# Package 'encoDnaseI'

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<b>Title</b> data provided by UCSC for Cd4 raw measures of DnaseI hypersensitivity
Description data provided by UCSC for Cd4 raw measures of DnaseI hypersensitivity
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Author VJ Carey <stvjc@channing.harvard.edu></stvjc@channing.harvard.edu>
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Suggests GGdata
Maintainer VJ Carey <stvjc@channing.harvard.edu></stvjc@channing.harvard.edu>
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# **R** topics documented:

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# hg18track-class Class "hg18track" container for hg18 annotation found in genome browser track files; class "chrnum" extends numeric for an indexing application on hg18track objects.

## Description

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container for hg18 annotation found in genome browser track files

#### **Objects from the Class**

Objects can be created by calls of the form new("hg18track", assayData, featureData, experimentData, annotatio These are single-sample eSet instances.

Note that demoTrk19 is a restriction of the rawCD4 structure to the interval of chromosome 19 that was assayed in the ENCODE project for DnaseI hypersensitivity.

#### Slots

assayData: Object of class "AssayData" ~~

phenoData: Object of class "AnnotatedDataFrame" ~~

featureData: Object of class "AnnotatedDataFrame" ~~

experimentData: Object of class "MIAME" ~~

annotation: Object of class "character" ~~

.\_\_classVersion\_\_: Object of class "Versions" ~~

#### Extends

Class "eSet", directly. Class "VersionedBiobase", by class "eSet", distance 2. Class "Versioned", by class "eSet", distance 3.

#### Methods

[ signature(x = "hg18track"): select using numeric, logical, or chrnum indices.

**chrnum** signature(object = "hg18track"): extract numeric tokens for chromosome number at which data values are obtained; note that chrnum is also used as name of a class.

**dataVals** signature(object = "hg18track"): actual data values

- **getTrkXY** signature(object = "hg18track", type = "character"): obtain a list with components x, y indicating location and data value respectively; location is within chromosome; default type is 'midpoint' of locations given as intervals
- getTrkXY signature(object = "hg18track", type = "missing"): take default midpoint x
  values corresponding to data values
- rangeLocs signature(object = "hg18track"): if measures from only one chromosome are present, this returns low and high values of chromStart and chromEnd respectively, otherwise error.
- clipTrk signature(obj = "hg18track", low="numeric", hi="numeric", attr="ANY"):
   create a restriction of the track using an interval specification. by default the chromStart fea tureData component is used for coordinates to clip; if attr is non-missing, the featureData
   component named by attr will be used.

initialize signature(.Object = "hg18track"): create a new instance

#### Author(s)

VJ Carey <stvjc@channing.harvard.edu>

### juxtaPlot

#### Examples

```
showClass("hg18track")
data(rawCD4)
rawCD4
rawCD4.chr1 = rawCD4[ chrnum(1), ]
rangeLocs(rawCD4.chr1)
plot(getTrkXY(rawCD4.chr1), ylab="data value", xlab="interval midpt on chr 1" )
c52 = clipTrk(rawCD4[ chrnum(5), ], 1.30e8, 1.33e8 )
plot(getTrkXY(c52))
```

juxtaPlot

two-panel plot with track info and snp screen t-values

### Description

two-panel plot with track info and snp screen t-values

#### Usage

juxtaPlot(trk, ssr, locstr)

#### Arguments

trk	instance of hg18track
ssr	instance of GGtools snpScreenResult
locstr	matrix with 2 rows: rsid (numeric component of dbSNP id) and loc

#### Details

xyplot of lattice package is used.

#### Value

xyplot output; use print in Sweave.

## Author(s)

VJ Carey <stvjc@channing.harvard.edu>

### Examples

```
## Not run:
# see vignette
data(sOSR2)
data(c19g) # track excerpt
juxtaPlot(c19g, sOSR2)
```

## End(Not run)

rawCd4DnaseI

#### Description

A data frame with information on the UCSC browser track related to DNaseI hypersensitivity; the rawCD4 object is an eSet extension representing the same information; rawHelaDnaseI is like rawCD4 but results on Hela cells.

#### Usage

```
data(rawCd4DnaseI)
data(rawHelaDnaseI)
data(rawCD4)
```

#### Details

Obtained from a MySQL representation of the data distributed at the Genome Browser FTP site

#### Value

a data.frame

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

#### References

hgdownload.cse.ucsc.edu ... it appears that they do not offer the MYD/MYI representations, just the txt.gz and sql files now. So if you obtain the encodeNhgriDnaseHsChipRawCd4.txt and .sql files at goldenPath/currentGenomes/Homo\\_Sapiens/encode/database, you can reconstruct the underlying data for this data.frame (hg18, Nov 2007).

#### Examples

```
data(rawCd4DnaseI)
dim(rawCd4DnaseI)
rawCd4DnaseI[1:5,]
library(lattice)
xyplot(dataValue~chromStart|chrom, data=rawCd4DnaseI, subset=chrom %in%
    c("chr1", "chr10", "chr19", "chr20"), scales=list(x=list(relation="free")))
```

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