

Package ‘PostChicago’

June 22, 2026

Type Package

Title PostChicago - visualization and integration of Capture-(Hi)C data

Version 0.99.2

Description PostCHiCAGO (stylized from here as PostChicago) is a toolbox for visualising and assessing the output from the CHiCAGO pipeline (SOURCE).
The plots created by PostChicago show reads or CHiCAGO scores over different regions.
PostChicago can integrate different experiments with other types of datasets and compare separate conditions.

biocViews Software, GeneRegulation, Epigenetics, Visualization

License Artistic-2.0

URL <https://github.com/FeldmannLabDKFZ/PostChicago>

Encoding UTF-8

RoxygenNote 7.3.3

Imports GenomicRanges, pheatmap, S4Vectors, IRanges, ggplot2, RColorBrewer, Chicago, utils, graphics, grDevices, stringr, gridExtra, BiocFileCache, matrixStats, rtracklayer, patchwork, dplyr, tidyr, rlang

Suggests knitr, rmarkdown, testthat (>= 3.0.0), data.table, BiocGenerics, codetools, BiocStyle, ragg

Config/testthat/edition 3

VignetteBuilder knitr

NeedsCompilation no

Depends R (>= 4.4.0)

git_url <https://git.bioconductor.org/packages/PostChicago>

git_branch devel

git_last_commit 9927510

git_last_commit_date 2026-06-05

Repository Bioconductor 3.24

Date/Publication 2026-06-21

Author Angelika Feldmann [aut, cre] (ORCID:
<<https://orcid.org/0000-0001-7094-8081>>),
Samuel Krall [aut],
Belinda Blum [aut]

Maintainer Angelika Feldmann <angelika.feldmann@dkfz-heidelberg.de>

Contents

| | |
|------------------------------------|-----------|
| PostChicago-package | 2 |
| aggregatePeaks | 3 |
| aggregatePeaks_regions | 4 |
| annotateInts | 5 |
| baitmap2baited_genes | 6 |
| boxplotsCapC | 7 |
| getMatrix | 8 |
| loadCdList | 10 |
| loadGrl | 11 |
| makeBedGraphs | 12 |
| makeIntsTable | 13 |
| makeOneGeneOnePeak | 15 |
| makeQCplots | 17 |
| plotAggregatePeaks | 18 |
| plotInteractions | 19 |
| plotSigIntsStats | 22 |
| quantifyWithinPeaks | 23 |
| rmap2rmapgr | 24 |
| runChicagoForPostChicago | 25 |
| Index | 27 |

| | |
|---------------------|---|
| PostChicago-package | <i>PostChicago: PostChicago - visualization and integration of Capture-(Hi)C data</i> |
|---------------------|---|

Description

PostCHiCAGO (stylized from here as PostChicago) is a toolbox for visualising and assessing the output from the CHiCAGO pipeline (SOURCE). The plots created by PostChicago show reads or CHiCAGO scores over different regions. PostChicago can integrate different experiments with other types of datasets and compare separate conditions.

PostCHiCAGO (stylized from here as PostChicago) is a toolbox for visualising and assessing the output from the CHiCAGO pipeline (SOURCE). The plots created by PostChicago show reads or CHiCAGO scores over different regions. PostChicago can integrate different experiments with other types of datasets and compare separate conditions.

PostCHiCAGO (stylized from here as PostChicago) is a toolbox for visualising and assessing the output from the CHiCAGO pipeline (SOURCE). The plots created by PostChicago show reads or CHiCAGO scores over different regions. PostChicago can integrate different experiments with other types of datasets and compare separate conditions.

Author(s)

Maintainer: Angelika Feldmann <angelika.feldmann@dkfz-heidelberg.de> ([ORCID](#))

Authors:

- Samuel Krall <sammikrall@gmail.com>
- Belinda Blum <belinda.blum@dkfz.de>

See Also

Useful links:

- <https://github.com/FeldmannLabDKFZ/PostChicago>

Useful links:

- <https://github.com/FeldmannLabDKFZ/PostChicago>

Useful links:

- <https://github.com/FeldmannLabDKFZ/PostChicago>

aggregatePeaks

*Aggregate Interactions to Interaction Peaks***Description**

Takes a table of interactions (ints) and aggregates significant interactions within the distance `dis` in restriction fragments for each bait. Works similarly to the `reduce()` function from `GenomicRanges`. Caution: All interactions, regardless the sample from which they originated, are aggregated, creating a 'consensus' interaction peak set. If interaction peaks should be aggregated for individual samples, we recommend a prior sample-specific filtering of ints. Only creates the table if it is not yet saved, otherwise loads the existing table.

Usage

```
aggregatePeaks(ints, dis, samples, fileprefix = "ints", outdir = NULL)
```

Arguments

| | |
|-------------------------|---|
| <code>ints</code> | Table containing all interactions that should be aggregated. |
| <code>dis</code> | Numeric, distance in restriction fragments over which interactions should be aggregated. <code>dis=1</code> will merge two adjacent fragments with significant interactions. |
| <code>samples</code> | Character vector of sample names, should correspond to the sample names used in ints. Typically corresponding to the names provided in the list of <code>ChicagoData</code> objects <code>cd</code> from which ints has been generated. |
| <code>fileprefix</code> | Default: <code>ints</code> ; Prefix of the filename for the <code>aggregatePeaks</code> table file, will be added the following extension: <code>paste0('_aggregatePeaks_in_regions_',dis,'bp.txt')</code> . |
| <code>outdir</code> | Default: current directory; Directory where the final table is saved. Will not create a copy if already present. |

Value

A table containing the positions of aggregated interactions. Redefines `intIDs` so that the ID contains all significant interactions that were aggregated. Data is automatically saved in a 'aggregate-Peaks_dis' text file whose name is defined by `fileprefix` and `dis`. Also saved as a bedfile.

Examples

```
# example code:
extdata <- system.file("extdata", package="PostChicago")
outputDir <- file.path(extdata,'postchicago')

#read in the interactions table:
ints=read.delim(paste0(outputDir,'/ints_all.txt'),stringsAsFactors=FALSE)

#aggregate all peaks within a distance of 5 restriction fragments
dis <- 5
samples <- names(ints)[grep('_N$',names(ints))]
samples <- unlist(strsplit(samples,split='_N'))
aggr <- aggregatePeaks(ints,dis,samples)
head(aggr)

dim(ints)
dim(aggr)
```

aggregatePeaks_regions

Aggregate Interactions to Interaction Peaks by Distance in bp

Description

Takes a table of interactions (ints) and aggregates significant interactions within the distance dis in base pairs (bp) for each bait. Works similarly to the reduce() function from GenomicRanges. Caution: All interactions, regardless the sample from which they originated, are aggregated, creating a 'consensus' interaction peak set. If interaction peaks should be aggregated for individual samples, we recommend a prior sample-specific filtering of ints. Only creates the table if it is not yet saved, otherwise loads the existing table.

Usage

```
aggregatePeaks_regions(ints, dis, samples, fileprefix = "ints", outdir = NULL)
```

Arguments

| | |
|------------|---|
| ints | Table containing all interactions that should be aggregated. |
| dis | Numeric, distance in bp over which interactions should be aggregated. dis=1 will merge two adjacent fragments with significant interactions. |
| samples | Character vector of sample names, should correspond to the sample names used in ints. Typically corresponding to the names provided in the list of ChicagoData objects cd from which ints has been generated. |
| fileprefix | Default: ints; Prefix of the filename for the aggregatePeaks table file, will be added the following extension: paste0('_aggregatePeaks_in_regions_',dis,'bp.txt'). |
| outdir | Default: current directory; Directory where the final table is saved. Will not create a copy if already present. |

Value

A table containing the position of aggregated interactions. Redefines intIDs so that the ID of the first interacting fragment in a peak stands in as otherEnd. Data is automatically saved in a 'aggregatePeaks_dis' text file whose name is defined by fileprefix and dis. Also saved as a bedfile.

Examples

```
# example code:
extdata <- system.file("extdata", package="PostChicago")
outputDir <- file.path(extdata,'postchicago')

#read in the interactions table:
ints=read.delim(paste0(outputDir,'/ints_all.txt'),stringsAsFactors=FALSE)

#aggregate all peaks within a distance of 5 restriction fragments
dis <- 5000
samples <- names(ints)[grep('_N$',names(ints))]
samples <- unlist(strsplit(samples,split='_N'))
aggr <- aggregatePeaks_regions(ints,dis,samples)
head(aggr)

dim(ints)
dim(aggr)
```

| | |
|--------------|------------------------------|
| annotateInts | <i>Annotate Interactions</i> |
|--------------|------------------------------|

Description

Annotates interaction tables containing CaptureC data with features (typically ChIP-Seq peaks) from a GRangesList object (grl). Annotation is based on feature overlap.

Usage

```
annotateInts(ints, grl, minov = NULL, maxgp = NULL)
```

Arguments

- ints a table with interactions derived from ChicagoData objects by makeIntsTable(). Assumes six named columns: 'seqnames_bait','start_bait','end_bait' (coordinates of the baited restriction fragment) and 'seqnames_otherEnd','start_otherEnd','end_otherEnd' (coordinates of the interacting restriction fragment).
- grl GRangesList containing features with which ints should be annotated. Names of the grl correspond to the names that will be given to table columns.
- minov Default:NULL, minimum overlap (minoverlap in findOverlaps) between otherEnd or bait fragment and the feature.
- maxgp Default:NULL, maximum gap (maxgap in findOverlaps) between otherEnd or bait fragment and the feature.

Value

interactions file with annotations from GRanges file

Examples

```
# basic usage of annotateInts

##Create an interactions table
ints <- data.frame(seqnames_bait='chr1', start_bait=1, end_bait=1000,
                  seqnames_otherEnd='chr1', start_otherEnd=10000, end_otherEnd=20000)
ints

##create a GRangesList with annotations
gr1 <- GenomicRanges::GRanges('chr1',IRanges::IRanges(10500,11000))
gr2 <- GenomicRanges::GRanges('chr1',IRanges::IRanges(900,1100))
grl <- GenomicRanges::GRangesList(gr1,gr2)
names(grl)=c('mytestpeaks1','mytestpeaks2')
grl

##Annotate interactions with intervals from grl
annotateInts(ints,grl)
```

baitmap2baited_genes *Convert baitmap to baited_genes*

Description

Searches for the Chicago baitmap file in ‘designDir’ and converts it to a GRanges object. Only executed if the output file ‘baited_genes.rds’ doesn’t exist yet, otherwise loads the existing file.

Usage

```
baitmap2baited_genes(designDir = "designDir", save = FALSE)
```

Arguments

| | |
|-----------|--|
| designDir | Path to the directory with Chicago design files. Default: ‘designDir’ |
| save | If TRUE, saves baited_genes as baited_genes.rds in designDir. Default: FALSE |

Value

Returns a GenomicRanges object with the positions of the baits, annotated with fragment IDs (re_id) and gene names

Examples

```
# Example running baitmap2baited_genes
extdata <- system.file("extdata", package="PostChicago")
designDir <- file.path(extdata,'designDir')
baitmap2baited_genes(designDir,save=FALSE)
```

 boxplotsCapC

Boxplots of summarized interactions data.

Description

Boxplots of normalized interaction data quantified from either interaction peaks or oneGeneOne-Peak tables. Returns 4 boxplots, containing downsampled read and Chicago score counts, total and size-normalized values. Function assumes that replicate data are saved in a column with the name structure '_rep1'.

Usage

```
boxplotsCapC(
  annotatedCapCTable,
  reps = FALSE,
  col = NULL,
  show_outliers = TRUE,
  show_notch = TRUE,
  name = NULL
)
```

Arguments

| | |
|--------------------|--|
| annotatedCapCTable | Table containing Capture-C quantification either over intervals or interaction peaks. Assumes the output from <code>quantifyWithinPeaks()</code> . |
| reps | TRUE/FALSE(default); should replicate or summary data be plotted? Assumes replicate data are stored in column names '_rep1', '_rep2' etc. |
| col | Custom plot color. |
| show_outliers | TRUE(default)/FALSE; Should outliers be shown? |
| show_notch | TRUE(default)/FALSE |
| name | optional, plot names |

Value

Boxplots of scores and reads of each experiment of a specific feature

Examples

```
# example code

##Create an rmapgr object
rmapgr <- GenomicRanges::GRanges(rep(1,100), IRanges::IRanges( 1:100,2:101 ))
rmapgr$id=1:length(rmapgr)

##create interaction data table with two interactions
data <- data.frame(intID=c('1;2', '1;3,1;5'),
                  baitID = as.character(c(1,1)),
                  otherEndID=as.character(c(2, '3,5')))

##create a ChicagoData object with two samples
```

```

cdlist <- list(
  sample1 = data.frame(baitID=rep(1,11),
                      otherEndID=c(2:12),
                      N=c(sample(1:100,10,replace=TRUE),1000),
                      score=c(sample(0:10,10,replace=TRUE),100)),

  sample2 = data.frame(baitID=rep(1,11),
                      otherEndID=c(2:12),
                      N=sample(90:110,11,replace=TRUE),
                      score=sample(0:10,11,replace=TRUE))
)

##create replicate ChicagoData object from cdlist:
cdlist.reps <- c(cdlist,cdlist)
f<-function(x){
  x$N=x$N*sample((7:13)/10,11,replace=TRUE)
  x$score=x$score*sample((7:13)/10,11,replace=TRUE)
  return(x)
}
cdlist.reps=lapply(cdlist.reps,FUN=f)
names(cdlist.reps)=paste0(rep(names(cdlist),2),'_rep',rep(1:2,each=2))
cdlist.reps=cdlist.reps[order(names(cdlist.reps))]

##run quantifyWithinPeaks
annotatedCapCTable <- quantifyWithinPeaks(data, cdlist, cdlist.reps, rmapgr=rmapgr)

##plots
col=c('grey','red')
boxplotsCapC(annotatedCapCTable, col=col)
boxplotsCapC(annotatedCapCTable, col=col,show_notch=FALSE)
boxplotsCapC(annotatedCapCTable, col=rep(col,each=2),reps=TRUE)

```

getMatrix

Create Interaction Matrices

Description

Function that creates and normalizes matrices containing reads or scores of fragment-to-fragment interactions surrounding the interactions that are provided in the interactions table 'd'.

Output is a list of matrices for the samples supplied in 'L', including distance-matched control matrices. The matrix list is automatically saved in the 'resfolder'.

Matrices are stranded, such that the promoter is to the right (+ direction) of the interaction, which is important because the read numbers are increasing with increased promoter proximity. A new list is only created and saved as '.rds' file if the matrix cannot be found in the specified 'resfolder', otherwise the preexisting matrix list is loaded. If the summit of an interaction peak or oneGeneOnePeak file is provided, getMatrix() extracts the summit info from the data, otherwise the function simply determines the central fragment of an interaction peak.

If norm = TRUE, normalized matrices are calculated from pre-existing unnormalized matrices. All values are normalized to the interaction in the reference sample defined by 'normsam', which is recommended to be a positive control sample.

Usage

```
getMatrix(
  L,
  zoom = 100,
  d,
  resfolder = "matrices/",
  type = "reads",
  readnorm = "downsampling",
  name = "",
  norm = FALSE,
  normsam = 1,
  pseudo = 1,
  rmapgr = rmapgr
)
```

Arguments

| | |
|-----------|---|
| L | list of ChicagoData tables. |
| zoom | Distance around interaction in restriction fragments for which matrices should be generated. |
| d | A table with cis-chromosomal interactions. Should contain the columns 'baitID', 'otherEndID', defined as in Chicago and can have a column 'intID', which is expected to be in the format 'baitID;otherEndID'. 'otherEndID' should be the center of interaction, from which the matrices will be calculated. |
| resfolder | Default: 'matrices/'. Path to the directory, where matrices should be stored. Will not create a copy if one is already saved in the given directory. Creates the directory if nonexistent. |
| type | 'reads' (default)/'scores'. If reads, the function performs downsampling relative to the sample with the smallest amount of reads that are mapped to baits. |
| readnorm | Default: 'downsampling'. Read normalization, either 'downsampling' or 'scaling'. Scaling means the same type of normaliyation as for the lineplots in plot-Interactions(). |
| name | Name of the matrix. |
| norm | TRUE/FALSE (default). Should the data be normalized? |
| normsam | numeric; defines to which sample in the list L should the data be normalized. Default: 1 (the first sample!). Only used if norm=TRUE. |
| pseudo | Default: 1; pseudocount to be added during the normalization between matrices. |
| rmapgr | GenomicRanges object containing restriction fragments within the reference genome. IDs must be saved in a column called 'id'. |

Value

List of interaction matrices.

Examples

```
# example code

##create ints table with 4 interactions
ints <- data.frame(intID=c('11;4', '11;7', '11;8', '12;14'),
```

```

        baitID = as.character(c(11,11,11,12)),
        otherEndID=as.character(c(4,7,8,14)),
        seqnames_bait=as.character((paste0('chr',rep(1,4)))),
        start_bait=as.character(c(21,21,21,23)),
        end_bait=as.character(c(22,22,22,23)),
        seqnames_otherEnd=as.character((paste0('chr',rep(1,4)))),
        start_otherEnd=as.character(c(7,13,15,27)),
        end_otherEnd=as.character(c(8,14,16,28))
    )

##make rmapgr
rmapgr <- GenomicRanges::GRanges(rep('chr1',20),
IRanges::IRanges( c(1,3,5,7,9,11,13,15,17,19,21,23,25,27,29,31,33,35,37,39),
c(2,4,6,8,10,12,14,16,18,20,22,24,26,28,30,32,34,36,38,40) ) )
rmapgr$id=1:length(rmapgr)

cdlist <- list(
sample1 = data.frame(baitID=c(rep(11,20),rep(12,20)),
                    otherEndID=rep(c(1:20),2),
                    N=sample(1:20,40,replace=TRUE),
                    N.1=sample(1:20,40,replace=TRUE),
                    N.2=sample(1:20,40,replace=TRUE),
                    score=sample(1:5,40,replace=TRUE)),

sample2 = data.frame(baitID=c(rep(11,20),rep(12,20)),
                    otherEndID=rep(c(1:20),2),
                    N=sample(15:45,40,replace=TRUE),
                    N.1=sample(15:45,40,replace=TRUE),
                    N.2=sample(15:45,40,replace=TRUE),
                    score=sample(1:20,40,replace=TRUE))
)

##getMatrix
getMatrix(L = cdlist, d = ints, zoom = 1, rmapgr=rmapgr)

##normalized matrices
getMatrix(L = cdlist, d = ints, zoom = 1, rmapgr=rmapgr,norm=TRUE,normsam=2)

##Add summit info to interactions:
ints$summit_re_id=c('4','7','8','13,14')
getMatrix(L = cdlist, d = ints, zoom = 1, rmapgr=rmapgr, resfolder='matrices_summits/')

```

loadCdList

Loads List with Interaction Tables

Description

Reads in all ChicagoData objects in resultsDir and converts them to a list of ChicagoData tables. ChicagoData objects must be saved during the Chicago pipeline, which can be done by running Chicago via runChicagoForPostChicago()

Usage

```
loadCdList(resultsDir = "results", baited_genes = NULL)
```

Arguments

| | |
|--------------|---|
| resultsDir | Default: 'results'; Directory containing ChicagoData output. |
| baited_genes | Optional; Used to control for the compatibility between loaded ChicagoData and current experiment; GenomicRanges object with at least two metadata columns: 1) 're_id' (captured fragment ID (bait), corresponding to fragment numbers indicated in rmapgr), 2) 'genenames' (names of the corresponding baits, typically gene names). Can be created automatically from the Chicago baitmap table (see R Bioconductor package 'Chicago') using the function baitmap2baited_genes(). |

Value

List containing ChicagoData tables.

Examples

```
# example code:
extdata <- system.file("extdata", package="PostChicago")
chicagoOutputDir <- file.path(extdata, 'results')
designDir <- file.path(extdata, 'designDir')

#create baited_genes:
baited_genes=baitmap2baited_genes(designDir, save=FALSE)

#load ChicagoData object list:
cdlist <- loadCdList(resultsDir = chicagoOutputDir, baited_genes = baited_genes)
```

loadGrl

Load Intervals into a GenomicRangesList

Description

Reads in .rds and .bed files in intDir into a GRangesList object.

Usage

```
loadGrl(intDir = "intervals")
```

Arguments

| | |
|--------|---|
| intDir | Directory containing .rds or .bed files with intervals. |
|--------|---|

Value

GenomicRanges object grl containing all intervals

Examples

```
# example code

#requires precomputed matrices in the resfolder (see getMatrix())
extdata <- system.file("extdata", package="PostChicago")
intDir <- file.path(extdata,'intervals')
dir(intDir)

#Load intervals for which the boxplots should be quantified into a GRangesList
grl <- loadGrl(intDir = intDir)
```

makeBedGraphs

Create Bedgraphs From ChicagoData

Description

Takes ChicagoData objects supplied in the list L, normalizes them to read coverage mapped to baits. Creates two file types: 1) a bedgraph file containing normalized reads for each sample mapping to the bait defined by id, 2) a bed file of the view point (bait).

Usage

```
makeBedGraphs(
  id,
  L_data,
  rmapgr,
  baited_genes,
  ints = NULL,
  L_norm = NULL,
  outputDir = "bedgraph",
  smoothingWindow = "off"
)
```

Arguments

| | |
|--------------|--|
| id | baitID, must correspond to the baitIDs defined in baited_genes and the supplied rmapgr object |
| L_data | List of ChicagoData (cd) objects. |
| rmapgr | GenomicRanges object containing restriction fragments. IDs must be saved in a column called 'id'. |
| baited_genes | GenomicRanges object with at least two metadata columns: 1) 're_id' (captured fragment ID (bait), corresponding to fragment numbers indicated in rmapgr), 2) 'genenames' (names of the corresponding baits, typically gene names). Can be created automatically from the Chicago baitmap table (see R Bioconductor package 'Chicago') using the function baitmap2baited_genes(). |
| ints | optional; table containing all significant interactions across L_data, usually created using makeIntsTable(). If supplied, a separate bed file containing all significant interactions mapping to the id is created |

| | |
|-----------------|--|
| L_norm | Separate list of ChicagoData tables which should be used to normalize samples. Useful if bedgraphs should be created only for a subset of promoters or genomic locations, but the normalization performed based on the whole dataset. If not provided, L_data is used for normalization. |
| outputDir | Default: 'bedgraph'. Output directory for the .bedgraphs, automatically created if not currently present. |
| smoothingWindow | Default: 'off'. Over how many restriction fragments should the data be smoothed? Not recommended for UCSC visualization, but can improve visualization on IGV. |

Value

Saves bedgraphs and annotation bedfiles of interactions with id as view point for each sample in L_data.

Examples

```
# example code:

extdata <- system.file("extdata", package="PostChicago")
chicagoOutputDir <- file.path(extdata,'results')
designDir <- file.path(extdata,'designDir')

#create baited_genes and rmapgr files:
baited_genes <- baitmap2baited_genes(designDir,save=FALSE)
rmapgr <- rmap2rmapgr(designDir, save = FALSE)

#load ChicagoData object list:
cdlist <- loadCdList(resultsDir = chicagoOutputDir, baited_genes = baited_genes)

## extract all view points (baits):
baits <- unique(unlist(lapply(cdlist,
                             FUN = function(x) unique(x$baitID))))

## create bedgraphs for selected view points and store them in outdir:
outdir <- "bedgraphs"
dir.create(outdir)
makeBedGraphs(baits[1], cdlist, rmapgr,
               baited_genes = baited_genes,
               outputDir = outdir
)

list.files(outdir)
```

makeIntsTable

Create Interactions Table

Description

Integrates all significant interactions from multiple datasets, as supplied by the list of Chicago-Data tables in L. The table is populated with scores, raw and downsampled reads from pooled and (optionally) replicate data.

Usage

```

makeIntsTable(
  L,
  baited_genes,
  repscores = FALSE,
  LL = NULL,
  ngroups = 1,
  scorecut = 5,
  readcut = 1,
  mode = "score",
  intervals = NULL,
  outfolder = NULL,
  overwrite = FALSE,
  saveFile = TRUE
)

```

Arguments

| | |
|--------------|--|
| L | list of ChicagoData objects containing summarized interactions. |
| baited_genes | GenomicRanges object with at least two metadata columns: 1) 're_id' (captured fragment ID (bait), corresponding to fragment numbers indicated in rmapgr), 2) 'genenames' (names of the corresponding baits, typically gene names). Can be created automatically from the Chicago baitmap table (see R Bioconductor package 'Chicago') using the function baitmap2baited_genes(). |
| repscores | TRUE/FALSE (default): Should replicate scores be added? if TRUE, LL must be supplied, Default: FALSE |
| LL | optional: List of ChicagoData tables containing individual replicates of summarized interactions. Names are expected to follow the naming convention '_rep1', '_rep2', ..., '_repn' |
| ngroups | Default: 1; number of groups in which unique view points (baits) should be processed. Efficient computing is achieved for a maximum of ca. 100-200 entries per group. |
| scorecut | Default: 5 (as recommended by Chicago); Chicago score cutoff for significant interactions. Sometimes useful to lower to 3. |
| readcut | Default: 1; Reads cutoff for significant interactions. |
| mode | Options: 'score', 'read' or 'both'; Default: 'score'; Which parameter should be used for the extraction of significant interactions? |
| intervals | optional: A GenomicRanges object with intervals with which the otherEnds should overlap. Useful if mode='reads', to reduce the number of interactions in ints. |
| outfolder | Default: current directory; Path to the directory in which the output is saved. |
| overwrite | Default: FALSE. If TRUE runs the function even if an ints_all.txt is present in the outputdir, otherwise loads ints_all.txt. |
| saveFile | Default: TRUE; saves ints_all.txt in outfolder. |

Value

Table of all significant interactions.

Examples

```
# example code

##Create an rmapgr object
rmapgr <- GenomicRanges::GRanges(rep(1,5), IRanges::IRanges( 1:5,2:6))
rmapgr$id=1:5

##create baited_genes with one gene
baited_genes <- GenomicRanges::GRanges(1,IRanges::IRanges(1,2))
baited_genes$re_id=1
baited_genes$genename='testgene'

##create a list of ChicagoData objects with two samples
cdlist <- list(
  sample1 = data.frame(baitID=rep(1,4),
                      otherEndID=c(2,3:5),
                      N=c(2,5,10,8),
                      N.1=c(2,5,10,8),
                      N.2=c(3,4,9,7),
                      score=c(5, 1, 2, 4)),

  sample2 = data.frame(baitID=rep(1,4),
                      otherEndID=c(2,3:5),
                      N=c(20,3,1,6),
                      N.1=c(20,3,1,6),
                      N.2=c(17,4,2,3),
                      score=c(3, 5, 3, 5))

)

##run makeIntsTable
ints = makeIntsTable(cdlist,baited_genes,saveFile=FALSE)
head(ints)
```

makeOneGeneOnePeak *Make one-Gene-one-Peak Interaction Tables.*

Description

This function takes the interaction table `ints` (typically containing all significant interactions) and intersects them with intervals provided in the `GRangesList` `grl`.

Each entry in `grl` receives a unique ID and is assigned only one interaction with each bait with which at least one significant interaction could be detected.

Within the resulting `OneGeneOnePeak` table, `start_otherEnd` and `end_otherEnd` correspond to the start of the first and the end of the last restriction fragment that overlap a particular interval. `otherEndID` corresponds to all overlapped restriction fragments from `rmapgr` separated by commas ('1,2,3'). `summit_re_id` corresponds to the restriction ID of the central restriction fragment within `rmapgr` that overlaps an interval. If two intervals overlap exactly the same set of restriction fragments from `rmapgr`, this will be visible in the output table column `'interval_id_<my_interval>'` as two comma-separated ids.

`OneGeneOnePeak` tables for each interval within `grl` are saved in the output directory.

Usage

```
makeOneGeneOnePeak(gr1, rmapgr, ints, maxgap = -1, folder = "oneGeneOnePeak")
```

Arguments

| | |
|---------------------|--|
| <code>gr1</code> | GRangesList containing features for which oneGeneOnePeak tables should be made. Names of the <code>gr1</code> correspond to the names that will be given to table columns. |
| <code>rmapgr</code> | GenomicRanges object containing restriction fragments from the reference genome. IDs must be saved in a column called 'id'. |
| <code>ints</code> | Interactions table containing all significant interactions and typically derived by <code>makeIntsTable()</code> . |
| <code>maxgap</code> | Default: -1, full overlap; Maximum gap between the otherEnd and an interval to count as overlap (see <code>findOverlaps()</code>). |
| <code>folder</code> | Output folder. Default: 'oneGeneOnePeak' |

Value

A list with oneGeneOnePeak interactions. Saves a total of four files, for each of the overlapping intervals in `gr1`: 1) An annotated OneGeneOnePeak interaction table as `.txt`. 2) An interaction table containing all information about interacting intervals, added to each row of the original interaction table `ints`, (not oneGeneOnePeak yet!): `'ints_<myintervaltype>_interacting.txt'` 3) A reference table containing all interacting intervals, their ids (`interval_id`) and covered `re_ids`. In this table, intervals are summarized to one interval if they overlap exactly the same restriction fragments: `'<myintervaltype>_interacting_rmapFrag_annotated.txt'` 4) A reference table containing all intervals with `interval_id` and `summit_re_id`: `'<myintervaltype>_withID.txt'` oneGeneOnePeak tables can next be annotated with quantitative data using `quantifyWithinPeaks()`.

Examples

```
# example code

##create ints table with 4 interactions
ints <- data.frame(intID=c('1;2', '1;3', '1;4','2;8'),
  baitID = as.character(c(1,1,1,2)),
  otherEndID=as.character(c(2,3,4,8)),
  seqnames_bait=as.character((paste0('chr',rep(1,4)))),
  start_bait=as.character(c(1,1,1,2)),
  end_bait=as.character(c(2,2,2,3)),
  seqnames_otherEnd=as.character((paste0('chr',rep(1,4)))),
  start_otherEnd=as.character(c(3,5,7,15)),
  end_otherEnd=as.character(c(4,6,8,16))
)

##make rmapgr
rmapgr <- GenomicRanges::GRanges(rep('chr1',8),
  IRanges::IRanges( c(1,3,5,7,9,11,13,15),
    c(2,4,6,8,10,12,14,16) ) )
rmapgr$id=1:length(rmapgr)

##make gr1 with one interval type and two intervals
gr1 <- GenomicRanges::GRangesList(testintervals=GenomicRanges::GRanges(rep('chr1',3),
  IRanges::IRanges(c(2,5,6), c(4,7,8))),
```

```
testintervals2=GenomicRanges::GRanges(rep('chr1',2),
IRanges::IRanges(c(5,11), c(10,13))),
testintervals3 = GenomicRanges::GRanges('chr1',IRanges::IRanges(1,2)))

##run makeOneGeneOnePeak where intervals have to fully overlap
ogopList <- makeOneGeneOnePeak(grl = grl,rmapgr = rmapgr, ints = ints, maxgap=-1)
```

makeQCplots

Quality Control Plots

Description

Creates and saves correlative heatmaps of reads and scores from interaction tables.

Usage

```
makeQCplots(ints, folder = NULL, fontsize = 10)
```

Arguments

| | |
|----------|--|
| ints | Interactions Table containing all significant interactions. Typically created using makeIntsTable(). |
| folder | Default: Current folder; Path to the directory in which plots are saved. |
| fontsize | Default: 10. |

Value

A pdf file containing QC heatmaps: pheatmaps_QC.pdf.

Examples

```
# example code:
extdata <- system.file("extdata", package="PostChicago")
inDir <- file.path(extdata,'postchicago')

#read in interactions:
ints=read.delim(paste0(inDir,'/ints_all.txt'),stringsAsFactors=FALSE)

#assess interactions:
makeQCplots(ints)
```

plotAggregatePeaks *Plot Aggregate Interaction Profiles*

Description

Takes a list of matrices and plots aggregate interaction signal, similar for metaprofiles in ChIP-Seq.

Usage

```
plotAggregatePeaks(
  Lm,
  mainprefix = NULL,
  col = NULL,
  ylim = NULL,
  lty = NULL,
  xlim = NULL,
  k = 3,
  ylab = "normalised reads",
  plotwhat = "all"
)
```

Arguments

| | |
|------------|---|
| Lm | List of matrices containing interaction reads or scores. |
| mainprefix | Optional; text added to the plot title. |
| col | Custom colors for aggregate plots. Default: 4 colors. |
| ylim | Optional |
| lty | Optional; Line type for the aggregate plots |
| xlim | Optional; should be added in restriction fragments surrounding the interval/peak center. Example: c(-50,50). Default: All restriction fragments represented in the matrix. |
| k | Numeric; over how many fragments should the running mean be calculated? Default: 3 |
| ylab | Optional; Default: 'normalised reads' |
| plotwhat | Defines what types of plots will be plotted. Options are 'all', 'mean', 'median' 'mean' and 'median' plot running averages over k fragments. Default: 'all': Four plots representing mean and median read counts both smoothed and unsmoothed over k fragments. |

Value

Plot containing aggregate profiles of interactions.

Examples

```
# example code

##create list of matrices:
```

```

##matrix1
m1<-matrix(c(c(1:10,11,9:0)*sample((8:1.2)/10,21,replace=TRUE),
            c(1:10,11,9:0)*sample((8:1.2)/10,21,replace=TRUE),
            c(1:10,11,9:0)*sample((8:1.2)/10,21,replace=TRUE)),nrow=3)

##matrix2
m2<-matrix(c(c(21:30,39,29:20)*sample((8:1.2)/10,21,replace=TRUE),
            c(21:30,39,29:20)*sample((8:1.2)/10,21,replace=TRUE),
            c(21:30,39,29:20)*sample((8:1.2)/10,21,replace=TRUE)),nrow=3)
rownames(m1)=c('1;2','2;3','3;4')
rownames(m2)=rownames(m1)

##matrix list:
Lm=list(m1=m1,m2=m2,m1_con=m1*0.2,m2_con=m2*0.1)

##plot smoothed median enrichment profiles:
plotAggregatePeaks(Lm,plotwhat='median', ylab='reads',
                  col=rep(c('red','blue'),2), lty=c(1,1,3,3))

##plot smoothed median profiles, increasing the smoothing window:
plotAggregatePeaks(Lm,plotwhat='median',k=5,ylab='reads',
                  col=rep(c('red','blue'),2), lty=c(1,1,3,3))

##plot smoothed mean profiles:
plotAggregatePeaks(Lm,plotwhat='mean',k=3,ylab='reads',
                  col=rep(c('red','blue'),2), lty=c(1,1,3,3))

##plot median and mean, smoothed and unsmoothed profiles:
par(mfrow=c(2,2))
plotAggregatePeaks(Lm,plotwhat='all',k=3,ylab='reads',
                  col=rep(c('red','blue'),2), lty=c(1,1,3,3))

```

plotInteractions

Plot Lineplots of Interactions

Description

Plots lineplots of normalized Capture-C read counts around one view point (bait) from one or many samples. Optional annotation with genomic intervals of choice and significant interactions.

Usage

```

plotInteractions(
  L,
  id,
  k,
  zoom = 2e+05,
  rmapgr,
  ylim = NULL,
  show.legend = TRUE,
  d = NULL,
  preselected = FALSE,
  name = NULL,

```

```

intervals = NULL,
show.legend.intervals = TRUE,
xlim = NULL,
col = NULL,
colintervals = NULL,
colints = NULL,
troubleshooting = FALSE,
lwd = 2,
lty = NULL,
show.legend.outside = FALSE,
cex.intervals = 0.7,
cex.legend = 1
)

```

Arguments

| | |
|-----------------------|--|
| L | List with the ChicagoData tables |
| id | View point (baitID), corresponds to a fragment id in rmapgr. |
| k | Smoothing factor across k restriction fragments (running average). |
| zoom | Default: 200000; Area around the bait in +/-bp to plot. |
| rmapgr | GenomicRanges object containing restriction fragments from the reference genome. IDs must be saved in a column called 'id'. |
| ylim | optional; Plotting limit of the y axis. |
| show.legend | TRUE(default)/FALSE; should the sample legend be shown inside the plot area? |
| d | Default: FALSE; interactions table to highlight significant interactions as transparent rectangles (shading). By default, if d is given, highlights any interaction. If different categories of interactions should be highlighted, these are specified in a column called 'clusters_refined'. Example usage would be 'lost', 'gained' and 'stable' interactions. Up to 8 categories can be defined which then receive a different color each. |
| preselected | , Default: FALSE; are interactions in d preselected for the correct baitID? Useful in cases when ids in the interactions table do not match the ids in L. |
| name | optional; Name of the view point, Default: name = baitID. |
| intervals | optional; Intervals to be plotted, provided as a named GenomicRangesList, useful for annotation of the plot with ChIP-Seq data. |
| show.legend.intervals | TRUE(default)/FALSE; should the legend for intervals be shown? |
| xlim | optional; Plotting limit of the x axis, as xlim=c(30000,200000), overrides zoom. |
| col | Custom colors of the lines. Default: 'Okabe-Ito' with a max of 8 different samples. |
| colintervals | Custom colors for interval annotation. |
| colints | Custom colors for significant interaction shading. Per default, interactions are plotted using the transparency parameter alpha of 0.3. However, this can be overridden by providing the alpha parameter together with the color: col <- rgb(0,0,0,0.2) will use black with an opacity of 0.2. |
| troubleshooting | if TRUE will return messages showing how far the process is, Default: FALSE |
| lwd | line width, Default 2pt |

lty line type, Default: 1
 show.legend.outside If TRUE, shows the legend outside of the plot. Note that the total plot area then becomes wider. Default: FALSE
 cex.intervals Defines the size of the intervals legend. Default: 0.7
 cex.legend Defines the size of the samples legend. Default: 1

Value

Line plots of Capture-C reads surrounding one view point.

Examples

```
# example code (See package Vignette for more examples!)

#define directories
extdata <- system.file("extdata", package="PostChicago")
chicagoOutputDir <- file.path(extdata,'results')
designDir <- file.path(extdata,'designDir')

#create baited_genes and rmapgr:
baited_genes=baitmap2baited_genes(designDir,save=FALSE)
rmapgr=rmap2rmapgr(designDir,save=FALSE)

#load list with ChicagoData objects:
cdlist <- loadCdList(resultsDir = chicagoOutputDir, baited_genes = baited_genes)
cdlist=cdlist[-grep('rep',names(cdlist))]

#standard plot:
col=c('indianred1','seagreen')
name='Hoxb3'
id=baited_genes[baited_genes$genename==name]$re_id
k=15
ylim=c(0,200)
plotInteractions(cdlist,id,k,ylim=ylim,show.legend = TRUE,name=name, rmapgr=rmapgr,col=col)

#vary line type and width:
plotInteractions(cdlist,id,k,ylim=ylim,show.legend = TRUE,name=name, rmapgr=rmapgr,col=col,
  lwd=1,lty=c(1,2))

#change zoom around the bait:
plotInteractions(cdlist,id,k,ylim=ylim,show.legend = TRUE,name=name, rmapgr=rmapgr,col=col,
  zoom=500000)

#change zoom and the number of restriction fragments over which the running mean is calculated:
plotInteractions(cdlist,id,k=61,ylim=ylim,show.legend = TRUE,name=name, rmapgr=rmapgr,col=col,
  zoom=500000)

#display significant interactions in the plot

#read in interactions:
inDir <- file.path(extdata,'postchicago')
ints <- read.delim(paste0(inDir,'/ints_all.txt'),stringsAsFactors=FALSE)

#choose the color with which interactions should be highlighted
colints <- 'yellow'
```

```
#plot
plotInteractions(cdlist,id,k=31,ylim=ylim,show.legend = TRUE,name=name, rmapgr=rmapgr,col=col,
  zoom=200000, d = ints, colints=colints)
```

plotSigIntsStats *Plot Interaction Statistics*

Description

Takes a list of ChicagoData tables and plots a per sample statistical summary.

The summary includes the total number of reads mapped to baits, total genes with significant interactions, total unique and significant interactions, the corresponding distributions per bait.

Usage

```
plotSigIntsStats(
  L,
  col = NULL,
  lty = NULL,
  filename = NULL,
  plotDistribution = TRUE,
  plotExamples = FALSE
)
```

Arguments

| | |
|------------------|---|
| L | list of ChicagoData tables for which to extract statistics. |
| col | Custom colors for the plots. Default: Okabe-Ito, 8 colors. |
| lty | optional; vector defining line types. |
| filename | optional; if provided, the plots will be saved under filename; must have the extension '.pdf' |
| plotDistribution | if TRUE (default), distributions of reads and interactions per bait are plotted. |
| plotExamples | Default: FALSE; if TRUE, example distributions will be plotted based on the data from Mahara et al. |

Value

Plots with statistics of interactions for each sample and bait.

Examples

```
# example code:
extdata <- system.file("extdata", package="PostChicago")
dataPath <- file.path(extdata, 'dataPath')
chicagoOutputDir <- file.path(extdata, 'results')
designDir <- file.path(extdata, 'designDir')
intDir <- file.path(extdata, 'intervals')
outputDir <- file.path(extdata, 'postchicago')
```

```

baited_genes=baitmap2baited_genes(designDir,save=FALSE)
cdlist <- loadCdList(resultsDir = chicagoOutputDir, baited_genes = baited_genes)
L <- cdlist[!grep('rep',names(cdlist))]
plotSigIntsStats(L,plotDistribution=FALSE,plotExamples=TRUE) ##this plots example distributions

```

quantifyWithinPeaks *Quantification of Reads and Scores within oneGeneOnePeak or AggregatedPeak Interactions.*

Description

Quantifies reads and scores for interaction tables containing interactions overlapping multiple restriction fragments, such as oneGeneOnePeak or aggregatePeak tables.

The function sums up all read and score information within the interaction, including restriction fragments that are not detected as significant interactions.

Output contains two types of quantifications:

- 1) total sum of reads (raw: `_N` and downsampled: `_N_downsampled`) and scores (`_score`) for each interaction
- 2) normalized quantifications (`_N_downsampled_mean` and `_score_mean`) that normalize for the total number of valid restriction fragments within an interaction peak.

Usage

```

quantifyWithinPeaks(
  data,
  cdlist,
  cdlist.reps = NULL,
  rmapgr,
  minsize = NULL,
  maxsize = NULL,
  file = NULL
)

```

Arguments

| | |
|--------------------------|---|
| <code>data</code> | Table with interactions derived either through <code>oneGeneOnePeak()</code> or <code>aggregatePeaks()</code> . Usually contains the following columns: "intID", "baitID", "otherEndID", "seqnames_bait", "s". minimal requirement: "intID", "baitID", "otherEndID". |
| <code>cdlist</code> | List of ChicagoData tables from which to make quantifications. |
| <code>cdlist.reps</code> | Optional: List of ChicagoData tables from which to make quantifications for individual replicates. |
| <code>rmapgr</code> | GenomicRanges object containing restriction fragments. IDs must be saved in a column called 'id'. |
| <code>minsize</code> | Optional, specifies the minimal size of valid restriction fragments and should correspond to the parameter chosen for creating the ChicagoData objects (<code>mySettings\$minFragLen</code>) within the <code>cdlist</code> . |
| <code>maxsize</code> | Optional, specifies the maximal size of valid restriction fragments and should correspond to the parameter chosen for creating the ChicagoData objects (<code>mySettings\$maxFragLen</code>) within the <code>cdlist</code> . |

file Optional; Path to the file in which the data should be saved. If specified, the final output will be saved under 'file', while up to four intermediate outputs will be saved under file1...4.txt. Note: Output will not be saved if file is not specified.

Value

Interaction table annotated with the sum and mean of read and score counts for each interaction.

Examples

```
# example code

##Create an rmapgr object
rmapgr <- GenomicRanges::GRanges(rep(1,5), IRanges::IRanges( 1:5,2:6 ))
rmapgr$id=1:5

##create data with two interactions
data <- data.frame(intID=c('1;2', '1;3;1;5'),
                  baitID = as.character(c(1,1)),
                  otherEndID=as.character(c(2, '3,5')))

##create a ChicagoData object with two samples
cdlist <- list(
  sample1 = data.frame(baitID=rep(1,4),
                      otherEndID=c(2,3:5),
                      N=c(2,5,10,8),
                      score=c(5, 1, 2, 4)),
  sample2 = data.frame(baitID=rep(1,4),
                      otherEndID=c(2,3:5),
                      N=c(20,3,1,6),
                      score=c(3, 5, 3, 5))
)

##run quantifyWithinPeaks
quantifyWithinPeaks(data, cdlist, rmapgr=rmapgr)
```

rmap2rmapgr

Convert .rmap to the GenomicRanges object 'rmapgr'

Description

Searches for the .rmap Chicago file in designDir and converts it to a GenomicRanges object rmapgr. Only executed if the GenomicRanges object doesn't exist in the design directory yet, otherwise loads the existing file.

Usage

```
rmap2rmapgr(designDir = "designDir", save = FALSE)
```

Arguments

| | |
|-----------|--|
| designDir | Default: 'designDir'; Path to the directory with Chicago design files. |
| save | Default: FALSE; If TRUE, saves the converted object as .rds in the designDir, using the same name as the .rmap file. |

Value

Returns a GenomicRanges object derived from .rmap.

Examples

```
# Example running rmap2rmapgr
extdata <- system.file("extdata", package="PostChicago")
designDir <- file.path(extdata, 'designDir')
rmap2rmapgr(designDir, save=FALSE)
```

runChicagoForPostChicago

Run Chicago and save ChicagoData objects.

Description

Runs Chicago such that the saved output files are compatible with our pipeline. Default settings are only executed if the files are not yet saved in outputDir.

Usage

```
runChicagoForPostChicago(
  df,
  mySettings = NULL,
  outputDir = "results",
  DataPath = "dataPath",
  DesignDir = "designDir",
  overwrite = FALSE
)
```

Arguments

| | |
|------------|---|
| df | data.frame containing at least two columns: 'filename' (.chinput filename) and 'type' (sample type) |
| mySettings | Chicago Settings for the pipeline. If not specified, default Chicago settings will be used. |
| outputDir | Default 'results'; Output directory in which ChicagoData objects are saved. |
| DataPath | Default 'dataPath'; Directory of input files (typically .chinput). |
| DesignDir | Default 'designDir'; Directory of Experiment Design Files, see Chicago documentation. |
| overwrite | Default: FALSE; Overwrites ChicagoData objects that are present in outputDir if set to TRUE. |

Value

Saves complete ChicagoData objects containing score and read counts for all ditags, including non-significant interactions and named after df\$type. Also saves standard Chicago outputs in outputDir.

Examples

```
##Example code using package-supplied data
extdata <- system.file("extdata", package = "PostChicago")
chicagoOutputDir <- file.path(extdata, 'results')
designDir <- file.path(extdata, 'designDir')
dataPath <- file.path(extdata, "dataPath")
chicagoOutputDir <- file.path(extdata, "results")

## checking which .chinput files we have stored:
dir(dataPath)

##Adjust weights settings

weightsPath <- file.path(
  system.file("extdata", package = "Chicago"),
  "weights"
)
weightSettings <- file.path(weightsPath, "mESC-2reps.settings")
weightSettings <- read.delim(weightSettings, header = FALSE)

mySettings <- Chicago::defaultSettings()
mySettings[grep("weight", names(mySettings))] <- weightSettings[, 2]
mySettings$minFragLen <- 100

##Create a table with sample info

fls <- list.files(dataPath)
fls

df <- data.frame(
  filename = fls,
  RA = stringr::str_remove(fls, "_rep.*"),
  rep = stringr::str_remove(stringr::str_remove(fls, "^.*h_"), "\\\\.chinput$")
)
df$type <- df$RA
head(df)

##Now you can run Chicago!

runChicagoForPostChicago(
  df = df, mySettings = mySettings, outputDir = chicagoOutputDir,
  DataPath = dataPath, DesignDir = designDir
)
```

Index

* internal

PostChicago-package, 2

aggregatePeaks, 3

aggregatePeaks_regions, 4

annotateInts, 5

baitmap2baited_genes, 6

boxplotsCapC, 7

getMatrix, 8

loadCdList, 10

loadGr1, 11

makeBedGraphs, 12

makeIntsTable, 13

makeOneGeneOnePeak, 15

makeQCplots, 17

plotAggregatePeaks, 18

plotInteractions, 19

plotSigIntsStats, 22

PostChicago-package, 2

quantifyWithinPeaks, 23

rmap2rmapgr, 24

runChicagoForPostChicago, 25