

Package ‘convert’

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Title Convert Microarray Data Objects

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Depends R (>= 2.6.0), Biobase (>= 1.15.33), limma (>= 1.7.0), marray, utils, methods

Description Define coerce methods for microarray data objects.

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URL <http://bioinf.wehi.edu.au/limma/convert.html>

biocViews Infrastructure, Microarray, TwoChannel

R topics documented:

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coerce

Description

Objects can be converted (coerced) from one class to another using `as(object, Class)` where `object` is an object to convert and `Class` is the name of the class to convert to. The following

From:	To:
RGLList	marrayRaw
marrayRaw	RGLList
MAList	marrayNorm
marrayNorm	MAList
RGLList	NChannelSet
marrayRaw	NChannelSet
MAList	ExpressionSet
marrayNorm	ExpressionSet

RGLList and marrayRaw are coerced to NChannelSet. Channel values are not transformed.

MAList and marrayNorm are coerced so that the ExpressionSet slot contains log-ratios (M-values) and the ExpressionSet object has the same number of columns as the original object. In this case, information on the A-values is lost.

There is intentionally no conversion from RGLList or marrayRaw to ExpressionSet, as ExpressionSet is intended for expression values, not intensities.

Author(s)

Gordon Smyth and others

See Also

[as](#) in the methods package.

Examples

```
##first set up some fake intensity matrices
testRed <- matrix(rnorm(5*2),5,2,
  dimnames=list(paste("gene",1:5, sep=""), c("S1", "S2")))
testGreen <- matrix(rnorm(5*2),5,2,
  dimnames=list(paste("gene",1:5, sep=""), c("S1", "S2")))

##some sample/target info
testTarget <- data.frame(slide=c("S1", "S2"), Cy3=c("T", "C"),
  Cy5=c("C", "T"), row.names=c("S1", "S2"))

maT <- new("marrayInfo", maLabels=c("S1", "S2"),
  maInfo= testTarget)

##now create instances and convert
x <- new("RGLList")
x$R <- testRed
x$G <- testGreen
y <- as(x, "marrayRaw")
z <- as(x, "NChannelSet")

x <- new("marrayRaw")
x@maGf <- testGreen
```

```
x@maRf <- testRed
x@maTargets = maT
y <- as(x, "RGList")
z <- as(x, "NChannelSet")

x <- new("MAList")
y <- as(x, "marrayNorm")

##we construct a reasonably complete fake, small
##instance of the marrayNorm class
x <- new("marrayNorm")
x@maM <- testRed
x@maA <- testGreen
maTargets(x) = maT
y <- as(x, "MAList")
y <- as(x, "ExpressionSet")

x <- new("MAList")
x$M <- testRed
x$A <- testGreen
x$targets <- testTarget
y <- as(x, "ExpressionSet")
```

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