# Package 'davidTiling'

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| <b>Version</b> 1.49.0   |
| <b>Title</b> Data and analysis scripts for David, Huber et al. yeast tiling array paper   |
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| <b>Depends</b> R (>= 2.10), Biobase (>= 2.5.5), tilingArray, GO.db  |
| Description This package contains the data for the paper by L. David et al. in PNAS 2006 (PMID 16569694): 8 CEL files of Affymetrix genechips, an ExpressionSet object with the raw feature data, a probe annotation data structure for the chip and the yeast genome annotation (GFF file) that was used. In addition, some custom-written analysis functions are provided, as well as R scripts in the scripts directory. |
| <b>biocViews</b> ExperimentData, Genome, Saccharomyces_cerevisiae_Data, MicroarrayData, ReproducibleResearch  |
| Reference A high-resolution map of transcription in the yeast genome.  David L, Huber W, Granovskaia M, Toedling J, Palm CJ, Bofkin L,  Jones T, Davis RW, Steinmetz LM. Proc Natl Acad Sci U S A. 2006  Apr 4;103(14):5320-5.  |
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| Contents  |
| data:davidTiling       2         data:gff       3         data:probeAnno       3         getAllGO       5         getAttributeField       5         GOHyperG       6  |

2 data:davidTiling

| scatterWithHist |  | • |  | • |  |  | • | • |  | <br> | • | • | • |  |  |  |  | • | • | • | • | • |  |
|-----------------|--|---|--|---|--|--|---|---|--|------|---|---|---|--|--|--|--|---|---|---|---|---|--|
| scoreSegments   |  |   |  |   |  |  |   |   |  | <br> |   |   |   |  |  |  |  |   |   |   |   |   |  |
| showDens        |  |   |  |   |  |  |   |   |  | <br> |   |   |   |  |  |  |  |   |   |   |   |   |  |
| yeastFeatures . |  |   |  |   |  |  |   |   |  | <br> |   |   |   |  |  |  |  |   |   |   |   |   |  |

Index 11

data:davidTiling

Dataset of class eSet with the raw 'CEL file' intensities

## Description

The data are from an experiment that used Affymetrix Scerevisiaetiling chips from 2004, which where custom-made for the Stanford Genome Center. The chips tile the complete genome of S. cerevisiae in steps of 8 bases, separately for each strand of each chromosome. The two tiles for one chromosome (Watson and Crick strands) are offset by 4 bases.

Note that the class eSet was used instead of AffyBatch since the additional overhead of 'CDF environments' in the latter is not needed here.

## Usage

```
data("davidTiling")
```

#### **Format**

Intensity data for 8 arrays. The phenoData slot contains the file names and the nucleic acid type.

## Author(s)

W. Huber <huber@ebi.ac.uk>

#### Source

Lior David and Lars Steinmetz, both from the Stanford Genome Center. Lars Steinmetz is also at EMBL Heidelberg.

## **Examples**

```
data("davidTiling")
dim(exprs(davidTiling))
```

data:gff 3

| data:gff | Genomic features of Saccharomyces cerevisiae |  |
|----------|--|--|
| S        | y y  |  |

## **Description**

A data frame with genomic features of Saccharomyces cerevisiae.

## Usage

```
data("gff")
```

## **Format**

Object of class data.frame. GFF is a file format for annotating genomes, see <insert the URL to the documentation page for GFF at Sanger here>. The format is essentially a rectangular table, and here it is represented as a data frame.

#### Author(s)

W. Huber <huber@ebi.ac.uk>

#### **Source**

Two GFF files were downloaded: saccharomyces\_cerevisiae.gff from ftp://genome-ftp.stanford.edu/pub/yeon 7 Aug 2005, 18:16 BST, and IGR\_v24.2.p001.allowoverlap.GFF from http://jura.wi.mit.edu/fraenkel/dow upon suggestion from the SGD curators on 30 Aug 2005. (Future versions of SGD's GFF files are likely to include the latter as well). They were parsed, combined and written into the gff data.frame with the script makeProbeAnno.R in the inst/scripts directory of this package.

## **Examples**

```
data("gff")
str(gff)
```

| data:probeAnno | An environment with probe mapping information for the Scerevisiae tiling array |
|----------------|--|
|                | uung array   |

## **Description**

The environment contains probe mapping information for the Affymetrix Scerevisiaetiling chip from 2004, which was custom-made for the Stanford Genome Center. The chips tile the complete genome of S. cerevisiae in steps of 8 bases, separately for each strand of each chromosome. The two tiles for one chromosome (Watson and Crick strands) are offset by 4 bases.

In the following a brief description of the 138 elements of the probeAnno environment.

probeReverse: a list of 8 factors, each of length 6553600, corresponding to the rows of davidTiling. For example, if the probe corresponding to the j-th row in davidTiling maps to the coding sequence of a gene, then the factor level of probeRevers\$CDS[j] is the name of the gene, and the

4 data:probeAnno

empty string "" otherwise. This applies to samples that were hybridized to the chip after a reverse transcription step.

probeDirect: analogous to probeReverse, but for samples that were hybridized to the chip without a reverse transcription step. The probes map to the opposite chromosomal strand compared to experiments with reverse transcription.

- 1.+. index: indices (from 1...6553600, corresponding to the rows of davidTiling) of probes mapping to the Watson strand of chromosome 1.
- 1.+.start, 1.+.end: start and end positions in genomic coordinates of the alignments of the probes (in the same order as in 1.+.index) to the Watson strand of chromosome 1. For 25-mers, the values in 1.+.end are those in 1.+.start plus 24, but not all probes on the array are 25-mers.
- 1.-. unique: specificity of the probe:
- 0 has exactly one perfect match (PM) and no near-matches in the genome
- 1 has exactly one PM and some near-matches
- 2 has no PM but one or more near-matches
- 3 has multiple PMs in the genome
- 1.-.index, 1.-.start, 1.-.end, 1.-.unique: analogous to the above, but for the Crick strand of chromosome 1. 2.+.index, 2.+.start, 2.+.end, 2.-.unique: analogous to the above, but for the Watson strand of chromosome 2; and so forth. "Chromosome 17" is mitochondrial DNA.

## Usage

```
data("probeAnno")
```

## Author(s)

W. Huber <huber@ebi.ac.uk>

#### Source

Probe sequences were obtained from Affymetrix in a file called S.cerevisiae\_tiling.1lq. The genomic sequences of the S.cerevisiae chromosomes were downloaded from ftp://genome-ftp.stanford.edu/pub/y on 7 Aug 2005, 18:16 BST in 17 files chr01.fsa—chr16.fsa, and chrmt.fsa. The probe sequences were matched against the chromosomal sequencences with the program MUMmer, see the script mapProbesToGenome.sh (in the inst/scripts directory of this package). MUMmer results were parsed and processed into the probeAnno environment with the script makeProbeAnno.R (in the inst/scripts directory of this package).

## **Examples**

```
data("probeAnno")
ls(probeAnno)
str(probeAnno$"1.+.start")
```

getAlIGO 5

| getAllGO | Get all GO categories for a list of genes. |
|----------|--|
|          |  |

## **Description**

The function uses the GO categories in the data.frame gff to obtain annotated GO categories, then the GO\*\*ANCESTOR data in the GO package to add all parent terms as well.

## Usage

```
getAllGO(x, gff)
```

## **Arguments**

x character vector.

gff data.frame with columns feature, Name, and (Ontology\_term or attributes),

see details

## **Details**

The elements of x are matched against the column gene in gff. All are required to match. A list of GO terms is then extracted from the corresponding rows in the Ontology\_term column. A gene may be annotated by several terms, separated by ",". Then the GO package is used to augment this by all ancestor terms.

## Value

List of character vectors.

## Author(s)

W. Huber <huber@ebi.ac.uk>

| getAttributeField | Extract the value of a certain field out of a character vector such as in |
|-------------------|---|
|                   | the "attributes" column of a GFF table.                                   |

## **Description**

Extract the value of a certain field out of a character vector such as in the "attributes" column of a GFF table.

## Usage

```
getAttributeField(x, field, attrsep=";")
```

## **Arguments**

field character vector of length 1, containing the field name.

attrsep character vector of length 1, containing the separator name.

6 GOHyperG

## **Details**

See example.

## Value

Character vector.

## Author(s)

W. Huber <huber@ebi.ac.uk>

## **Examples**

```
acol = c("ID=46891;Name=TEL01L-TR;Note=Bla",
   "ID=46892;Name=TEL01L;Note=Di",
   "ID=46893;Name=TEL01L-XR;Note=Bla")
getAttributeField(acol, "Name")
getAttributeField(acol, "ID")
```

GOHyperG

Hypergeometric test for GO category enrichment.

## **Description**

This function is adapted from the function of the same name in GOstats package. Main difference is that it draws the GO annotations of the genes in candidates from the data.frame gff. It also draws a plot.

## Usage

```
GOHyperG(candidates, gff, plotmain)
```

## **Arguments**

candidates character vector

gff see getAllGO

plotmain character, plot title

## **Details**

The elements of x are matched against the column gene in gff. All are required to match. A list of GO terms is then extracted from the corresponding rows in the Ontology\_term column. A gene may be annotated by several terms, separated by ",". Then the GO package is used to augment this by all ancestor terms.

## Value

List of character vectors.

scatterWithHist 7

## Author(s)

W. Huber <huber@ebi.ac.uk>

## See Also

```
getAllGO
```

scatterWithHist

Scatterplot with histograms of marginal distributions

## Description

Scatterplot with histograms of marginal distributions.

## Usage

```
scatterWithHist(x, breaks, barcols, xlab, ylab, ...)
```

## Arguments

```
x numeric matrix with 2 columns.
breaks numeric vector with histogram breaks, see hist.
barcols character vector of length 2, colors for the histogram filling.
xlab characterof length 1, label for x-axis.
ylab characterof length 1, label for y-axis.
... further arguments that get passed on to plot.
```

## Value

The function is called for its side effect.

## Author(s)

W. Huber <huber@ebi.ac.uk>

## Examples

```
x = rexp(100)
x = cbind(x, x+0.6*rnorm(length(x)))
scatterWithHist(x,
    breaks=seq(min(x),max(x),length=20),
    barcols=c("mistyrose", "lightblue"),
    xlab="Daffodil", ylab="Petunia", pch=16)
```

8 showDens

|--|

## Description

Score the segments found by a previous call to findSegments by comparing to genome annotation

## Usage

```
scoreSegments(s, gff,
  nrBasePerSeg = 1500,
  probeLength = 25,
  params = c(overlapFraction = 0.5, oppositeWindow = 100, flankProbes=10),
  verbose = TRUE)
```

## **Arguments**

s environment. See details.

gff GFF dataframe.

nrBasePerSeg Numeric of length 1. This parameter determines the number of segments.

probeLength Numeric of length 1.

params vector of additional parameters, see details.

verbose Logical.

#### **Details**

This function scores segments. It is typically called after a *segmentation*. For an example segmentation script, see the script segment.R in the scripts directory of this package. For an example scoring script, which loads the data and then calls this function, see the script scoreSegments.R.

## Value

A dataframe with columns as described in the details section.

## Author(s)

W. Huber <huber@ebi.ac.uk>

showDens Plot function for more than one density

## **Description**

Plot function for more than one density

## Usage

```
showDens(z, breaks, xat, xtickLabels=paste(xat), col, ylab = "", ...)
```

yeastFeatures 9

## **Arguments**

z List: numeric vectors for computing histograms for

breaks

Numeric vector: breaks of the histogram

Numeric vector: where to put the x-axis ticks

xtickLabels

Character vector: what to write underneath them

col Character vector: colours of the histograms

ylab Character scalar: y-axis label

... futher arguments passed on to plot

#### **Details**

•••

## Value

returns scale factor

## Author(s)

Wolfgang Huber <huber@ebi.ac.uk>

#### See Also

hist

## **Examples**

```
showDens(list(x1=runif(100), x2=exp(runif(50))-1, x3=runif(20)),\\ breaks=seq(0, 2, 0.2), xat=seq(0, 2, 0.5), col=rainbow(3), xlab="Random Numbers")
```

yeastFeatures

A data.frame with metadata about the features in SGD GFF files

## **Description**

The rows of the data frame correspond to feature categories, such as "gene", "CDS", "telomere". The column isTranscribed is a logical vector that denotes whether this feature category is considered to be transcribable.

## Usage

data(yeastFeatures)

## **Format**

A data.frame

## Author(s)

W. Huber <huber@ebi.ac.uk>

10 yeastFeatures

## Examples

data(yeastFeatures)

## **Index**

```
* datasets
    data:davidTiling, 2
    data:gff, 3
    data:probeAnno, 3
    yeastFeatures, 9
* hplot
    scatterWithHist, 7
    showDens, 8
* manip
    getAllGO, 5
    {\tt getAttributeField, 5}
    GOHyperG, 6
    {\tt scoreSegments}, 8
AffyBatch, 2
data:davidTiling, 2
data:gff, 3
data:probeAnno, 3
davidTiling, 3, 4
davidTiling (data:davidTiling), 2
eSet, 2
getAllGO, 5, 6, 7
getAttributeField, 5
gff (data:gff), 3
GOHyperG, 6
hist, 7, 9
phenoData, 2
plot, 7
probeAnno (data:probeAnno), 3
scatterWithHist, 7
scoreSegments, 8
showDens, 8
yeastFeatures, 9
```