ACME
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aGFFCalc-class Class "aGFFCalc"

Description

Store results of ACME calculations

Objects from the Class

Objects can be created by calls of the form new("aGFFCalc", ...).

Slots

call: Object of class "call", contains the exact call to do.aGFF.calc, for historical purposes
threshold: Object of class "numeric", the threshold used in the calculation
cutpoints: Object of class "numeric", the data value above which probes were considered positive
vals: Object of class "matrix", equivalent in size to the original data matrix, containing the calculated p-values from the ACME algorithm
annotation: Object of class "data.frame", currently a copy of the original annotation, possibly reordered in chromosome order
data: Object of class "matrix", the original data, possibly reordered
samples: Object of class "data.frame", sample metadata
Extends

Class "aGFF", directly.

Methods

plot signature(x = "aGFFCalc", ask=FALSE): plot the results of an ACME calculation

print signature(x = "aGFFCalc"): brief overview of the object

show signature(object = "aGFFCalc"): brief overview of the object

Author(s)

Sean Davis <sdavis2@mail.nih.gov>

See Also

do.aGFF.calc, aGFF-class

Examples

data(example.agff)
example.agffcalc <- do.aGFF.calc(example.agff, window=1000, thresh=0.9)
example.agffcalc

aGFF-class Class for storing GFF-like data

Description

The GFF format is quite versatile while remaining simple. This class simply stores the annotation associated with a set of GFF files from the same regions of the genome along with some information about the samples from which the data came and the data (from the "score" column of the GFF file) themselves.

Objects from the Class

Objects can be created by calls of the form new("aGFF", ...). Also, the \code{read.resultsGFF()} function returns aGFF objects.

Slots

annotation: Object of class "data.frame" with two columns absolutely necessary, "Chromosone" and "Location". Other columns can be included.

data: Object of class "matrix" of the same number of rows as the annotation slot and the same number of columns as the number of rows in the samples slot, containing data for later analysis

samples: Object of class "data.frame" for describing the samples, one row per sample
do.aGFF.calc

Methods

plot signature(x = "aGFF") to plot a region along the genome.
print signature(x = "aGFF") to display summary of aGFF object
show signature(object = "aGFF") to display summary of aGFF object

Author(s)

Sean Davis

See Also

read.resultsGFF and aGFFCalc-class

Examples

# Load an example
data(example.agff)
example.agff

do.aGFF.calc

Perform ACME calculation

Description

This function performs the moving window chi-square calculation. It is written in C, so is quite fast.

Usage

do.aGFF.calc(x, window, thresh)

Arguments

x An aGFF class object
window An integer value, representing the number of basepairs to include in the windowed chi-square calculation
thresh The quantile of the data distribution for each sample that will be used to classify a probe as positive

Details

A window size on the order of 2-3 times the average size of fragments from sonication, digestion, etc. and containing at least 8-10 probes is the recommended size. Larger size windows are probably more sensitive, but obviously reduce the accuracy with which boundaries of signal can be called.

A threshold of between 0.9 and 0.99 seems empirically to be adequate. If one plots the histogram of data values and there is an obvious better choice (such as a bimodal distribution, with one peak representing enrichment), a more data-driven approach may yield better results.

Value

An object of class aGFFCalc
Author(s)
Sean Davis <sdavis2@mail.nih.gov>

Examples
```r
data(example.agff)
example.agffcalc <- do.aGFF.calc(example.agff, window=1000, thresh=0.9)
example.agffcalc
```

### Description
An aGFF data structure from two Nimblegen arrays, custom tiled to include multiple HOX genes.

### Usage
```r
data(example.agff)
```

### Format
The format is: `chr "example.agff"`

### Source
From Scacheri et al., Plot Genet, 2006. Pubmed ID 16604156

### Examples
```r
data(example.agff)
example.agff
```

---

**findClosestGene**

**Find closest refseq gene**

### Description
This function is used to find the nearest refseq transcript(s) to a point in the genome specified. Note that it is limited to the refseq transcripts listed at genome.ucsc.edu, where this function goes for information.

### Usage
```r
findClosestGene(chrom, pos, genome = "hg17", position = "txStart")
```
findRegions

Arguments

- `chrom` Usually specified like 'chr1', 'chr2', etc.
- `pos` A position in base pairs in the genome
- `genome` Something like 'hg16', 'hg17', 'mm6', etc.
- `position` The location to measure distance from: one of 'txStart', 'txEnd', 'cdsStart', 'cdsEnd'

Details

The first time the function is run, it checks to see if the refflat table for the given genome is present in the package environment. If not, it downloads it to the /tmp directory and gunzips it (using `getRefflat`). It is then stored so that in future calls, there is no re-download required.

Value

A data frame with the gene name, refseq id(s), txStart, txEnd, cdsStart, cdsEnd, exon count, and distance. Note that distance is measured as pos-position, so negative values mean that the point in the gene is to the left of the point specified in the function call (with the p-tel on the left).

Note

The function may return more than one transcript, as several transcripts may have the same start site

Author(s)

Sean Davis <sdavis2@mail.nih.gov>

Examples

```
findClosestGene('chr1',100000000,'hg17')
```

findRegions

Find all regions in data above p-value threshold

Description

After the ACME calculation, each probe is associated with a p-value of enrichment. However, one often wants the contiguous regions associated with runs of p-values above a given p-value threshold.

Usage

```
findRegions(calcobj, thresh = 1e-04)
```

Arguments

- `calcobj` An aGFFCalc object
- `thresh` The p-value threshold
getRefflat

Details

Runs of p-values above the p-value threshold will be reported as one "region". These can be used for downstream analyses, export to browsers, submitted for transcription factor binding enrichment analyses, etc.

Value

A data frame with these columns:

- **Length**: The length of the region in probes
- **TF**: Either TRUE or FALSE; TRUE regions represent regions of enrichment while FALSE regions are the regions between the TRUE regions
- **StartInd**: The starting Index of the region
- **EndInd**: The ending Index of the region
- **Sample**: The sample containing the region
- **Chromosome**: The Chromosome of the region
- **Start**: The starting basepair of the region
- **End**: The ending basepair of the region

Author(s)

Sean Davis <sdavis2@mail.nih.gov>

See Also

do.aGFF.calc, findClosestGene

Examples

data(example.agff)
extample.agffcalc <- do.aGFF.calc(example.agff, window = 1000, thresh = 0.9)
foundregions <- findRegions(example.agffcalc, thresh = 0.001)
foundregions[1:6,]

getRefflat

Get the refflat table from ucsc for the given genome

Description

Fetches the refflat table from ucsc, stores in temp dir and then gunzips it and reads it in.

Usage

getRefflat(genome = "hg17")

Arguments

genome: The genome code from ucsc, like 'hg16', 'mm6', etc.
**read.resultsGFF**

Value

A data frame mirroring the UCSC table structure.

Author(s)

Sean Davis <sdavis2@mail.nih.gov>

References

http://genome.ucsc.edu

See Also

findClosestGene

Examples

```r
rf <- getRefflat('hg17')
```

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**read.resultsGFF**  
Read Nimblegen GFF files

**Description**

A GFF format file is a quite flexible format for storing genomic data. Nimblegen uses these format files as one format for making chip-chip data available. This function reads these files, one per experiment and creates a resulting aGFF-class object.

**Usage**

```r
read.resultsGFF(fnames, path = ".", samples = NULL, notes = NULL, skip = 0, sep = \t, quote = "", ...)```

**Arguments**

- `fnames`: A vector of filenames
- `path`: The path to the filenames
- `samples`: A data.frame containing sample information, one row per sample, in the same order as the files in fnames
- `notes`: A character vector for notes–not currently stored
- `skip`: Number of lines to skip if the file contains a header
- `sep`: The field separator–should be a tab character for gff files, but can be set if necessary.
- `quote`: The text quote character–again not used for gff file, typically
- `...`: ...

**Details**

The output is an aGFF object.
write.sgr

Value
A single aGFF object.

Author(s)
Sean Davis <sdavis2@mail.nih.gov>

References
http://www.sanger.ac.uk/Software/formats/GFF/

See Also
aGFF-class

Examples
```r
datdir <- system.file('extdata',package='ACME')
fnames <- dir(datdir)
example.agff <- read.resultsGFF(fnames,path=datdir)
```

write.sgr

Write Affy IGB .sgr format files

Description
The affy Integrated Genome Browser (IGB) is a powerful, fast browser for genomic data. The file format is simple (three columns: chromosome, location, and score) to generate. This function will write the .sgr files associated with an aGFFCalc object. There will be either one or two files (default two) representing the raw data and the calculated data (which is output as -log10(val) for visualization purposes).

Usage
```
write.sgr(agff, raw = TRUE, vals = TRUE, directory = ".")
```

Arguments
- **agff**: An aGFFCalc object obtained after running do.aGFF.calc
- **raw**: Create a file for the raw data?
- **vals**: Create a file for the calculated p-values?
- **directory**: Give a directory for storing the files

Author(s)
Sean Davis

Examples
```r
data(example.agff)
write.sgr(example.agff)
```
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