

logitT

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logitTAffy

Testing for differential gene expression using the Logit-t algorithm

Description

This function takes an instance of AffyBatch and calculates t-statistics for tests of differential gene expression for oligonucleotide arrays using the Logit-t algorithm.

Usage

```
logitTAffy(object, group)
```

Arguments

object	an instance of AffyBatch
group	a vector specifying the group label for each array

Details

For more details see the package vignette.

Value

A named vector containing the t-statistics for each probe set for each array.

Author(s)

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References

William J Lemon, Sandya Liyanarachchi and Ming You (2003). A high performance test of differential gene expression for oligonucleotide arrays. *Genome Biology* 2003, 4:R67. <http://genomebiology.com/2003/4/10/R67>

See Also

[AffyBatch](#)

Examples

```
if(require(SpikeInSubset)) {
  library(SpikeInSubset)
  data(spikein95)
  logitTex<-logitTAffy(spikein95, group=c("A", "A", "A", "B", "B", "B"))
  logitTex[1:10]                                     # extract t-s
  logitTex[grep("AFFX-BioB-5_at", names(logitTex))] # extract t-stat
  pvals<-(1-pt(abs(logitTex), df=4))*2           # calculate t-
  signifgenes<-names(logitTex) [pvals<0.01]        # find signif
} else{
  stop("Please install the SpikeInSubset package to run the example.")
}
```

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