the intensity values.

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averageProbes

#### Author(s)

Diego Diez

### See Also

as.matrix

#### **Examples**

```
## Not run:
data(codelink.example)
mat <- as.matrix(codelink.example)</pre>
is(mat)
## End(Not run)
```

*averageProbes* averageProbes

### Description

Takes a CodelinkSet object and computes the average (mean) and sd of duplicated probes.

### Usage

```
averageProbes(object, parallel = FALSE)
```

#### Arguments

| object   | an object of class "CodelinkSet".                               |
|----------|---|
| parallel | whether to use a parallel version (requires package multicore). |

#### Details

This function will compute the mean() and sd() on each duplicated probe (i.e. identical probe id as for probeNames(). CodelinkSet objects use the per-array index to enable unique identifiers needed for ExpressionSet derived objects. Although the method probeNames() provides access to Codelink probe ids, this is inconvenient when dealing with other packages that make use of featureNames() to obtain probe ids and feed them to the corresponding annotation package.

In such cases CodelinkSet objects are not compatible with methods working on ExpressionSet objects. To avoid this limitation it is possible to construct a CodelinkSet object containing unique identifiers (CodelinkSetUnique class) by averaging the intensities of all replicated probes. This is done by computing the mean(). The sd() is also computed and stored in the slot sd.

The current implementation takes a lot of time so a parallelized version of lapply() may be used through the package 'multicore'. This is controlled by the argument 'parallel' which is FALSE by default.

According to the authors it is not actually possible to use 'multicore' in a GUI environment like the R.app Cocoa application in R, because this interferes with the events loop. Therefore it is adviced to use option 'parallel=TRUE' in an R session running in a shell.

### Author(s)

Diego Diez

### Examples

```
## End(Not run)
```

bkgdCorrect Background correction of intensity values.

# Description

Takes a Codelink object with Spot mean and Bkgd median values and corrects applying one of the methods available.

#### Usage

```
bkgdCorrect(object, method = "half", preserve = FALSE, verbose = FALSE,
offset = 0)
```

#### Arguments

| object   | character; an object of class "Codelink".   |
|----------|---|
| method   | character; the correction method to use, one of "none", "subtract", "half" and "normexp". |
| preserve | logical; if Smean and Bmedian slots should be preserved.                                  |
| verbose  | logical; if TRUE print some information with method normexp.                              |
| offset   | numeric; value to add to intensities.   |

### Details

Available methods are: . none: left intensities untouched. . subtract: simple subtraction of Bkgd median from Spot mean. . half: the same as above but aboid negative values setting all intensity values below zero to 0.5. . normexp: apply normexp background adjustment from package limma.

#### Value

An object of class Codelink with corrected intensity values, that is Ri slot.

#### Author(s)

Diego Diez

### codelink.example

### Examples

```
## Not run:
    data(codelink.example)
    codelink.example <- bkgdCorrect(codelink.example, method = "half")</pre>
```

## End(Not run)

codelink.example Dataset of class 'Codelink'

#### Description

Dataset from a h20kcod (Codelink Human UniSet I 20k) array containing 2 samples and ~20000 probes.

### Usage

```
data(codelink.example)
```

## Format

A Codelink object containing 20469 probes and 2 samples.

#### Author(s)

Diego Diez

codelink.exprset Dataset of class 'CodelinkSet'

### Description

Dataset from a h20kcod (Codelink Human UniSet I 20k) array containing 4 samples and ~20000 probes.

# Usage

```
data(codelink.example)
```

### Format

A Codelink object containing 20469 probes and 4 samples.

### Author(s)

Diego Diez

createWeights

### Description

Create a weight matrix to be used in limma.

### Usage

```
createWeights(object, type=NULL)
```

#### Arguments

| object | an object of class "Codelink".       |
|--------|--------------------------------------|
| type   | weight assigned to each Probe\_type. |

# Author(s)

Diego Diez

### Examples

```
## Not run:
    data(codelink.example)
    w <- createWeights(codelink.example, type = list(FIDUCUAL = 0.1))
## End(Not run)
```

cutCV

Calculate cutoff based in CV.

### Description

Takes a Codelink object and calculate cutoff based in CV.

### Usage

```
cutCV(object, subset=c(1:dim(object)[2]))
```

### Arguments

| object | an object of class "Codelink".            |
|--------|---|
| subset | subset of arrays to calculate cutoff with |

### Details

First it computes the median of CV for each gene over all arrays. Then it computes the mean and sd of all medians. Finally:

cutoff = mean + 3 \* sd

#### decDetect

### Author(s)

Diego Diez

### Examples

```
## Not run:
    # data: Normalized Codelink object merged.
    cutoff <- cutCV(data)
## End(Not run)
```

decDetect

Determine decimal type of Codelink files

# Description

Determine decimal type of Codelink files.

# Usage

```
decDetect(file, nlines)
```

### Arguments

| file   | the file to be read.     |
|--------|--------------------------|
| nlines | number of lines to skip. |

#### Value

Decimal type.

### Author(s)

Diego Diez

dim.Codelink Return the dimension of a Codelink object.

### Description

Takes a Codelink object and returns the dimension (genes x samples).

#### Usage

```
## S3 method for class 'Codelink':
dim(x)
```

# Arguments

x an object of class "Codelink".

### Value

A numeric vector with the dimensions.

### Author(s)

Diego Diez

### Examples

```
## Not run:
data(codelink.example)
dim(codelink.example)
```

## End(Not run)

fc2Cond

Select probes based on fold change calculation

### Description

Takes a Codelink object and calculate fold changes (M) between two conditions (samples). Then select genes based on those who pass the passed cutoff.

# Usage

```
fc2Cond(object, cond1=NULL, cond2=NULL, fc=1.0, verbose=FALSE)
```

### Arguments

| object  | an object of class "Codelink".                          |
|---------|---|
| cond1   | numeric or character; First condition to compute M.     |
| cond2   | numeric or character; Second condition to compute M.    |
| fc      | value of the fold change cutoff                         |
| verbose | logical; if some information is printed on the console. |

# Details

Conditions can be passed as characters or as numeric index from the sample slot. The intensities are internally transformed to log2 if needed. The M value is computed as:

M = cond1 - cond2

### Value

A logical vector indicating which genes pass the cutoff

### Author(s)

Diego Diez

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imageCodelink Image plot of Codelink arrays

# Description

Plot and image of a Codelink array if the layout information is found.

# Usage

# Arguments

| object | an object of class "Codelink".                          |
|--------|---|
| array  | array to be used.                                       |
| what   | with data plot: bg, smean, ri, ni.                      |
| low    | color used for low intensities.                         |
| high   | color used for high intensities-                        |
| mar    | character vector specifiying margings.                  |
| gc     | numerical; number of grid columns.                      |
| gr     | numerical; number of grid rows.                         |
| log.it | logical; if TRUE data is log2 transformed (if not yet). |
|        | additional arguments passed to image.                   |

# Author(s)

Diego Diez

# Examples

```
## Not run:
    data(codelink.example)
    imageCodelink(codelink.example)
```

## End(Not run)

logCodelink

#### Description

Takes a Codelink object and apply logCodelink to intensity values.

### Usage

```
logCodelink(object)
```

### Arguments

object an object of class "Codelink" or a list of genes.

# Value

A Codelink object with logCodelink intensities.

#### Author(s)

Diego Diez

### See Also

log2

### Examples

```
## Not run:
    data(codelink.example)
    codelink.example <- logCodelink(codelink.example)
## End(Not run)
```

mergeArray Merge Codelink Bioarrays Data

# Description

Merge data in a Codelink Object corresponding to same samples. Need a vector indicating the classes and an optional vector indicating the labels of the mer- ged samples.

#### Usage

#### na2false

### Arguments

| object    | an object of class "Codelink".                                   |
|-----------|--|
| class     | a numeric vector indicating the classes.                         |
| names     | an optional character vector indicating labels for each class.   |
| method    | the method used to summarize. Currently only "mean" supported.   |
| log.it    | logical; a logical indicating if log2 values should be returned. |
| merge.snr | logical; a logical indicating if SNR values should be merged.    |

### Value

An object of class "Codelink".

# Author(s)

Diego Diez

# Examples

```
## Not run:
data(codelink.example)
codelink.example <- bkgdCorrect(codelink.example)
codelink.example <- normalize(codelink.example, log.it = FALSE)
codelink.example <- mergeArray(codelink.example, class = c(1,1),
names = "SAMPLE", log.it = TRUE)
```

## End(Not run)

na2false Set NAs to FALSE

# Description

Takes a logical vector as input and set all NAs to FALSE. This may happens when comparison is done on NA values.

### Usage

na2false(x)

# Arguments

x a logical vector.

### Value

A logical vector without NAs.

### Author(s)

Diego Diez

normalize

### Examples

```
## Not run:
    a <- c(1, 2, 3, NA, 5)
    b <- c(5, 4, NA, 2, 1)
    sel <- a > b
    sel <- na2false(sel)
## End(Not run)
```

normalize Normalization wrapper for Codelink objects.

# Description

Takes a Codelink object and applies normalization to intensity values.

# Usage

```
normalize(object, method = "quantiles", log.it = TRUE, preserve = FALSE,
weights = NULL, verbose = FALSE)
```

### Arguments

| object   | an object of class "Codelink".           |
|----------|--|
| method   | method to use in normalization.          |
| log.it   | logical; if data should be log2.         |
| preserve | logical; if Ri slot should be preserved. |
| weights  | weights vector for method CyclicLoess.   |
| verbose  | should informative output be printed.    |

### Details

Currently supported methods include "loess", "quantiles" and "median". Median normalization is analogous to the default method applied for the manufacturer in the Codelink software. Loess is a modified version of CyclicLoess implemented in the affy package, allowing missing values and weights. Quantile normalization uses the normalizeQuantiles() function in the limma package.

# Value

A Codelink object with normalized intensity values.

### Author(s)

Diego Diez

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#### normalize.loess

### Examples

normalize.loess Normalized chips using cyclic loess.

### Description

Takes a matrix and apply cyclic loess normalization. It is based in normalize.loess from package affy but supports NA.

### Usage

```
normalize.loess(mat, subset = sample(1:(dim(mat)[1]),
    min(c(5000,nrow(mat)))), epsilon = 10^-2, maxit = 1, log.it = TRUE,
verbose = FALSE, span = 2/3, family.loess = "symmetric", weights = NULL)
```

### Arguments

| mat          | a matrix with columns containing the values of the chips to normalize.   |
|--------------|--|
| subset       | a subset of the data to fit a loess to.  |
| epsilon      | a tolerance value (supposed to be a small value - used as a stopping criterium).                               |
| maxit        | maximum number of iterations.  |
| log.it       | logical. If TRUE it takes the log2 of mat  |
| verbose      | logical. If TRUE displays current pair of chip being worked on.  |
| span         | parameter to be passed the function loess  |
| family.loess | parameter to be passed the function loess. "gaussian" or "symmetric" are acceptable values for this parameter. |
| weights      | a vector of weights for the individual measurements.   |

# Value

A matrix of normalized values.

#### Author(s)

Diego Diez

# Examples

```
## Not run:
mat <- matrix(sample(500), 100, 5)
mat <- normalize.loess(mat)
## End(Not run)
```

plotCV

Plot of CV

### Description

Takes a Codelink object and plot de distribution of CV after applying mergeCodelink.

### Usage

```
plotCV(object, subset=c(1:dim(object)[2]), cutoff=NULL, title=NULL, legend.cex=1
```

#### Arguments

| object     | an object of class "Codelink".                     |
|------------|--|
| subset     | subset of arrays to plot                           |
| cutoff     | cutoff of CV to be shown.                          |
| title      | title of the plot.                                 |
| legend.cex | factor to apply to the fonts in the legend to fit. |

### Author(s)

Diego Diez

plotCorrelation *Plot correlation scatterplot between two arrays* 

#### Description

Takes a Codelink object as argument and plot Correlation scatterplot of two arrays.

# Usage

```
plotCorrelation(object, x=1, y=2, cutoff=FALSE, label="type", title=NULL, xlim=N
```

# Arguments

| object | an object of class "Codelink".   |
|--------|----------------------------------|
| Х      | array to be used in x axis.      |
| У      | array to be used in y axis.      |
| cutoff | cutoff used to show fold change. |
| label  | labels to shown.                 |
| title  | The title of the plot.           |
| xlim   | range for the X axis.            |
| ylim   | range for the Y axis.            |
|        |                                  |

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#### plotDensities

### Author(s)

Diego Diez

### See Also

plot

# Examples

```
## Not run:
    data(codelink.example)
    plotCorrelation(codelink.example)
```

```
## End(Not run)
```

plotDensities Plot Densities

# Description

Takes a Codelink object and plot the distributions of intensities.

#### Usage

### Arguments

| object     | an object of class "Codelink".                               |
|------------|--|
| subset     | subset of arrays to be plotted (default: all).               |
| title      | title of the plot.   |
| legend.cex | font factor use in legend to fit.                            |
| what       | what data to plot, may be "bg", "smean", "snr", "ri" or "ni" |

#### Author(s)

Diego Diez

# Examples

```
## Not run:
    data(codelink.example)
    plotDensities(codelink.example)
```

## End(Not run)

plotMA

MA plot

### Description

Takes a Codelink object and plot M vs A.

# Usage

```
plotMA(object, array1 = 1, array2 = NULL, cutoff = c(-1, 1), label = NULL,
    type = NULL, high.list = NULL, high.col = "blue", high.pch = 21,
    high.bg = "cyan",snr = NULL, snr.cutoff = 1, legend.x = NULL, pch = ".",
    subset = NULL, title = NULL, xlim = NULL, ylim = NULL)
```

#### Arguments

| array1first array to be used.array2second array to be used.cutoffcutoff to be used as fold change markeer.labeltype of labeling used in legend.typespot type information.high.listlist of genes highlighted.high.colcolor used for high genes.high.pchpch used for high genes.snrbackground color used for high genes.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchsubset of spots used to plot based on 'type' slot.titlesubset of spots used to plot based on 'type' slot.titlerange for the X axis.ylimrange for the Y axis. | object     | an object of class "Codelink" or "MAarrayLM".                         |
|---|------------|---|
| cutoffcutoff to be used as fold change markeer.labeltype of labeling used in legend.typespot type information.high.listlist of genes highlighted.high.colcolor used for high genes.high.pchpch used for high genes.high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.       | arrayl     | first array to be used.   |
| labeltype of labeling used in legend.typespot type information.high.listlist of genes highlighted.high.colcolor used for high genes.high.pchpch used for high genes.high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsettitle of the plot.xılimrange for the X axis.  | array2     | second array to be used.  |
| typespot type information.high.listlist of genes highlighted.high.colcolor used for high genes.high.pchpch used for high genes.high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.   | cutoff     | cutoff to be used as fold change markeer.                             |
| high.listlist of genes highlighted.high.colcolor used for high genes.high.pchpch used for high genes.high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.   | label      | type of labeling used in legend.                                      |
| high.colcolor used for high genes.high.pchpch used for high genes.high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | type       | spot type information.  |
| high.pchpch used for high genes.high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | high.list  | list of genes highlighted.  |
| high.bgbackground color used for high genes.snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | high.col   | color used for high genes.  |
| snrvector with SNR values, usually, taking rowMeans() from a SNR matrix.snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | high.pch   | pch used for high genes.  |
| snr.cutoffSNR cutoff used for label spots.legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | high.bg    | background color used for high genes.                                 |
| legend.xrelative position of the legend.pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | snr        | vector with SNR values, usually, taking rowMeans() from a SNR matrix. |
| pchpch style used to main spots.subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | snr.cutoff | SNR cutoff used for label spots.                                      |
| subsetsubset of spots used to plot based on 'type' slot.titletitle of the plot.xlimrange for the X axis.  | legend.x   | relative position of the legend.                                      |
| titletitle of the plot.xlimrange for the X axis.  | pch        | pch style used to main spots.   |
| xlim range for the X axis.  | subset     | subset of spots used to plot based on 'type' slot.                    |
| -   | title      | title of the plot.  |
| ylim range for the Y axis.  | xlim       | range for the X axis.   |
|   | ylim       | range for the Y axis.   |

### Details

This function has suffered recent re-working, to increase the usability and to clean a little bit the code.

If array2 is NULL a median array is computed using all available arrays. Then the values of M and A are computed using the following formula:

M = array2 - array1

A = (array2 + array1) / 2

If type information is available in the Codelink object, or provided throught the 'type' argument, spots are colored based on that. DISCOVERY spots are plotted black with pch = "." whereas the

### printHead

other classes are plotted with different background colors, using gray as border to increase contrasts. For that pch = 21 is used. If snr is specified as label option, the SNR is used to label spots, if available in the Codelink object. In this case, the mean SNR across all arrays is used when array2 = NULL.

Some parameters may not be working right now, as the new function is using a different method to labels spots.

The legend is 'automagically' located, but this can be overrided with the legend.x argument.

In addition, a subset of the spots can be plotted based on type information when available. This allows, for example, to plot only DISCOVERY spots.

### Author(s)

Diego Diez

#### Examples

```
## Not run:
    data(codelink.example)
    plotMA(codelink.example)
```

```
## End(Not run)
```

printHead

Print briefly a Codelink object

### Description

Takes a Codelink object and print a summary information of the data estored. It is based on print-Head() from package limma.

#### Usage

```
printHead(x)
```

#### Arguments

x an object of class "Codelink".

### Author(s)

Diego Diez

readCodelink

### Description

Read data exported as text by Codelink Software. It reads values (normalized by Codelink Software or not) flags and information about probes.

# Usage

```
readCodelink(files=list.files(pattern="TXT"), sample.name=NULL,
flag, dec=NULL, type="Spot",preserve=FALSE,verbose=2,
file.type="Codelink", check=TRUE, fix=FALSE)
```

### Arguments

| files       | list of files to read.   |
|-------------|--|
| sample.name | vector of same length as files with sample names.  |
| flag        | list with values to assign based on Flag quality values.   |
| dec         | character indicating the decimal character used in the files.  |
| type        | character indicating which base value to read from files.  |
| preserve    | logical, if TRUE Bkgd\_stdev slot is not removed (if present).   |
| verbose     | numerical, set the level of information. Level 2 set as old behaviour. Level > 2 output some debug info. |
| file.type   | exported file type, currently Codelink or XLS file formats supported.                                    |
| check       | logical, check for probe order consistency.  |
| fix         | logical, try to fix probe order consistency.   |

### Value

An object of class "Codelink".

### Author(s)

Diego Diez

### See Also

read.table

### Examples

```
## Not run:
    # reading default extension (TXT).
    data <- readCodelink()
    # specify a different one.
    files <- list.files(pattern = "txt")
    data <- readCodelink(files = files)</pre>
```

#### readHeader

```
# example.
data(codelink.example)
```

```
## End(Not run)
```

readHeader

Read Header from Codelink Bioarrays Files

### Description

Read the header of Codelink files and obtein usefull information.

#### Usage

readHeader(file, dec=FALSE)

# Arguments

| file | File to read.                             |
|------|---|
| dec  | logical; If TRUE determine decimal point. |

### Value

A list with header and other usefull information.

### Author(s)

Diego Diez

### Examples

```
## Not run:
    files <- list.files(pattern = "TXT")
    head <- readHeader(files[1])
## End(Not run)
```

readHeaderXLS Read Header from XLS exported Codelink Bioarrays Files

# Description

Read the header of Codelink files and obtain useful information.

# Usage

```
readHeaderXLS(file, dec=FALSE)
```

### Arguments

| file | File to read.                             |
|------|---|
| dec  | logical; If TRUE determine decimal point. |

### Details

This function is not meant to be used by normal users.

# Value

A list with header and other usefull information.

### Author(s)

Diego Diez

### Examples

```
## Not run:
    files <- list.files(pattern = "TXT")
    head <- readHeaderHeader(files[1])</pre>
```

```
## End(Not run)
```

reportCodelink Write a report of genes selected in HTML

### Description

Takes a list of genes as argument and writes an HTML page containing informa- tion about these genes: Unigene, Genbank, Entrez Gene, etc.

### Usage

# Arguments

| object     | an object of class "Codelink" or a list of genes.                 |
|------------|---|
| chip       | the chip description package.                                     |
| filename   | file name used in the report.                                     |
| title      | title used in the report.   |
| probe.type | logical; if TRUE Probe type information is writen.                |
| other      | list of vectors containing aditional values to add to the report. |
| other.ord  | slot name in other to order genes by.                             |

# Value

Nothing, only the HTML file generated.

### selCV

### Author(s)

Diego Diez

# See Also

htmlpage

```
selCV
```

Select based on CV cutoff.

# Description

Takes a Codelink object and select genes based on CV cutoff.

# Usage

selCV(object, cutoff)

### Arguments

| object | an object of class "Codelink".         |
|--------|--|
| cutoff | cutoff normaly calculated with cutCV() |

#### Value

A logical vector.

# Author(s)

Diego Diez

writeCodelink Write a Codelink object to file.

# Description

Export of the data from a codelink object to a text file.

# Usage

```
writeCodelink(object, file, dec = ".", sep = "\t", flag = FALSE, chip)
```

# Arguments

| object | an object of class "Codelink".         |
|--------|--|
| file   | filename to write object to.           |
| dec    | decimal character to use.              |
| sep    | delimiter character to use.            |
| flag   | should the Codelink flags be written.  |
| chip   | chip package to use, normally guessed. |

### Details

By default, intensities and SNR are wrote to the file. If set, the flag are also output. The header have "INTENSITY\\_", "SNR\\_" and "FLAG\\_" respectibely appendend to the sample name for those values. The default delimiter is the tab character, but that can be set with the sep argument. The default decimal character is the point.

## Author(s)

Diego Diez

# Examples

```
## Not run:
    data(codelink.example)
    writeCodelink(codelink.example, file = "foo.txt")
## End(Not run)
```

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