Package ‘mariner’

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Title       Mariner: Explore the Hi-Cs
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Description Tools for manipulating paired ranges and
             working with Hi-C data in R. Functionality includes
             manipulating/merging paired regions, generating
             paired ranges, extracting/aggregating interactions
             from `.hic` files, and visualizing the results.
             Designed for compatibility with plotgardener for
             visualization.

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mariner-package

Description

‘mariner’ is an R/Bioconductor package for exploring Hi-C data. It enables users to flexibly manipulate, extract, and aggregate chromatin interaction data quickly and efficiently.

Details

Key Features:

- Manipulating Paired Ranges - Convert, bin, and shift paired genomic ranges.
- Clustering & Merging Interactions - Group nearby interactions and select one as representative.
- Extracting & Aggregating Interactions - Pull Hi-C pixels or matrices, then aggregate by files or interactions.
- Calculating Loop Enrichment - Determine loop enrichment to local background with selection functions to flexibility select foreground and background.

For more details on the features of ‘mariner’, read the vignette: ‘browseVignettes(package=“mariner”)’

Author(s)

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See Also

Useful links:

aggHicMatrices

Aggregate count matrices from InteractionArray objects

Description

Aggregation of count matrices is done blocks to avoid large memory usage. Use ‘nBlocks’ to control the number of blocks read into memory at once. Blocks are defined as ‘length(interactions(x))/nBlocks’.

Usage

aggHicMatrices(
  x,
  by = NULL,
  FUN = sum,
  nBlocks = 5,
  verbose = TRUE,
  BPPARAM = bpparam(),
  compressionLevel = 0
)

## S4 method for signature 'InteractionArray'
aggHicMatrices(
  x,
  by = NULL,
  FUN = sum,
  nBlocks = 5,
  verbose = TRUE,
  BPPARAM = bpparam(),
  compressionLevel = 0
)

Arguments

x
  InteractionArray object.

by
  String (length one character vector) describing whether to aggregate by interactions, files, or neither (i.e. NULL as default).

FUN
  Function to use for aggregating.

nBlocks
  Number of blocks for block-processing arrays. Default is 5. Increase this for large datasets. To read and process all data at once, set this value to 1.

verbose
  Boolean (TRUE or FALSE) describing whether to report block-processing progress.

BPPARAM
  Parallelization params (passed to ‘BiocParallel::bplapply’). Default is the result of ‘BiocParallel::bpparams()’. Parallel processing is not available when ‘by=interactions’.

compressionLevel
  Number (length one numeric vector) between 0 (Default) and 9 indicating the compression level used on HDF5 file.
Details

Since interactions are typically the largest dimension in an InteractionArray, using ‘by=interactions’ creates an HDF5-backed array to store these large arrays. Currently parallel processing for HDF5-backed arrays are not supported regardless of the value of ‘BPPARAM’. Both ‘by=NULL’ and ‘by=files’ support parallel processing.

Value

An aggregated ‘DelayedArray’ object. If ‘by=interactions’ or ‘by=files’ then a 3-dimensional ‘DelayedArray’ is returned. If ‘by=NULL’ (default) then A 2-dimensional ‘DelayedMatrix’ is returned.

Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loops as GIinteractions object
loops <-
  WT_5kbLoops.txt() |
    setNames("WT") |
    read.table(header=TRUE) |
    as_ginteractions(keep.extra.columns=FALSE)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(loops) <- 'ENSEMBL'

## Expand pixel ranges with a 5 pixel buffer on either side
loops <-
  binPairs(loops, binSize=100e3) |
    pixelsToMatrices(buffer=5)

## Extract 10, 11x11 count matrices from 2 hic files
iarr <-
  loops[1:10] |
    pullHicMatrices(binSize=100e3, files=hicFiles)

## Aggregate all, by files, or by interactions
aggHicMatrices(x=iarr)
aggHicMatrices(x=iarr, by="files")
aggHicMatrices(x=iarr, by="interactions")
```
aggPairMcols

Aggregate the metadata columns of merged pairs

Description

Aggregate the metadata columns of merged pairs

Usage

aggPairMcols(x, columns, funs)

## S4 method for signature
## 'MergedGInteractions,character,character_OR_function_OR_list'
 aggPairMcols(x, columns, funs)

Arguments

x
MergedGInteractions object.

columns
Character vector of columns to aggregate.

funs
Character vector of functions to apply to 'columns'.

Value

'x' with aggregated metadata columns

Examples

## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

bedpeFiles <- c(
  marinerData::FS_5kbLoops.txt(),
  marinerData::WT_5kbLoops.txt()
)
names(bedpeFiles) <- c("FS", "WT")

## Read in bedpeFiles as a list of GInteractions
## Use only first 1000 rows for fast example
giList <-
  lapply(bedpeFiles, read.table, header=TRUE, nrows=1000) |> 
  lapply(as_ginteractions) |> 
  setNames(gsub("^.*extdata/(.{2}).*$", \1, bedpeFiles))

## Add names describing the source and loop
giList <- lapply(seq_along(giList), \(i\) {
  x <- giList[[i]]
  x$name <- paste0(names(giList)[i], ".loop_", length(x))
})
## Cluster & merge pairs
\[
x \leftarrow \text{mergePairs}(x = \text{giList}, \radius = 5e03)
\]

## List loop names
\[
\text{aggPairMcols}(x, \columns = \text{"name"}, \text{fun} = \text{"list"})
\]

## Aggregate values
\[
\text{aggPairMcols}(x, \columns = \text{c("APScoreAvg"), fun = \text{"mean"})
\]
\[
\text{aggPairMcols}(x, \columns = \text{c("APScoreAvg", "avg"), fun = \text{"mean"})
\]
\[
\text{aggPairMcols}(x, \columns = \text{c("APScoreAvg"), fun = c("mean", "median"))
\]

## Custom functions
\[
\text{aggPairMcols}(x, \columns = \text{c("APScoreAvg"), fun = \lambda(x) \left\{ 
\text{ifelse(is.na(sd(x)), 0, sd(x))}
\right\})
\]

---

### as_ginteractions

**Convert DataFrames to GInteraction objects**

#### Description

`as_ginteractions` takes a paired-interaction (i.e. BEDPE) formatted data-frame-like object and converts it to a GInteractions object. For convenience, `makeGInteractionsFromDataFrame` can be used as an alias.

#### Usage

```r
as_ginteractions(
  df,
  keep.extra.columns = TRUE,
  starts.in.df.are.0based = FALSE,
  ...
)
```

```r
makeGInteractionsFromDataFrame(
  df,
  keep.extra.columns = TRUE,
  starts.in.df.are.0based = FALSE,
  ...
)
```

```
## S4 method for signature
## 'DF_OR_df_OR_dt,logical_OR_missing,logical_OR_missing'
makeGInteractionsFromDataFrame(df, keep.extra.columns, starts.in.df.are.0based)
```
## S4 method for signature
## 'DF_OR_df_OR_dt,logical_OR_missing,logical_OR_missing'
as_ginteractions(df, keep.extra.columns, starts.in.df.are.0based)

**Arguments**

- **df**
  A data.table, data.frame, or DataFrame object. Assumes that the first 6 columns are in the format chr1, start1, end1 and chr2, start2, end2, representing each pair of interactions.

- **keep.extra.columns**
  TRUE or FALSE (the default). If TRUE, the columns in df that are not used to form the genomic ranges of the returned GRanges object are then returned as metadata columns on the object. Otherwise, they are ignored. If df has a width column, then it's always ignored.

- **starts.in.df.are.0based**
  TRUE or FALSE (the default). If TRUE, then the start positions of the genomic ranges in df are considered to be 0-based and are converted to 1-based in the returned GRanges object. This feature is intended to make it more convenient to handle input that contains data obtained from resources using the "0-based start" convention. A notorious example of such resource is the UCSC Table Browser (http://genome.ucsc.edu/cgi-bin/hgTables).

... Additional arguments.

**Value**

GInteraction object

**Examples**

- **data.frame**
  ```r
def <- data.frame(chr1 = "chr1", x1 = 10000, x2 = 20000,
                    chr2 = "chr1", y1 = 30000, y2 = 40000)
makeGInteractionsFromDataFrame(df)
  ```

- **data.frame**
  ```r
def <- data.frame(chr1 = "chr1", x1 = 10000, x2 = 20000,
                    chr2 = "chr1", y1 = 30000, y2 = 40000)
as_ginteractions(df)
  ```

- **data.table**
  ```r
library(data.table)
df <- data.table::data.table(chr1 = "chr1", x1 = 10000, x2 = 20000,
                             chr2 = "chr1", y1 = 30000, y2 = 40000)
as_ginteractions(df)
  ```

- **DataFrame**
  ```r
library(S4Vectors)
df <- DataFrame(chr1 = "chr1", x1 = 10000, x2 = 20000,
                 chr2 = "chr1", y1 = 30000, y2 = 40000)
  ```
as_ginteractions(df)

## Alias
df <- data.frame(chr1 = "chr1", x1 = 10000, x2 = 20000,
                 chr2 = "chr1", y1 = 30000, y2 = 40000,
                 pval = 0.05, dist = 10000)
makeGInteractionsFromDataFrame(df)

## Additional metadata
df <- data.frame(chr1 = "chr1", x1 = 10000, x2 = 20000,
                 chr2 = "chr1", y1 = 30000, y2 = 40000,
                 pval = 0.05, dist = 10000)
as_ginteractions(df)

## Remove additional metadata
as_ginteractions(df, keep.extra.columns = FALSE)

## Add 1 to starts (for 0-based programs)
as_ginteractions(df, starts.in.df.are.0based = TRUE)

---

**binPairs**

*Flexibly bin paired ranges*

**Description**

Paired range objects (like `GInteractions` or BEDPE-formatted `data.frame`-like objects) can be binned separately for each set of ranges.

**Usage**

```r
binPairs(x, binSize, pos1 = "center", pos2 = "center", ...)
```

## S4 method for signature
## 'DF_OR_df_OR_dt, numeric, character_OR_numeric_OR_missing, character_OR_numeric_OR_missing'
```r
binPairs(x, binSize, pos1, pos2)
```

## S4 method for signature
## 'GInteractions, numeric, character_OR_numeric_OR_missing, character_OR_numeric_OR_missing'
```r
binPairs(x, binSize, pos1, pos2)
```
Arguments

x  ‘GInteractions’ or ‘data.frame’-like object with paired interactions.

binSize  Integer (numeric) vector describing the new size of each pair of ranges. Accepts up to 2 values for adjusting each pair.

pos1, pos2  Position within anchors to resize the bin. Can be a character or integer vector of length 1 or ‘length(x)’ designating the position for each element in ‘x’. Character options are “start”, “end” and “center”. Integers are referenced from the start position for ‘+’ and ‘*’ strands and from the end position for the ‘-’ strand.

...  Additional arguments.

Value

GInteractions-like object binned to ‘binSize’ by ‘pos1‘ and ‘pos2‘.

Examples

```r
## Construct interactions as data.frame
df1 <-
data.frame(chr1 = "chr1", x1 = 10000, x2 = 20000,
        chr2 = "chr1", y1 = 30000, y2 = 40000)

## Assign each range to 20-kb bins from the start positions
binPairs(x = df1,
        binSize = 20000,
        pos1 = 'start',
        pos2 = 'start')

## Construct GInteractions
library(InteractionSet)
gi1 <-
data.frame(chr1 = "chr1", x1 = 10000, x2 = 20000,
        chr2 = "chr1", y1 = 30000, y2 = 40000) |> as_ginteractions()

## Assign each range to 20-kb bins from the start positions
binPairs(x = gi1,
        binSize = 20000,
        pos1 = 'start',
        pos2 = 'start')
```

binRanges  *Flexibly bin ranges*

Description

Flexibly bin ranges
Usage

binRanges(x, binSize, pos = "center")

## S4 method for signature 'GRanges,numeric,character_OR_numeric_OR_missing'
binRanges(x, binSize, pos = "center")

Arguments

x 'GRanges' object
binSize Integer (numeric) describing the new size of each range.
pos Position within range to resize the bin. Can be a character or integer vector of length 1 or 'length(x)' designating the position for each element in ‘x’. Character options are "start", "end" and "center". Integers are referenced from the start position for '+' and '*' strands and from the end position for the '-' strand.

Value

'GRanges' object that has been shifted by ‘pos’ and assigned to bins of ‘binSize’.

Examples

library(GenomicRanges)

## Create example GRanges
gr1 <- GRanges(seqnames = "chr1",
   ranges = IRanges(start = rep(5000,3),
   end = rep(6000,3)),
   strand = c('+', '-', '*'))
gr2 <- gr1 |> promoters(upstream = 2000, downstream = 200)

## Binning the results
binRanges(x = gr1, binSize = 1000, pos = 'start')
binRanges(x = gr1, binSize = 1000, pos = 'end')
binRanges(x = gr1, binSize = 1000, pos = 'center')

## Bin after shifting back to TSS
binRanges(x = gr2, binSize = 1000, pos = 2000)

calcLoopEnrichment

Calculate loop enrichment over background.

Description

Pulls Hi-C pixels and calculates the enrichment of the selected foreground ('fg') over the selected background ('bg').
Usage

calcLoopEnrichment(
  x,
  files,
  fg = selectCenterPixel(mhDist = 1, buffer = 5),
  bg = selectTopLeft(n = 4, buffer = 5) + selectBottomRight(n = 4, buffer = 5),
  FUN = function(fg, bg) median(fg + 1)/median(bg + 1),
  nBlocks = 5,
  verbose = TRUE,
  BPPARAM = bpparam(),
  ...
)

## S4 method for signature 'GInteractions,character'
calcLoopEnrichment(
  x,
  files,
  fg = selectCenterPixel(mhDist = 1, buffer = 5),
  bg = selectTopLeft(n = 4, buffer = 5) + selectBottomRight(n = 4, buffer = 5),
  FUN = function(fg, bg) median(fg + 1)/median(bg + 1),
  nBlocks = 5,
  verbose = TRUE,
  BPPARAM = bpparam(),
  ...
)

Arguments

x
  GInteractions object.

files
  Character file paths to '.hic' files.

fg
  Integer vector of matrix indices for the foreground.

bg
  Integer vector of matrix indices for the background.

FUN
  Function taking two parameters (i.e., 'fg', 'bg') defining how enrichment should be calculated. Must produce a single value (numeric of length one).

nBlocks
  Number of blocks for block-processing arrays. Default is 5. Increase this for large datasets. To read and process all data at once, set this value to 1.

verbose
  Boolean (TRUE or FALSE) describing whether to report block-processing progress.

BPPARAM
  Parallelization params (passed to ‘BiocParallel::bplapply’). Default is the result of ‘BiocParallel::bpparams()’. Parallel processing is not available when ‘by=interactions’.

... Additional arguments passed to ‘pullHicMatrices’. See ?['pullHicMatrices'].

Value

A DelayedMatrix of enrichment scores where rows are interactions (i.e. loops) and columns are Hi-C files.
Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loops as GInteractions object
loops <-
  WT_5kbLoops.txt() |> 
    setNames("WT") |> 
    read.table(header=TRUE) |> 
    as_ginteractions(keep.extra.columns=FALSE)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(loops) <- 'ENSEMBL'

## Expand binSize of loops
loops <- binPairs(x=loops, binSize=100e3)

## Calculate loop enrichment
calcLoopEnrichment(x=loops[1:10],
  files=hicFiles)

## Customize different foreground/background
## with selection functions
buffer <- 10 # choose pixel radius around center
fg <- selectCenterPixel(mhDist=seq(0,4), buffer=buffer)
bg <- selectCorners(n=6, buffer=buffer) +
  selectOuter(n=2, buffer=buffer)
calcLoopEnrichment(x=loops[1:10],
  files=hicFiles,
  fg=fg,
  bg=bg)
```

---

**Description**

A GInteractions object containing pixels of interest is resized to the ‘from’ resolution (if its not already), then count matrices are extracted for each interaction and Hi-C file using the new ‘to’
resolution. Count matrices are aggregated by interactions with the supplied 'aggFUN', and a new pixel is selected with the supplied 'selectFUN'. For large datasets, increase 'nBlocks' to allow for smaller blocks of data to be processed in memory.

Usage

changePixelRes(
  x,
  files,
  from,
  to,
  aggFUN = sum,
  selectFUN = "which.max",
  nBlocks = 5,
  verbose = TRUE,
  norm = "KR",
  half = "upper",
  ...
)

## S4 method for signature 'GInteractions,character'
changePixelRes(
  x,
  files,
  from,
  to,
  aggFUN = sum,
  selectFUN = "which.max",
  nBlocks = 5,
  verbose = TRUE,
  norm = "KR",
  half = "upper",
  ...
)

Arguments

x  GInteractions object.
files  Character file paths to ‘.hic’ files.
from  Number (length one numeric vector) describing the resolution of ‘x’. Data will be binned to this value if it is not already binned.
to  Number (length one numeric vector) describing the new resolution for the pixels.
aggFUN  Function to use for aggregating across Hi-C files. Must be passable to ‘which.max’ or ‘which.min’. Default is “sum”.
selectFUN  Function to use for selecting among aggregated interactions. Must be one of "which.max" or "which.min".
nBlocks  Number of blocks for block-processing arrays. Default is 5. Increase this for large datasets. To read and process all data at once, set this value to 1.
changePixelRes

verbose  Boolean (TRUE or FALSE) describing whether to report block-processing progress. Default is TRUE.

norm    String (length one character vector) describing the Hi-C normalization to apply. Use `strawr::readHicNormTypes()` to see accepted values for each file in `files`.

half    String (character vector of length one) indicating whether to keep values for the upper triangular (`half="upper"`) where `start1 < start2`, lower triangular (`half="lower"`) where `start1 > start2`, or both (`half="both"`, default). When `half="upper"` all lower triangular values are `NA`. When `half="lower"` all upper triangular values are `NA`. When `half="both"` there are no `NA` values. For interchromosomal interactions there is no inherent directionality between chromosomes, so data is returned regardless of specified order.

... Additional arguments passed to `pullHicMatrices()`. See ?`pullHicMatrices`.

Value

A GInteractions object with the updated pixel interactions, along with a column with the aggregated max/min value for that pixel.

Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BioCManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loops as GInteractions object
loops <-
  WT_5kbLoops.txt() |
  setNames("WT") |
  read.table(header=TRUE) |
  as_ginteractions(keep.extra.columns=FALSE)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(loops) <- 'ENSEMBL'

## Rebin loops to 2.5e6 resolution
loops <- binPairs(x=loops, binSize=2.5e06)

## Change pixel resolution from 2.5e6 to 500e3
changePixelRes(x=loops[1:5],
  files=hicFiles,
  from=2.5e6,
  to=500e3)
```
CountMatrix-class  CountMatrix Class

Description
A class for displaying dimnames associated with the count matrices resulting from pullHicMatrices() |> counts(showDimnames=TRUE).

Usage
## S4 method for signature 'CountMatrix'
show(object)

Arguments
object A CountMatrix object.

Details
This class is used only for attaching a "show" method.

Value
A CountMatrix object (clone of DelayedArray)
A DelayedArray object with dimnames for the first two dimensions.

Slots
object InteractionArray object

Examples
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK.HEK_PJA27_inter_30.hic(),
  marinerData::LEUK.HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loop pixels as GInteractions object
pixels <-
  WT_5kbLoops.txt() |> 
  setNames("WT") |> 
  read.table(header=TRUE) |> 
  as_ginteractions(keep.extra.columns=FALSE) |>

binPairs(binSize=10^3)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(pixels) <- 'ENSEMBL'

## Expand pixels to regions for pulling
## Hi-C submatrices
regions <- pixelsToMatrices(x=pixels, buffer=5)

## Extract 11x11 count matrices from the
## first 100 regions and 2 Hi-C files
iarr <- pullHicMatrices(x=regions[1:100],
                       files=hicFiles,
                       binSize=10^3)

## Display the start bin of each
## interaction in the count
## matrices
counts(iarr, showDimnames=TRUE)

---

counts,InteractionArray-method

```
Access count matrices from InteractionArray or InteractionMatrix
```

Description

Access count matrices from InteractionArray or InteractionMatrix
Access count matrices from InteractionArray or InteractionMatrix
Replace method for counts

Usage

```
## S4 method for signature 'InteractionArray'
counts(object, showDimnames = FALSE)

## S4 method for signature 'InteractionMatrix'
counts(object)

## S4 replacement method for signature 'InteractionMatrix'
counts(object) <- value
```

Arguments

- **object**  
  InteractionMatrix object
- **showDimnames**  
  Logical vector of length-one indicating whether to show dimensions of count matrices (default FALSE). Only applicable for InteractionArray objects.
- **value**  
  Value for replacement
Value

For InteractionArray, a 4-dimensional DelayedArray of Hi-C submatrices is returned with the following dimensions: rows of count matrix, columns of count matrix, Interactions in ‘object’, Hi-C ‘files’.

For InteractionMatrix, a 2-dimensional DelayedArray is returned with rows representing interactions in ‘object’ and columns for each Hi-C file in ‘files’.

For InteractionMatrix, the replace matrix replaces the counts assay with matrix-like objects supplied in ‘value’.

Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

######################################
## Accessing Hi-C count submatrices ##
######################################

## Create example interactions
x <- read.table(text="
  9 14435000 14490000 9 14740000 14795000
  9 89540000 89595000 9 89785000 89840000
  9 23700000 23755000 9 23760000 23815000"
)
x <- as_ginteractions(x)

## Extract 3, 11x11 count matrices from 2 hic files
iarr <- pullHicMatrices(x, hicFiles, 5e03)

## Access count matrices
counts(iarr)
counts(iarr, FALSE)

## Create example interactions
x <- read.table(text="
  9 14000000 14500000 9 14500000 24000000
  9 89500000 90000000 9 89500000 24000000
  9 23500000 24000000 9 23500000 24000000"
)
x <- as_ginteractions(x)
```
## Extract 3 pixels from 2 hic files
mat <- pullHicPixels(x, hicFiles, 500e03)

## Access count matrix
counts(mat)

#################################
## Replacing Hi-C count matrix ##
#################################

## Realize as in-memory matrix
counts(mat) <- as.matrix(counts(mat))
counts(mat)

DelegatingGIInteractions-class

Virtual class for delegating GIInteractions

Description

Uses a delegate ‘GIInteractions’ object during initialization to assign its ‘GIInteractions’ slots.

Value

DelegatingGIInteractions virtual class

Slots

delegate  A ‘GIInteractions’ object used to initialize ‘GIInteractions’-specific slots.

See Also

[InteractionSet::GIInteractions]

getPairClusters  Get clustered pairs from MergedGIInteractions object

Description

Returns the clustered pairs associated with each range in the ‘MergedGIInteractions’ object. Order always follows the indices of the ‘MergedGIInteractions’ object.
Usage

getPairClusters(x, ...)

## S4 method for signature 'MergedGInteractions'
getPairClusters(x)

Arguments

x

MergedGInteractions object.

...

Additional arguments.

Value

A list of data.tables corresponding to each pair in 'x'.

Examples

```r
## Load required packages
library(data.table, include.only="fread")

## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Reference BEDPE files (loops called with SIP)
bedpeFiles <- c(
  marinerData::FS_5kbLoops.txt(),
  marinerData::WT_5kbLoops.txt()
)
names(bedpeFiles) <- c("FS", "WT")

## Read in bedpeFiles as a list of GInteractions
## Use only first 1000 rows for fast example
giList <-
  lapply(bedpeFiles, fread, nrows = 1000) |> lapply(as_ginteractions)

## Cluster & merge pairs
x <- mergePairs(x = giList,
  radius = 10e03,
  column = "APScoreAvg")

## Access pair clusters
getPairClusters(x[1:3])
getPairClusters(x[3:1])
getPairClusters(x[c(3, 1, 2)])
gapPairClusters(x) |> length()
```
hdf5BlockApply

HDF5-backed blockApply

Description

Read in array data in blocks, apply function, and write back to an HDF5 file.

Usage

hdf5BlockApply(x, FUN, sink, grid, sink_grid, verbose = TRUE)

## S4 method for signature 'DelayedArray'

hdf5BlockApply(x, FUN, sink, grid, sink_grid, verbose = TRUE)

Arguments

x
  Delayed Array object.

FUN
  Function that takes one argument 'block' and processes it.

sink
  HDF5RealizationSink object.

grid
  ArrayGrid over array 'x'.

sink_grid
  ArrayGrid over 'sink'.

verbose
  Logical - whether block processing progress should be displayed.

Details

Implements an HDF5-backed option for block processing on DelayedArray objects.

Value

An HDF5Array object.

Examples

## This function is intended for advanced users.
## To learn more about using DelayedArray
## or HDF5-backed objects, see ?DelayedArray or
## ?HDF5Array

library(DelayedArray)
library(HDF5Array)
library(rhdf5)

dl <- c(11L, 11L, 100L, 2L)

a <- array(data=seq(1, prod(dims)), dim=dims)
a <- DelayedArray(a)

## Define spacings, breaking up the longest dim
## Here we are processing in blocks of 10
spacings <- dim(a)

## Define storage dimensions (all except those
## over which the function is being applied)
storageDims <- dims[c(1,2,3)]

## Define chunk dimensions for writing to HDF5
chunkDims <- storageDims

## Create grid for applying the data (grid)
## and grid for writing to the sink (sink_grid)
grid <- RegularArrayGrid(dims, spacings)
sink_grid <- RegularArrayGrid(storageDims, chunkDims)

## Create HDF5 file for writing
h5 <- tempfile(fileext = ".h5")
h5createFile(h5)

## Define compression for HDF5
compressionLevel <- 0

## Create HDF5-backed realization sink
sink <- HDF5RealizationSink(filepath=h5,
  name="counts",
  type="integer",
  dim=storageDims,
  chunkdim=chunkDims,
  level=compressionLevel)

## Wrap function that operates on each block
## this can be anything, here it is sum
FUN <- \(block) apply(block, c(1,2,3), sum)

## Read, apply, and write to HDF5
ans <- hdf5BlockApply(x=a,
  FUN=FUN,
  sink=sink,
  grid=grid,
  sink_grid=sink_grid,
  verbose=TRUE)

ans
**initialize,DelegatingGInteractions-method**

*Initialization method for DelegatingGInteractions*

---

**Description**

Initialization method for DelegatingGInteractions

**Usage**

```r
## S4 method for signature 'DelegatingGInteractions'
initialize(.Object, ..., delegate = GInteractions())
```

**Arguments**

- `.Object`: GInteractions object
- `...`: Additional arguments
- `delegate`: GInteractions object

**Value**

DelegatingGInteractions object

---

**initialize,MergedGInteractions-method**

*Initialize MergedGInteractions*

---

**Description**

Initialize MergedGInteractions

**Usage**

```r
## S4 method for signature 'MergedGInteractions'
initialize(.Object, ..., delegate = GInteractions())
```

**Arguments**

- `.Object`: GInteractions object
- `...`: Additional arguments
- `delegate`: GInteractions object

**Value**

MergedGInteractions
InteractionArray-class  

InteractionArray Class

Description

The `InteractionArray` class extends `InteractionSet` to provide an interface for accessing submatrices pulled from Hi-C data.

Usage

```
InteractionArray(assays, interactions, ...)
```

## S4 method for signature 'ANY,GInteractions'

```
InteractionArray(assays, interactions, ...)
```

## S4 method for signature 'missing,missing'

```
InteractionArray(assays, interactions, ...)
```

## S4 method for signature 'InteractionArray'

```
show(object)
```

## S4 method for signature 'InteractionArray'

```
rbind(..., deparse.level = 1)
```

## S4 method for signature 'InteractionArray'

```
cbind(..., deparse.level = 1)
```

Arguments

- `assays`, `interactions`
  See `?InteractionSet`

- `...`
  InteractionArray objects to be combined column-wise. All objects must be the same class.

- `object`
  InteractionArray object.

- `deparse.level`
  An integer scalar; see `?base::cbind` for a description of this argument.

Details

This class is constructed with the `pullHicMatrices()` function when all paired ranges have equal dimensions.

Value

An InteractionArray (see description)
Description

The `InteractionMatrix` class extends the `InteractionSet` to provide an interface for accessing the count matrix pulled from Hi-C data.

Usage

```r
InteractionMatrix(assays, interactions, ...) 
## S4 method for signature 'ANY,GInteractions'
InteractionMatrix(assays, interactions, ...) 
## S4 method for signature 'missing,missing'
InteractionMatrix(assays, interactions, ...) 
## S4 method for signature 'InteractionMatrix'
show(object) 
## S4 method for signature 'InteractionMatrix'
rbind(..., deparse.level = 1) 
## S4 method for signature 'InteractionMatrix'
cbind(..., deparse.level = 1)
```

Arguments

- `assays, interactions`
  
  See `?InteractionSet`

- `...` InteractionMatrix objects to be combined column-wise. All objects must be the same class.

- `object` InteractionMatrix object.

- `deparse.level` An integer scalar; see `?base::cbind` for a description of this argument.
MatrixSelection-class

Details

This class is constructed with the ‘pullHicPixels()’ function when all paired ranges define a single pixel.

Value

An InteractionMatrix (see description)

See Also

[InteractionSet::InteractionSet]

Examples

InteractionMatrix()

Description

An object containing the selected indices of a matrix.

Value

A MatrixSelection object (see description)

Slots

- `x` Vector of selected indices from a matrix of `dim = buffer*2+1`.
- `buffer` Integer indicating the buffer size, or number of pixels around a matrix.

Examples

selectCenterPixel(0, 5)
Description

The ‘MergedGIInteractions’ class extends the ‘GIInteractions’ to contain additional information about the pairs being merged.

Details

The ‘MergedGIInteractions’ class uses a delegate object during initialization to assign its ‘GIInteractions’ slots. In addition to containing information from all pairs, it also behaves as a ‘GIInteractions’ object. ‘mergePairs()’ builds this object.

Value

A MergedGIInteractions object (see description)

Slots

delegate A ‘GIInteractions’ object used to initialize ‘GIInteractions’-specific slots. This is the mergedPairs set of interactions.

dbscan

A 'data.table' containing all input pairs combined. Also contains all metadata for each pair and 1) the source of the file, 2) an id, 3) which chromosome pair it belongs to (i.e. ‘grp’), and 4) the assigned cluster from ‘dbscan’ (i.e. ‘clst’).

selectionMethod Character describing which method was used to select the final pair from the cluster of merged pairs.

See Also

[InteractionSet::GIInteractions]

Examples

```r
## Load required packages
library(data.table, include.only="fread")

## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Reference BEDPE files (loops called with SIP)
bedpefiles <- c(
  marinerData::FS_5kbLoops.txt(),
```
mergePairs

Merge sets of paired interactions

Description

Sets of paired range objects (i.e., ‘GInteractions’) are first clustered by genomic distance with ‘db-scan’, then a representative interaction is selected for each cluster.

Usage

mergePairs(
  x,
  radius,
  method = "manhattan",
  column = NULL,
  selectMax = TRUE,
  pos = "center"
)

## S4 method for signature 'list_OR_SimpleList_OR_GInteractions,numeric'
mergePairs(
  x,
  radius,
  method = "manhattan",
  column = NULL,
  selectMax = TRUE,
  pos = "center"
)
**mergePairs**

**Arguments**

- **x**: List of `GInteractions` or `data.frame`-like objects.
- **radius**: Numeric describing the distance in base pairs used to define a cluster or pairs.
- **method**: Character describing the distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". Any unambiguous substring can be given. Default is "manhattan".
- **column**: Character denoting the column to be used to select among clustered interactions.
- **selectMax**: Logical. TRUE (default) uses `which.max()` to select the interaction pair. FALSE uses `which.min()`. Only applicable when `column` is specified.
- **pos**: Positions used for clustering pairs. Must be one of "start", "end" or "center". Default is "center".

**Details**

Interactions are clustered into groups using the provided base pair `radius`, and distance `method` with `dbscan()`. Representative interactions are selected for each group by one of two methods. If `column` and `selectMax` arguments are provided, the representative interaction with the maximum (or minimum) value in `column` is returned for each cluster. If these parameters are missing, new ranges for each pair are returned by calculating the median of modes for each cluster.

**Value**

Returns a `MergedGInteractions` object.

**Examples**

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

bedpeFiles <- c(
  marinerData::FS_5kbLoops.txt(),
  marinerData::WT_5kbLoops.txt() )
names(bedpeFiles) <- c("FS", "WT")

## Read in bedpeFiles as a list of GInteractions
## Use only first 1000 rows for fast example
giList <-
  lapply(bedpeFiles, read.table, header=TRUE, nrows=1000) |>
  lapply(as_ginteractions)

## Cluster & merge pairs
x <- mergePairs(x = giList,
  radius = 10e03,
  column = "APScoreAvg")
x
```
names.MatrixSelection  Names S3 method for autocomplete

Description

Names S3 method for autocomplete

Extract `$` operator for MatrixSelection

Extract `[[` operator for MatrixSelection

Concatenate MatrixSelection objects

Concatenate MatrixSelection objects

Usage

```r
## S3 method for class 'MatrixSelection'
names(x)

## S4 method for signature 'MatrixSelection'
x$name

## S4 method for signature 'MatrixSelection,ANY,ANY'
x[[i]]

## S4 method for signature 'MatrixSelection,MatrixSelection'
e1 + e2

## S4 method for signature 'MatrixSelection,MatrixSelection'
e1 - e2
```

Arguments

- `x` A `MatrixSelection` object.
- `name` Name of slot.
- `i` A character or numeric to extract.
- `object` A `MatrixSelection` object.

Value

different data or metadata concerning a MatrixSelection object
parallel_slot_names.MergedGInteractions-method

Method for parallel slots for MergedGInteractions

Description

Method for parallel slots for MergedGInteractions

Usage

```r
## S4 method for signature 'MergedGInteractions'
parallel_slot_names(x)
```

Arguments

- `x` See ?parallel_slot_names

Value

MergedGInteractions with parallel slots

path,InteractionMatrix-method

Accessor for h5File path from an InteractionMatrix

Description

Returns the file path describing where the on-disk HDF5 data associated with the InteractionMatrix object is stored.

This method circumvents the `assays<-` and `path<-` methods for updating the HDF5 path because they are not accessible when the file path is broken.

Usage

```r
## S4 method for signature 'InteractionMatrix'
path(object)
```

```r
## S4 replacement method for signature 'InteractionMatrix'
path(object) <- value
```

Arguments

- `object` InteractionMatrix object
- `value` String (length-one character vector) to use for path replacement.
Details

If the file no longer exists, the path is returned along with a warning. This allows the file path to be updated even if the original linked data no longer exists.

Value

The path to the HDF5 file associated with the InteractionMatrix object.

Updates path to HDF5 file for the InteractionMatrix object.

Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Accessing path to HDF5 data

## Create example interactions
x <- read.table(text="
9 14000000 14500000 9 14500000 15000000
9 89500000 90000000 9 89500000 90000000
9 23500000 24000000 9 23500000 24000000"
)x <- as_ginteractions(x)

## Extract 3 pixels from 2 hic files
imat <- pullHicPixels(x, hicFiles, 500e03)

## Access path
path(imat)
```

```r
## Updating path to HDF5 data

## Create example interactions
x <- read.table(text="
9 14000000 14500000 9 14500000 15000000
9 89500000 90000000 9 89500000 90000000
9 23500000 24000000 9 23500000 24000000"
)x <- as_ginteractions(x)

## Extract 3 pixels from 2 hic files
```
h5File <- tempfile(fileext=".h5")
imat <- pullHicPixels(x, hicFiles, 500e03, h5File=h5File)

## Move file to new location
newFile <- tempfile(fileext="_new.h5")
file.rename(from=h5File, to=newFile)

## Update path
path(imat) <- newFile
path(imat)

---

**pixelsToMatrices**

*Expand pixels to submatrices*

**Description**

Pixels are defined as paired-ranges with starts & ends equal to their ‘binSize’. This function takes GIInteractions fitting this description and expands the ranges such that there is a ‘buffer’ of pixels around each range.

**Usage**

pixelsToMatrices(x, buffer)

## S4 method for signature 'GIInteractions,numeric'
pixelsToMatrices(x, buffer)

**Arguments**

- **x**: GIInteractions object.
- **buffer**: Number (length one numeric vector) of pixels around the pixels in ‘x’.

**Details**

For example, a buffer of 3 would return a GIInteractions object with 3 pixels surrounding the original pixel ranges.

After using ‘pullHicMatrices()’, the result will return a matrix of row and column dimensions of buffer*2+1.

Note, this function does not handle out-of-bound ranges.

**Value**

‘x’ with updated ranges.
## Define example 100bp pixel
library(InteractionSet)

```r
pixel <- GInteractions(
    anchor1=GRanges("chr1:500-600"),
    anchor2=GRanges("chr1:2000-2100")
)
```

## Expand pixel to matrix with
## 3 pixels surrounding the center
## pixel

```r
region <- pixelsToMatrices(x=pixel, buffer=3)
```

## Examples

```r
plotEnrichment

Adjust loop enrichment to remove distance-dependent effect.
```

### Description

Adjust loop enrichment to remove distance-dependent effect.

Show diagnostic plot of loop enrichment before and after distance adjustment.

### Usage

```r
plotEnrichment(scores, interactions, k = 25, nknots = 10, plot = TRUE)
```

```r
adjustEnrichment(x, interactions, k = 25, nknots = 10)
```

```r
## S4 method for signature 'DelayedMatrix_OR_matrix,GInteractions'
adjustEnrichment(x, interactions, k = 25, nknots = 10)
```

```r
## S4 method for signature 'numeric,GInteractions'
plotEnrichment(scores, interactions, k = 25, nknots = 10, plot = TRUE)
```

### Arguments

- **scores**: Numeric vector of enrichment scores.
- **interactions**: A GInteractions Object containing the interactions used to calculate enrichment scores.
- **k**: Number of observations for rolling window.
- **nknots**: integer or function giving the number of knots to use see `?smooth.spline` for more info.
- **plot**: Boolean (default=FALSE), of whether to show diagnostic plot.
- **x**: A DelayedMatrix or matrix with enrichment scores.
Value

A DelayedMatrix of enrichment scores where rows are loops and columns are Hi-C files. A plot (and associated data) for visualizing loop enrichment before and after distance adjustment.

Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loops as GIinteractions object
loops <-
  WT_5kbLoops.txt() |> setNames("WT") |> read.table(header=TRUE, nrows=1000) |> as_ginteractions(keep.extra.columns=FALSE)

## Removes the "chr" prefix for compatibility with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(loops) <- 'ENSEMBL'

## Calculate loop enrichment
enrich <- calcLoopEnrichment(
  x=binPairs(loops, 100e03),
  files=hicFiles)

adjustEnrichment(enrich, loops)

plotEnrichment(enrich[,1], loops)
```

Description

Used to plot single or aggregate matrix such as aggregate peak analysis.
Usage

plotMatrix(
  data,
  params = NULL,
  x = NULL,
  y = NULL,
  width = NULL,
  height = NULL,
  just = c("left", "top"),
  default.units = "inches",
  draw = TRUE,
  palette = colorRampPalette(RColorBrewer::brewer.pal(9, "YlGnBu")),
  zrange = NULL,
  na.color = "grey"
)

## S4 method for signature 'DelayedMatrix_OR_matrix'
plotMatrix(
  data,
  params = NULL,
  x = NULL,
  y = NULL,
  width = NULL,
  height = NULL,
  just = c("left", "top"),
  default.units = "inches",
  draw = TRUE,
  palette = colorRampPalette(RColorBrewer::brewer.pal(9, "YlGnBu")),
  zrange = NULL,
  na.color = "grey"
)

Arguments

data 'DelayedMatrix', 'matrix', list of matrices, or 3 column 'data.frame' of APA results.
params Optional 'pgParams' object containing relevant function parameters.
x Numeric or unit object specifying the x-location of plot.
y Numeric or unit object specifying the y-location of plot.
width Numeric or unit object specifying the width of plot.
height Numeric or unit object specifying the height of plot.
just String or numeric vector specifying the justification of the viewport relative to its (x, y) location.
default.units String indicating the default units to use if 'x', 'y', 'width', or 'height' are only given as numeric vectors.
draw Logical value indicating whether graphics output should be produced.
`plotMatrix`

- **palette**
  - 'colorRampPalette' function to use for mapping values to colors.
- **zrange**
  - Vector of length 2; max and min values to set color scale
- **na.color**
  - String indicating the color to use for mapping NA values.

**Value**

Function will draw a color-mapped matrix and return an S3 object of class ‘MatrixPlot’.

**Examples**

```r
library(plotgardener)
library(RColorBrewer)

## Create divergent matrix ####
m <- matrix(data=rnorm(n=21*21, mean=0, sd=2), nrow=21, ncol=21)

## Define parameters
p <- pgParams(width=3, height=3, default.units="inches")

## Create page
pageCreate(params=p)

## Plot apa
plot <- plotMatrix(data=m,
                    x=p$width/2,
                    y=p$height/2,
                    width=p$width*0.5, height = p$width*0.5,
                    just=c("center", "center"),
                    palette=colorRampPalette(c("blue", "white", "red")),
                    zrange=NULL)

## Annotate legend
annoHeatmapLegend(plot=plot,
                   x=2.3,
                   y=0.75,
                   width=0.1,
                   height=0.75)

## Create sequential matrix
m <- matrix(data=sample(0:100, 21*21, replace=TRUE), nrow=21, ncol=21)

## Define parameters
p <- pgParams(width=3, height=3, default.units="inches")

## Create page
pageCreate(params=p)

## Plot apa
plot <- plotMatrix(data=m,
                    x=p$width/2,
                    y=p$height/2,
                    width=p$width*0.5, height = p$width*0.5,
                    just=c("center", "center"),
                    palette=colorRampPalette(c("blue", "white", "red")),
                    zrange=NULL)
```

## Annotate legend
```r
annoHeatmapLegend(plot=plot,
                   x=2.3,
                   y=0.75,
                   width=0.1,
                   height=0.75)
```
pullHicMatrices

Pull submatrices from `.hic` files

Description

Pull submatrices from `.hic` files

Usage

```r
pullHicMatrices(
  x,
  files,
  binSize,
  ...,
  h5File = tempfile(fileext = "h5"),
  half = "both",
  norm = "NONE",
  matrix = "observed",
  blockSize = 248956422,
  onDisk = TRUE,
  compressionLevel = 0,
  chunkSize = 1
)
```

```r
## S4 method for signature 'GInteractions,character,numeric'
pullHicMatrices(
  x,
  files,
  binSize,
  h5File,
  half,
  norm,
  matrix,
  blockSize,
  ...,
)
pullHicMatrices

onDisk, compressionLevel, chunkSize
)

Arguments

x GInteractions object containing interactions to extract from Hi-C files.
files Character file paths to `.hic` files.
binSize Integer (numeric) describing the resolution (range widths) of the paired data.
... Additional arguments.
h5File Character file path to save `.h5` file.
half String (character vector of length one) indicating whether to keep values for the upper triangular (`half="upper"`) where `start1 < start2`, lower triangular (`half="lower"`) where `start1 > start2`, or both (`half="both"`, default). When `half="upper"` all lower triangular values are `NA`. When `half="lower"` all upper triangular values are `NA`. When `half="both"` there are no `NA` values. For interchromosomal interactions there is no inherent directionality between chromosomes, so data is returned regardless of specified order.
norm String (length one character vector) describing the Hi-C normalization to apply. Use `strawr::readHicNormTypes()` to see accepted values for each file in `files`.
matrix String (length one character vector) Type of matrix to extract. Must be one of "observed", "oe", or "expected". "observed" is observed counts, "oe" is observed/expected counts, "expected" is expected counts.
blockSize Number (length one numeric vector) describing the size in base-pairs to pull from each `.hic` file. Default is 248956422 (the length of the longest chromosome in the human hg38 genome). For large `.hic` files `blockSize` can be reduced to conserve the amount of data read in at a time. Larger `blockSize` values speed up performance, but use more memory.
onDisk Boolean (length one logical vector that is not NA) indicating whether extracted data should be stored on disk in an HDF5 file. Default is TRUE.
compressionLevel Number (length one numeric vector) between 0 (Default) and 9 indicating the compression level used on HDF5 file.
chunkSize Number (length one numeric vector) indicating how many values of `x` to chunk for each write to HDF5 stored data. This has downstream implications for accessing subsets later. For small `compressionLevel` values use smaller `chunkSize` values and for large `compressionLevel` values use large (i.e. `length(x)`) values to improve performance.

Value

InteractionSet object with a 4-dimensional array of Hi-C submatrices, rownames, and colnames. Array is stored with the following dimensions: Interactions in `x`, Hi-C `files`, rows of submatrix, columns of submatrix. The submatrices returned have rows cooresponding to anchor1 of `x` and columns correspond to anchor2 of `x`.
Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK.HEK_PJA27_inter_30.hic(),
  marinerData::LEUK.HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loop pixels as GIInteractions object
pixels <-
  WT_5kbLoops.txt() |> 
    setNames("WT") |> 
    read.table(header=TRUE) |> 
    as_ginteractions(keep.extra.columns=FALSE) |> 
    binPairs(binSize=100e3)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(pixels) <- 'ENSEMBL'

## Expand pixels to regions for pulling
## Hi-C submatrices
regions <- pixelsToMatrices(x=pixels, buffer=5)

## Extract 11x11 count matrices from the
## first 100 regions and 2 Hi-C files
iarr <- pullHicMatrices(x=regions[1:100],
  files=hicFiles,
  binSize=100e3)
iarr

## Access count matrices
counts(iarr)

## Display the start bin of each
## interaction in the count
## matrices
counts(iarr, showDimnames=TRUE)
```

## pullHicPixels

Pull contact frequency from `.hic` files

### Description

Pull contact frequency from `.hic` files.
pullHicPixels

Usage

pullHicPixels(
  x,
  files,
  binSize,
  ...
  h5File = tempfile(fileext = ".h5"),
  half = "both",
  norm = "NONE",
  matrix = "observed",
  blockSize = 248956422,
  onDisk = TRUE,
  compressionLevel = 0,
  chunkSize = 1
)

## S4 method for signature 'GInteractions,character,numeric'
pullHicPixels(
  x,
  files,
  binSize,
  h5File,
  half,
  norm,
  matrix,
  blockSize,
  onDisk,
  compressionLevel,
  chunkSize
)

Arguments

x     GInteractions object containing interactions to extract from Hi-C files.
files  Character file paths to '.hic' files.
binSize  Integer (numeric) describing the resolution (range widths) of the paired data.
...  Additional arguments.
h5File  Character file path to save '.h5' file.
half  String (character vector of length one) indicating whether to keep values for the upper triangular ("half="upper") where 'start1 < start2', lower triangular ("half="lower") where 'start1 > start2', or both ('half="both", default). When 'half="upper"' all lower triangular values are 'NA'. When 'half="lower"' all upper triangular values are 'NA'. When 'half="both"' there are no 'NA' values. For interchromosomal interactions there is no inherent directionality between chromosomes, so data is returned regardless of specified order.
norm  String (length one character vector) describing the Hi-C normalization to apply. Use 'strawr::readHicNormTypes()' to see accepted values for each file in 'files'.

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pullHicPixels

matrix

String (length one character vector) Type of matrix to extract. Must be one of "observed", "oe", or "expected". "observed" is observed counts, "oe" is observed/expected counts, "expected" is expected counts.

blockSize

Number (length one numeric vector) describing the size in base-pairs to pull from each `.hic` file. Default is 248956422 (the length of the longest chromosome in the human hg38 genome). For large `.hic` files `blockSize` can be reduced to conserve the amount of data read in at a time. Larger `blockSize` values speed up performance, but use more memory.

onDisk

Boolean (length one logical vector that is not NA) indicating whether extracted data should be stored on disk in an HDF5 file. Default is TRUE.

compressionLevel

Number (length one numeric vector) between 0 (Default) and 9 indicating the compression level used on HDF5 file.

chunkSize

Number (length one numeric vector) indicating how many values of `x` to chunk for each write to HDF5 stored data. This has downstream implications for accessing subsets later. For small `compressionLevel` values use smaller `chunkSize` values and for large `compressionLevel` values use large (i.e. `length(x)`) values to improve performance.

Value

InteractionSet object with a 2-dimensional array of Hi-C interactions (rows) and Hi-C sample (columns).

Examples

```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loop pixels as GInteractions object
pixels <-
  WT_5kbLoops.txt() |>
  setNames("WT") |>
  read.table(header=TRUE) |>
  as_ginteractions(keep.extra.columns=FALSE) |>
  binPairs(binSize=100e3)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(pixels) <- 'ENSEMBL'

## Extract the first 100 pixels
```
imat <- pullHicPixels(x=pixels[1:100],
                      files=hiCFiles,
                      binSize=100e3)
imat

## Access count matrix
counts(imat)

removeShortPairs

Remove interactions that would cross the Hi-C diagonal or a specified
distance from the diagonal.

Description

Note this is only applies to intrachromosomal pairs, as pair distance is meaningless for interchromosomal pairs. Therefore, all interchromosomal pairs are kept.

Usage

removeShortPairs(x, padding = 0)

## S4 method for signature 'GInteractions'
removeShortPairs(x, padding = 0)

Arguments

x A GInteractions object.
padding Minimum distance away from the diagonal.

Value

A GInteractions object with the short pairs removed.

Examples

## Example GInteractions object
gi <- as.ginteractions(read.table(
  text="
  seqnames1 start1 end1 seqnames2 start2 end2 keep
  chr1 300 400 chr1 300 400 'no'
  chr1 100 200 chr1 300 400 'yes'
  chr1 300 400 chr1 100 200 'yes'
  chr1 300 400 chr2 300 400 'yes'
  chr1 250 350 chr1 300 400 'only_with_padding_50'
  chr1 300 400 chr1 250 350 'only_with_padding_50'
  ",
  header=TRUE
))
### Remove pairs that would cross the diagonal
removeShortPairs(gi)

### Add 50bp of padding
removeShortPairs(gi, padding=50)

---

#### selectionMethod

Get `selectionMethod` from `MergedGInteractions` object

**Description**

Get `selectionMethod` from `MergedGInteractions` object

**Usage**

```
selectionMethod(x, ...)
```

### S4 method for signature 'MergedGInteractions'

```
selectionMethod(x, ...)
```

**Arguments**

- `x` MergedGInteractions object.
- `...` Additional arguments.

**Value**

A character vector describing which selection method was used for merging.

**Examples**

```r
## Load required packages
library(data.table, include.only="fread")

## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BioCManager::install("marinerData")

## Reference BEDPE files (loops called with SIP)
bedpeFiles <- c(
  marinerData::FS_5kbLoops.txt(),
  marinerData::WT_5kbLoops.txt()
)
names(bedpeFiles) <- c("FS", "WT")

## Read in bedpeFiles as a list of GInteractions
## Use only first 1000 rows for fast example
```
selectPixel <- lapply(bedpeFiles, fread, nrows=1000) |> lapply(as_ginteractions)

## Cluster & merge pairs
x <- mergePairs(x = giList,
                 radius = 10e03,
                 column = "APScoreAvg")

selectionMethod(x)

---

selectPixel

Get the pixel representing the strongest or weakest interaction in an InteractionArray

Description

Get the pixel representing the strongest or weakest interaction in an InteractionArray

Usage

selectPixel(
  x,
  aggFUN = sum,
  selectFUN = "which.max",
  nBlocks = 5,
  verbose = TRUE
)

## S4 method for signature 'InteractionArray'
selectPixel(
  x,
  aggFUN = sum,
  selectFUN = "which.max",
  nBlocks = 5,
  verbose = TRUE
)

Arguments

x InteractionArray object

aggFUN Function to use for aggregating across Hi-C files. Must be passable to 'which.max' or 'which.min'. Default is "sum".

selectFUN Function to use for selecting among aggregated interactions. Must be one of "which.max" or "which.min".
nBlocks Number of blocks for block-processing arrays. Default is 5. Increase this for large datasets. To read and process all data at once, set this value to 1.

verbose Boolean (TRUE or FALSE) describing whether to report block-processing progress. Default is TRUE.

Value
A GInteractions object with the updated pixel interactions, along with a column with the aggregated max/min value for that pixel.

Examples
```r
## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Read .hic file paths
hicFiles <- c(
  marinerData::LEUK_HEK_PJA27_inter_30.hic(),
  marinerData::LEUK_HEK_PJA30_inter_30.hic()
)
names(hicFiles) <- c("FS", "WT")

## Read in loops as GInteractions object
loops <-
  WT_5kbLoops.txt() |> 
  setNames("WT") |>
  read.table(header=TRUE) |> 
  as_ginteractions(keep.extra.columns=FALSE)

## Removes the "chr" prefix for compatibility
## with the preprocessed hic files
GenomeInfoDb::seqlevelsStyle(loops) <- 'ENSEMBL'

## Rebin loops to 2.5e6 resolution
loops <- binPairs(x=loops, binSize=2.5e06)

## Pull 5x5 matrices
iarr <- pullHicMatrices(x=loops[1:5],
  files=hicFiles,
  binSize=500e3,
  norm="KR",
  half='upper')

## Select pixel
selectPixel(iarr)
```
selectRadius

Visualize selection for a MatrixSelection object

Description

Note: that buffer must be the same as the selection functions to work appropriately
For `selectCoordinates`, `rowInd` and `colInd` are paired such that the selected position in the
matrix is `c(rowInd[1:i], colInd[1:j])` for `i` rows and `j` columns.

Usage

```r
selectRadius(x, buffer, invert = FALSE)
selectCenterPixel(mhDist, buffer, invert = FALSE)
selectSubmatrix(m, invert = FALSE)
selectCoordinates(rowInd, colInd, buffer, invert = FALSE)
selectBlock(rowInd, colInd, buffer, invert = FALSE)
selectTopLeft(n, buffer, inset = 0, invert = FALSE)
selectTopRight(n, buffer, inset = 0, invert = FALSE)
selectBottomRight(n, buffer, inset = 0, invert = FALSE)
selectBottomLeft(n, buffer, inset = 0, invert = FALSE)
selectCorners(n, buffer, inset = 0, invert = FALSE)
selectRows(rows, buffer, invert = FALSE)
selectCols(cols, buffer, invert = FALSE)
selectInner(n, buffer, invert = FALSE)
selectOuter(n, buffer, invert = FALSE)
```

## S4 method for signature 'MatrixSelection'
show(object)

## S4 method for signature 'numeric'
selectRadius(x, buffer, invert = FALSE)

## S4 method for signature 'numeric'
selectCenterPixel(mhDist, buffer, invert = FALSE)

## S4 method for signature 'matrix'
selectSubmatrix(m, invert = FALSE)

## S4 method for signature 'numeric'
selectCoordinates(rowInd, colInd, buffer, invert = FALSE)

## S4 method for signature 'numeric'
selectBlock(rowInd, colInd, buffer, invert = FALSE)

## S4 method for signature 'numeric'
selectTopLeft(n, buffer, inset = 0, invert = FALSE)

## S4 method for signature 'numeric'
selectTopRight(n, buffer, inset = 0, invert = FALSE)

## S4 method for signature 'numeric'
selectBottomRight(n, buffer, inset = 0, invert = FALSE)

## S4 method for signature 'numeric'
selectBottomLeft(n, buffer, inset = 0, invert = FALSE)

## S4 method for signature 'numeric'
selectCorners(n, buffer, inset = 0, invert = FALSE)

## S4 method for signature 'numeric'
selectRows(rows, buffer, invert = FALSE)

## S4 method for signature 'numeric'
selectCols(cols, buffer, invert = FALSE)

## S4 method for signature 'numeric'
selectInner(n, buffer, invert = FALSE)

## S4 method for signature 'numeric'
selectOuter(n, buffer, invert = FALSE)

Arguments

x Integer vector of manhattan distances to select.

buffer Integer describing the number of pixels surrounding the central pixel.

invert Boolean indicating whether to invert the selection.

mhDist Integer vector of manhattan distances to select along with center pixel.

m matrix with 1's indicating selected positions and 0's indicated unselected positions.

rowInd Integer describing the row indices.
colInd  Integer describing the column indices.

n  Integer describing the number of outer pixels to select. Must be length of one.

inset  Integer describing the number of pixels to inset the selection from the outer edge of the matrix. Default of 0 uses no inset.

rows  Integer describing which rows to select.

cols  Integer describing which cols to select.

object  A MatrixSelection object.

Value

A text-based visualization of the select matrix indices.

Examples

res <- selectCenterPixel(0, 3)
show(res)
selectRadius(x=c(2,3,4), buffer=5, invert=FALSE)
selectCenterPixel(0, 5)
selectSubmatrix(m = matrix(rep(c(1,0,1), 3), nrow=3, ncol=3))
selectCoordinates(rowInd=1:3, colInd=1:3, buffer=5)
selectBlock(rowInd=1:3, colInd=1:3, buffer=5)
selectTopLeft(n=3, buffer=5, inset=1, invert=FALSE)
selectTopRight(n=3, buffer=5, inset=1, invert=FALSE)
selectBottomRight(n=3, buffer=5, inset=1, invert=FALSE)
selectBottomLeft(n=3, buffer=5, inset=1, invert=FALSE)
selectCorners(n=3, buffer=5, inset=1, invert=FALSE)
selectRows(rows=1:3, buffer=5, invert=FALSE)
selectCols(cols=1:3, buffer=5, invert=FALSE)
selectInner(n=1, buffer=5, invert=FALSE)
selectOuter(n=1, buffer=5, invert=FALSE)
seqnames1

Access each portion of a GInteractions-like object

Description

Access each portion of a GInteractions-like object

Usage

seqnames1(x, ...)
seqnames2(x, ...)
start1(x, ...)
end1(x, ...)
start2(x, ...)
end2(x, ...)

## S4 method for signature 'GInteractions.OR.InteractionSet'
seqnames1(x)

## S4 method for signature 'GInteractions.OR.InteractionSet'
seqnames2(x)

## S4 method for signature 'GInteractions.OR.InteractionSet'
start1(x)

## S4 method for signature 'GInteractions.OR.InteractionSet'
end1(x)

## S4 method for signature 'GInteractions.OR.InteractionSet'
start2(x)

## S4 method for signature 'GInteractions.OR.InteractionSet'
end2(x)

Arguments

x GInteractions object.
...

Value

A vector of values corresponding to the requested component of a GInteractions-like object. For seqnames1 and seqnames2 the RLE is coerced to a character vector.
Examples

library(InteractionSet)
## Create example reference interactions objects
gi <- read.table(text="
    chr1 10 20 chr1 50 60
    chr2 30 40 chr2 60 70
    chr1 50 60 chr3 10 20"
) |> as_ginteractions()

iset <- InteractionSet(assays=matrix(nrow=3),
                      interactions=gi)

## Access vectors of values
seqnames1(gi)
start1(gi)
end1(gi)
seqnames2(gi)
start2(gi)
end2(gi)

## Also works for InteractionSet-like objects
seqnames1(iset)
start1(iset)
end1(iset)
seqnames2(iset)
start2(iset)
end2(iset)

shiftRanges

Flexibly shifting GRanges according to strand

Description

Flexibly shifting GRanges according to strand

Usage

shiftRanges(x, pos)

## S4 method for signature 'GRanges,character_OR_numeric'
shiftRanges(x, pos)

Arguments

x GRanges object

pos Position within anchors to resize the bin. Can be a character or integer vector
      of length 1 or ‘length(x)’ designating the position for each element in bedpe.
Character options are "start", "end" and "center". Integers are referenced from the start position for '+' and '*' strands and from the end position for the '-' strand.

Value

GRanges object with a single position range that has been shifted appropriately.

Examples

library(GenomicRanges)

## Create example GRanges
gr1 <- GRanges(seqnames = "chr1",
               ranges = IRanges(start = rep(5000,3),
                                end = rep(6000,3)),
               strand = c('+', '-', '*'))

gr2 <- gr1 |> promoters(upstream = 2000, downstream = 200)

## Shifting anchors by keyword
shiftRanges(gr1, 'start')
shiftRanges(gr1, 'end')
shiftRanges(gr1, 'center')

## Shifting anchors by position
shiftRanges(gr1, 100)
shiftRanges(gr1, c(100, 200, 300))

## Shifting back to TSS
shiftRanges(gr2, 2000)

---

snapToBins

**Snap GRanges or GInteractions to nearest bins**

Description

Snap GRanges or GInteractions to nearest bins
Snap paired-objects to nearest bins

Usage

snapToBins(x, binSize)

## S4 method for signature 'GRanges,numeric'

snapToBins(x, binSize)

## S4 method for signature 'GInteractions,numeric'

snapToBins(x, binSize)
Arguments

x 'GInteractions' object.
binSize Integer (numeric) describing the new size of each range.

Value

GRanges object snapped to the nearest 'binSize'.
Input object snapped to the nearest 'binSize'.

Examples

library(GenomicRanges)
## Example GRanges object
x <- GRanges(seqnames = c("chr1"),
              ranges = IRanges(start = c(1, 1, 25, 19, 21),
                               end = c(15, 11, 31, 31, 39)))
snapToBins(x, binSize = 5)
snapToBins(x, binSize = 10)
snapToBins(x, binSize = 20)

library(InteractionSet)
## Sample GInteractions object
x <- GInteractions(anchor1 = c(GRanges("chr1:1-15"),
                                GRanges("chr1:1-11")),
                   anchor2 = c(GRanges("chr1:25-31"),
                                GRanges("chr1:19-31")))
snapToBins(x, binSize = 5)
snapToBins(x, binSize = 10)
snapToBins(x, binSize = 20)

snpToBins(x, binSize = 5)
snapToBins(x, binSize = 10)
snapToBins(x, binSize = 20)

sources

Accessor for sources

Description

Access the names or source files of a 'MergedGInteractions' object.

Usage

sources(x)

## S4 method for signature 'MergedGInteractions'
sources(x)
subsetBySource

Arguments

x         MergedGInteractions object.

Value

A character vector of names or source files of a `MergedGInteractions` object.

Examples

```r
## Load required packages
library(data.table, include.only="fread")

## Load marinerData
if (!require("marinerData", quietly = TRUE))
  BiocManager::install("marinerData")

## Reference BEDPE files (loops called with SIP)
loopFiles <- c(
  marinerData::FS_5kbLoops.txt(),
  marinerData::WT_5kbLoops.txt()
)
names(loopFiles) <- c("FS", "WT")

## Read in loopFiles as a list of GIInteractions
## Use only first 1000 rows for fast example
giList <-
  lapply(loopFiles, fread, nrow=1000) |>
  lapply(as_ginteractions)

## Cluster & merge pairs
x <- mergePairs(x = giList,
                 radius = 10e03)

sources(x)
```

Description

Returns the subset of MergedGInteractions that belong to each input source object (see these with `sources(x)`). If the source pairs all come from the same object, their corresponding merged pair is returned. However, if at least one source pair comes from a different object, then that merged pair is not returned.
Usage

subsetBySource(x, include, exclude)

## S4 method for signature 'MergedGInteractions,missing,missing'
subsetBySource(x)

## S4 method for signature 'MergedGInteractions,character_OR_missing,missing'
subsetBySource(x, include)

## S4 method for signature 'MergedGInteractions,missing,character_OR_missing'
subsetBySource(x, exclude)

## S4 method for signature
## 'MergedGInteractions,character_OR_missing,character_OR_missing'
subsetBySource(x, include, exclude)

Arguments

x          MergedGInteractions object.
include (Optional) A character vector of sources in which a pair must be present. For a list of available sources use ‘sources(x)’.
exclude (Optional) A character vector of sources in which a pair must be absent. For a list of available sources use ‘sources(x)’.

Details

Optional ‘include’ and ‘exclude’ parameters modulate the behavior of ‘subsetBySource’ to return different subsets of originating pairs. For example, ‘include’ requires that the returned pairs be present in specific sources, while ‘exclude’ requires that returned pairs be absent from specific sources. Sources not listed in either ‘include’ or ‘exclude’ are ignored (they may or may not) be present in the returned ‘MergedGInteractions’ object. ‘include’ and ‘exclude’ can be used independently or in combination to return every possible set. If any of the same sources are used in both ‘include’ and ‘exclude’ the function will return a 0-length MergedGInteractions object.

Value

A list of sub-setted ‘MergedGInteractions’ objects or a ‘MergedGInteractions’ object (if ‘include’ and/or ‘exclude’ are used).

Examples

## Load required packages
library(GenomicRanges)
library(InteractionSet)

## Define example anchor regions
gr1 <-
   GRanges(seqnames = "chr1",
           ranges = IRanges(start = c(30,40,40,70,80),
           )
subsetBySource

end = c(40,50,50,80,90))

gr2 <-
  GRanges(seqnames = "chr1",
          ranges = IRanges(start = c(30,30,50,10,30),
                          end = c(40,40,60,20,40)))

## Form GInteractions and split into two files
gilist <- split(x = GInteractions(gr1, gr2),
                 f = c(rep(1,3), rep(2,2)))

## Merge pairs
x <- mergePairs(x = gilist, radius = 20)

subsetBySource(x)
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