**FDb.FANTOM4.promoters.hg19**

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**hg18ToHg19**

*UCSC liftOver chain for hg18 to hg19, used in build scripts*

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**Description**

In the subdirectory inst/build/, there are several scripts that rebuild this FeatureDb from scratch. Since this is lifted from hg18, a chain is included.

**Author(s)**

Tim Triche, Jr.

**Examples**

```r
data(hg18ToHg19)
hg18ToHg19
```

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**FDb.FANTOM4.promoters.hg19**

*Annotation package for FANTOM4 CAGE promoters from THP-1 cells*

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**Description**

This package loads one or more FeatureDb objects. Such FeatureDb objects are an R interface to prefabricated databases contained by this package. In the case of the FANTOM4 promoter database, it is FDb.FANTOM4.promoters.hg19 (hg18 is available directly from unibas.ch).

**Author(s)**

Tim Triche, Jr.
See Also

features makeFeatureDbFromUCSC import.bed

Examples

```r
## load the library
library(FDb.FANTOM4.promoters.hg19)

## list the contents that are loaded into memory
ls('package:FDb.FANTOM4.promoters.hg19')

## show the db object that is loaded by calling it's name
FDb.FANTOM4.promoters.hg19

## extract features for use in annotating data
FANTOM4.hg19 <- features(FDb.FANTOM4.promoters.hg19)

## we'd prefer if R would stop us from comparing across assemblies:
met <- metadata(FDb.FANTOM4.promoters.hg19) ## need to fetch genome
genome(FANTOM4.hg19) <- met[which(met[, 'name'] == 'Genome'), 'value']

## Plot the observed/expected CpG ratio look like across promoters
## (computed as Pr(CG) / Pr(C)Pr(G)Pr(CG|G,C) within a 3kb window)
## Conversion back to numeric is due to an artifact of features()
values(FANTOM4.hg19)$oecg <- as.numeric(values(FANTOM4.hg19)$oecg)
hist(values(FANTOM4.hg19)$oecg, breaks=200,
    xlab='Observed/expected CpG content from hg19',
    main='FANTOM4 promoter CpG content, 3kb windows')

## The function used for this is FDb.FANTOM4.promoters.hg19:::oecg()```
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