# Package 'PeakError'

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Title Compute the Label Error of Peak Calls			
Description Chromatin immunoprecipitation DNA sequencing results in genomic tracks that show enriched regions or peaks where proteins are bound.  This package implements fast C code that computes the true and false positives with respect to a database of annotated region labels.			
Suggests testthat, ggplot2			
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## Description

Check for a valid data.frame with chrom names.

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#### Usage

checkChrom(df)

## **Arguments**

df df

#### Author(s)

Toby Dylan Hocking

checkPositions

checkPositions

## **Description**

Check for a valid data.frame with chromStart, chromEnd.

## Usage

checkPositions(df)

## **Arguments**

df df

## Author(s)

Toby Dylan Hocking

PeakError

PeakError

## **Description**

Compute true and false positive peak calls, with respect to a database of annotated regions.

## Usage

PeakError(peaks, regions)

## Arguments

peaks data.frame with columns chrom, chromStart, chromEnd. NOTE: chromStart

should be 0-based and chromEnd should be 1-based. EXAMPLE: the first 100 base of of a chromosome are chromStart=0, chromEnd=100. The second 100

bases are chromStart=100, chromEnd=200.

regions data.frame with columns chrom, chromStart, chromEnd, annotation.

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#### Value

data.frame for each region with additional counts of true positives (tp, possible.tp), false positives (fp, possible.fp, fp.status), and false negatives (fn, fn.status).

#### Author(s)

Toby Dylan Hocking

## **Examples**

```
x < - seq(5, 85, by=5)
peaks <- rbind(</pre>
  Peaks("chr2", x, x+3),
  Peaks("chr3", c(25, 38, 57), c(33, 54, 75)),
  Peaks("chr4", c(5, 32, 38, 65), c(15, 35, 55, 85)),
  Peaks("chr5", c(12, 26, 56, 75), c(16, 54, 59, 85)))
regions.list <- list()
for(chr in 1:5){
  regions.list[[chr]] <- data.frame(</pre>
    chrom=paste0("chr", chr),
    chromStart=c(10, 30, 50, 70),
    chromEnd=c(20, 40, 60, 80),
    annotation=c("noPeaks", "peakStart", "peakEnd", "peaks"))
}
regions <- do.call(rbind, regions.list)</pre>
err <- PeakError(peaks, regions)</pre>
ann.colors <- c(</pre>
  noPeaks="#f6f4bf",
  peakStart="#ffafaf",
  peakEnd="#ff4c4c",
  peaks="#a445ee")
if(require(ggplot2)){
  ggplot()+
    geom_rect(aes(
      xmin=chromStart+1/2, xmax=chromEnd+1/2,
      ymin=-1, ymax=1,
      fill=annotation,
      linetype=fn.status,
      size=fp.status),
      data=err, color="black")+
    scale_y_continuous("", breaks=NULL)+
    scale_linetype_manual(
      values=c("false negative"="dotted", correct="solid"))+
    scale_size_manual(
      values=c("false positive"=3, correct=1))+
    scale_fill_manual(
      values=ann.colors,
      breaks=names(ann.colors))+
    facet_grid(chrom ~ .)+
    theme_bw()+
    guides(
      fill=guide_legend(order=1),
```

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```
linetype=guide_legend(order=2, override.aes=list(fill="white")),
    size=guide_legend(order=3, override.aes=list(fill="white")))+
    theme(panel.margin=grid::unit(0, "cm"))+
    geom_segment(aes(
        chromStart+1/2, 1/2, xend=chromEnd+1/2, yend=1/2),
        data=peaks, color="deepskyblue", size=2)+
    scale_x_continuous(
        "position on chromosome",
        breaks=seq(10, 90, by=10))+
    geom_text(aes(
        base, -1/2, label="N"),
        data.frame(base=10:90),
        color="deepskyblue")
}
```

PeakErrorChrom

PeakErrorChrom

## **Description**

Compute the PeakError assuming that peaks and regions are on the same chrom.

#### Usage

```
PeakErrorChrom(peaks,
    regions)
```

## **Arguments**

peaks

data.frame with columns chromStart, chromEnd. NOTE: chromStart should be 0-based and chromEnd should be 1-based. EXAMPLE: the first 100 base of of a chromosome are chromStart=0, chromEnd=100. The second 100 bases are

chromStart=100, chromEnd=200.

regions

data.frame with columns chromStart, chromEnd.

## Value

data.frame with 1 row for each region and error columns.

#### Author(s)

Toby Dylan Hocking

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## Description

Make a data.frame that represents a list of peaks.

## Usage

```
Peaks(chrom = factor(),
   base.before = integer(),
   last.base = integer())
```

## Arguments

chrom character or factor with chrom name for example "chr22"

base.before integer, base before peak.

last.base integer, last base of peak.

#### Value

data.frame with columns chrom, chromStart, chromEnd.

## Author(s)

Toby Dylan Hocking

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