

Package ‘HiCExperiment’

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Title Bioconductor class for interacting with Hi-C files in R

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Description R generic interface to Hi-C contact matrices in `.(m)cool` , ` .hic` or HiC-Pro derived formats, as well as other Hi-C processed file formats. Contact matrices can be partially parsed using a random access method, allowing a memory-efficient representation of Hi-C data in R. The `HiCExperiment` class stores the Hi-C contacts parsed from local contact matrix files. `HiCExperiment` instances can be further investigated in R using the `HiContacts` analysis package.

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URL <https://github.com/js2264/HiCExperiment>

BugReports <https://github.com/js2264/HiCExperiment/issues>

Depends R (>= 4.2)

Imports InteractionSet, strawr, GenomeInfoDb, GenomicRanges, IRanges, S4Vectors, BiocGenerics, BiocIO, BiocParallel, methods, rhdf5, Matrix, vroom, dplyr, stats

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 'AllGenerics.R' 'HiCExperiment-methods.R'
 'AggrHiCExperiment-methods.R' 'PairsFile-class.R'
 'ContactsFile-class.R' 'ContactsFile-methods.R'
 'CoolFile-class.R' 'CoolFile-methods.R' 'HicFile-class.R'
 'HicFile-methods.R' 'HicproFile-class.R' 'HicproFile-methods.R'
 'PairsFile-methods.R' 'import-methods.R' 'available.R' 'bin.R'
 'checks.R' 'coerce.R' 'data.R' 'export-methods.R' 'globals.R'

'parse-cool.R' 'parse-hic.R' 'parse-hicpro.R' 'parse-pairs.R'
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Author Jacques Serizay [aut, cre] (<<https://orcid.org/0000-0002-4295-0624>>)

Maintainer Jacques Serizay <jacquessserizay@gmail.com>

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Description

Pairs parsing functions

Usage

```
.pairs2gi(
  file,
  chr1.field = NULL,
  start1.field = NULL,
  chr2.field = NULL,
  start2.field = NULL,
  strand1.field = NULL,
  strand2.field = NULL,
  frag1.field = NULL,
  frag2.field = NULL,
  nThread = 1,
  nrows = Inf
)
```

Arguments

| | |
|--|---|
| file | pairs file. Default formatting is <readname>\t<chr1>\t<start1>\t<chr2>\t<start2>. |
| chr1.field, start1.field, chr2.field, start2.field, strand1.field, | |
| strand2.field, frag1.field, frag2.field | |
| nThread | Index of the column in which each field is contained in the pairs file. |
| nrows | Number of CPUs to use to import the pairs file in R |
| | Number of pairs to import |

Value

a GInteractions object

AggrHiCExperiment AggrHiCExperiment *S4 class*

Description

The AggrHiCExperiment extends HiCExperiment class.

Usage

```
AggrHiCExperiment(
  file,
  resolution = NULL,
  targets,
  flankingBins = 50,
  metadata = list(),
  topologicalFeatures = S4Vectors::SimpleList(),
  pairsFile = NULL,
  bed = NULL,
  maxDistance = NULL,
  BPPARAM = BiocParallel::bpparam()
)
```

```

## S4 method for signature 'AggrHiCExperiment,missing'
slices(x)

## S4 method for signature 'AggrHiCExperiment,character'
slices(x, name)

## S4 method for signature 'AggrHiCExperiment,numeric'
slices(x, name)

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

```

Arguments

| | |
|----------------------------------|---|
| <code>file</code> | CoolFile or plain path to a Hi-C contact file |
| <code>resolution</code> | Resolution to use with the Hi-C contact file |
| <code>targets</code> | Set of chromosome coordinates for which interaction counts are extracted from the Hi-C contact file, provided as a GRanges object (for diagonal-centered loci) or as a GInteractions object (for off-diagonal coordinates). |
| <code>flankingBins</code> | Number of bins on each flank of the bins containing input targets. |
| <code>metadata</code> | list of metadata |
| <code>topologicalFeatures</code> | topologicalFeatures provided as a named SimpleList |
| <code>pairsFile</code> | Path to an associated .pairs file |
| <code>bed</code> | Path to regions file generated by HiC-Pro |
| <code>maxDistance</code> | Maximum distance to use when compiling distance decay |
| <code>BPPARAM</code> | BiocParallel parameters |
| <code>x, object</code> | A AggrHiCExperiment object. |
| <code>name</code> | The name/index of slices to extract. |

Value

An AggrHiCExperiment object.

Slots

| | |
|----------------------------------|--|
| <code>fileName</code> | Path of Hi-C contact file |
| <code>resolutions</code> | Resolutions available in the Hi-C contact file. |
| <code>resolution</code> | Current resolution |
| <code>interactions</code> | Genomic Interactions extracted from the Hi-C contact file |
| <code>scores</code> | Available interaction scores. |
| <code>slices</code> | Available interaction slices. |
| <code>topologicalFeatures</code> | Topological features associated with the dataset (e.g. loops (<Pairs>), borders (<GRanges>), viewpoints (<GRanges>), etc...) |
| <code>pairsFile</code> | Path to the .pairs file associated with the Hi-C contact file |
| <code>metadata</code> | metadata associated with the Hi-C contact file. |

See Also

[HiCExperiment\(\)](#)

Examples

```
fpath <- HiContactsData::HiContactsData('yeast_wt', 'mcool')
data(centros_yeast)
x <- AggrHiCExperiment(
  file = fpath,
  resolution = 8000,
  targets = centros_yeast[c(4, 7)]
)
x
slices(x, 'count')[1:10, 1:10, 1]
```

AllGenerics

*Generic functions***Description**

Generics functions created in HiCExperiment package.

Arguments

| | |
|-------|--------------------------------|
| x | Passed to corresponding method |
| name | Passed to corresponding method |
| value | Passed to corresponding method |
| ... | Passed to corresponding method |

as

*Coercing functions***Description**

Coercing functions available for HiCExperiment objects.

Usage

```
## S4 method for signature 'HiCExperiment'
as.matrix(x, use.scores = "balanced", sparse = FALSE)

## S4 method for signature 'HiCExperiment'
as.data.frame(x)

gi2cm(gi, use.scores = "score")

cm2matrix(cm, replace_NA = NA, sparse = FALSE)

df2gi(
```

```

df,
seqnames1 = "seqnames1",
start1 = "start1",
end1 = "end1",
seqnames2 = "seqnames2",
start2 = "start2",
end2 = "end2"
)

```

Arguments

| | |
|--|--|
| x | HiCExperiment object |
| use.scores | Which scores to use to inflate GInteractions |
| sparse | Whether to return the contact matrix as a sparse matrix |
| gi | GInteractions object |
| cm | A ContactMatrix object |
| replace_NA | Replace NA values |
| df | A data.frame object |
| seqnames1, start1, end1, seqnames2, start2, end2 | Names (as strings) of columns containing corresponding information in a data.frame parsed into GInteractions (default: FALSE) |

Examples

```

mcoolPath <- HiContactsData::HiContactsData('yeast_wt', 'mcool')
contacts <- import(mcoolPath, focus = 'XVI', resolution = 16000, format = 'cool')
gis <- interactions(contacts)
cm <- gi2cm(gis, 'balanced')
cm
cm2matrix(cm)[1:10, 1:10]
df2gi(data.frame(
  chr1 = 'I', start1 = 10, end1 = 100,
  chr2 = 'I', start2 = 40, end2 = 1000,
  score = 12,
  weight = 0.234,
  filtered = TRUE
), seqnames1 = 'chr1', seqnames2 = 'chr2')

```

Description

HiCExperiment binning methods

Usage

```

## S4 method for signature 'GInteractions,numeric'
bin(x, resolution, seqinfo = NULL)

## S4 method for signature 'PairsFile,numeric'
bin(x, resolution, seqinfo = NULL)

```

Arguments

| | |
|------------|---|
| x | A PairsFile or GInteractions object |
| resolution | Which resolution to use to bin the interactions |
| seqinfo | Seqinfo object |

Examples

```
pairsf <- HiContactsData::HiContactsData('yeast_wt', 'pairs.gz')
pf <- PairsFile(pairsf)
```

checks*Checks functions*

Description

Internal functions to validate the nature/structure of (m)cool files or HiCEExperiment objects. All these check functions should return a logical.

Usage

```
.check_cool_file(path)

.check_cool_format(path, resolution, ...)

.is_mcool(path)

.is_cool(path)

.check_hic_file(path)

.check_hic_format(path, resolution, ...)

.is_hic(path)

.check_hicpro_files(path, bed)

.is_hicpro_matrix(path)

.is_hicpro_regions(bed)

.check_resolution(contacts, resolution)

.check_scores(contacts, use.scores)

.is_square(pair)
```

Arguments

| | |
|------------|--|
| path | Path of a (m)cool file |
| resolution | Resolution |
| ... | HiCExperiment object, arguments passed on by other functions |
| bed | Path to regions file generated by HiC-Pro |
| contacts | A HiCExperiment object |
| use.scores | Name of scores to check |
| pair | Pairs object with length of 1 |

Value

Logical

ContactsFile-class *ContactsFile S4 class*

Description

The **ContactsFile** class describes a **BiocFile** object, pointing to the location of an Hi-C matrix file (cool, mcool, hic, hicpro, ...) and containing additional slots:

1. resolution: at which resolution the associated mcool file should be parsed
2. pairsFile: the path (in plain character) to an optional pairs file (stored as a **PairsFile** object);
3. metadata: a list. If the CoolFile is created by **HiCool**, it will contain two elements: log (path to **HiCool** processing log file) and stats (aggregating some stats from **HiCool** mapping).

ContactsFile methods.

Arguments

| | |
|------------|--|
| path | String; path to an Hi-C matrix file (cool, mcool, hic, hicpro) |
| resolution | numeric; resolution to use with Hi-C matrix file |
| pairsFile | String; path to a pairs file |
| metadata | list. |
| object | A ContactsFile object. |
| x | A ContactsFile object. |

Slots

resolution numeric value or NULL
 pairsFile PairsFile object
 metadata list

See Also

[CoolFile\(\)](#), [HicFile\(\)](#), [HicproFile\(\)](#)

Description

The CoolFile class describes a BiocFile object, pointing to the location of an Hi-C matrix file (cool, mcool, hic, hicpro, ...) and containing additional slots:

1. resolution: at which resolution the associated mcool file should be parsed
2. pairsFile: the path (in plain character) to an optional pairs file (stored as a PairsFile object);
3. metadata: a list. If the CoolFile is created by HiCool, it will contain two elements: log (path to HiCool processing log file) and stats (aggregating some stats from HiCool mapping).

CoolFile methods.

Arguments

| | |
|------------|--|
| path | String; path to a (m)cool file |
| resolution | numeric; resolution to use with mcool file |
| pairsFile | String; path to a pairs file |
| metadata | list; if the CoolFile object was generated by HiCool::HiCool, this list contains the path to log file, some statistics regarding the number of pairs obtained by hicstuff as well as the arguments and the hash ID used by HiCool. |
| object | A CoolFile object. |

See Also

[HicFile\(\)](#), [HicproFile\(\)](#)

Examples

```
mcoolPath <- HiContactsData::HiContactsData('yeast_wt', 'mcool')
pairsPath <- HiContactsData::HiContactsData('yeast_wt', 'pairs.gz')
cf <- CoolFile(
  mcoolPath,
  resolution = 2000,
  pairsFile = pairsPath,
  metadata = list(info = 'Yeast WT Hi-C exp.')
)
cf
resolution(cf)
pairsFile(cf)
metadata(cf)
```

| | |
|------|--|
| data | <i>Example datasets provided in HiCExperiment & HiContactsData</i> |
|------|--|

Description

Example datasets provided in HiCExperiment & HiContactsData

Usage

```
data(centros_yeast)

contacts_yeast(full = FALSE)

contacts_yeast_eco1(full = FALSE)
```

Arguments

| | |
|------|------------------------------------|
| full | Whether to import all interactions |
|------|------------------------------------|

Format

An object of class "GRanges".

Source

HiContacts

Examples

```
data(centros_yeast)
centros_yeast
contacts_yeast()
```

| | |
|----------------|-------------------------------------|
| export-methods | <i>HiCExperiment export methods</i> |
|----------------|-------------------------------------|

Description

Export methods to save a HiCExperiment object into a set of HiC-Pro-style files (matrix & regions files)

Usage

```
## S4 method for signature 'HiCExperiment,missing,character'
export(object, prefix, format, ...)
```

Arguments

| | |
|--------|---|
| object | A HiCExperiment object |
| prefix | Prefix used when generating output file(s). |
| format | File format. Available: cool and HiC-Pro. |
| ... | Extra arguments to use when exporting to cool. Can be metadata <string> or chunksize <integer>. |

Value

Path to saved files

Examples

```
#####
## ----- Importing .(m)cool contact matrices ----- ##
#####

mcoolPath <- HiContactsData::HiContactsData('yeast_wt', 'mcool')
hic <- import(mcoolPath, format = 'mcool', resolution = 16000)
export(hic[["II"]], prefix = 'subset_chrII', format = 'cool')
export(hic[["II"]], prefix = 'subset_chrII', format = 'HiC-Pro')
```

HiCExperiment

HiCExperiment *S4 class*

Description

The HiCExperiment class describes Hi-C contact files imported in R, either through the HiCExperiment constructor function or using the import method implemented by HiCExperiment package.

Usage

```
HiCExperiment(
  file,
  resolution = NULL,
  focus = NULL,
  metadata = list(),
  topologicalFeatures = S4Vectors::SimpleList(compartments = GenomicRanges::GRanges(),
    borders = GenomicRanges::GRanges(), loops =
    InteractionSet::GInteractions(GenomicRanges::GRanges(), GenomicRanges::GRanges()),
    viewpoints = GenomicRanges::GRanges()),
  pairsFile = NULL,
  bed = NULL
)

makeHiCExperimentFromGInteractions(gi)

## S4 method for signature 'HiCExperiment'
resolutions(x)

## S4 method for signature 'HiCExperiment'
```

```

resolution(x)

## S4 method for signature 'HiCExperiment'
focus(x)

## S4 replacement method for signature 'HiCExperiment,character'
focus(x) <- value

## S4 method for signature 'HiCExperiment,numERIC'
zoom(x, resolution)

## S4 method for signature 'HiCExperiment,character'
refocus(x, focus)

## S4 method for signature 'HiCExperiment,missing'
scores(x)

## S4 method for signature 'HiCExperiment,character'
scores(x, name)

## S4 method for signature 'HiCExperiment,numERIC'
scores(x, name)

## S4 replacement method for signature 'HiCExperiment,character,numERIC'
scores(x, name) <- value

## S4 method for signature 'HiCExperiment,missing'
topologicalFeatures(x)

## S4 method for signature 'HiCExperiment,character'
topologicalFeatures(x, name)

## S4 method for signature 'HiCExperiment,numERIC'
topologicalFeatures(x, name)

## S4 replacement method for signature 'HiCExperiment,character,GRangesOrGInteractions'
topologicalFeatures(x, name) <- value

## S4 method for signature 'HiCExperiment'
pairsFile(x)

## S4 replacement method for signature 'HiCExperiment,character'
pairsFile(x) <- value

## S4 replacement method for signature 'HiCExperiment,list'
metadata(x) <- value

## S4 method for signature 'HiCExperiment,numERIC'
subsetByOverlaps(x, ranges)

## S4 method for signature 'HiCExperiment,logical'
subsetByOverlaps(x, ranges)

```

```
## S4 method for signature 'HiCExperiment,GRanges'
subsetByOverlaps(x, ranges, type = c("within", "any"))

## S4 method for signature 'HiCExperiment,GInteractions'
subsetByOverlaps(x, ranges)

## S4 method for signature 'HiCExperiment,Pairs'
subsetByOverlaps(x, ranges)

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

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

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

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

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

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

## S4 method for signature 'HiCExperiment'
fileName(object)

## S4 method for signature 'HiCExperiment'
interactions(x, fillout.regions = FALSE)

## S4 replacement method for signature 'HiCExperiment,GInteractions'
interactions(x) <- value

## S4 method for signature 'HiCExperiment'
length(x)

## S4 replacement method for signature 'HiCExperiment'
x$name <- value

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

## S4 method for signature 'HiCExperiment'
seqinfo(x)

## S4 method for signature 'HiCExperiment'
bins(x)
```

```
## S4 method for signature 'HiCExperiment'
anchors(x)

## S4 method for signature 'HiCExperiment'
regions(x)

## S4 method for signature 'HiCExperiment'
cis(x)

## S4 method for signature 'HiCExperiment'
trans(x)
```

Arguments

| | |
|---------------------|---|
| file | CoolFile or plain path to a Hi-C contact file |
| resolution | Resolution to use with the Hi-C contact file |
| focus | Chromosome coordinates for which interaction counts are extracted from the Hi-C contact file, provided as a character string (e.g. "II:4001-5000"). If not provided, the entire Hi-C contact file will be imported. |
| metadata | list of metadata |
| topologicalFeatures | topologicalFeatures provided as a named SimpleList |
| pairsFile | Path to an associated .pairs file (optional) |
| bed | Path to regions file generated by HiC-Pro (optional) |
| gi | GInteractions object |
| x | A HiCExperiment object. |
| value | Value to add to topologicalFeatures, scores, pairsFile or metadata slots. |
| name | Name of the element to access in topologicalFeatures or scores SimpleLists. |
| type | any of within or any, to subset interactions by overlap with a provided GRanges. |
| i, ranges | a GRanges, coordinates in character, or boolean vector to subset a HiCExperiment |
| object | A HiCExperiment object. |
| fillout.regions | Whehter to add missing regions to GInteractions' regions? |

Value

An *HiCExperiment* object.

Slots

fileName Path of Hi-C contact file
focus Chr. coordinates for which interaction counts are extracted from the Hi-C contact file.
resolutions Resolutions available in the Hi-C contact file.
resolution Current resolution
interactions Genomic Interactions extracted from the Hi-C contact file
scores Available interaction scores.

`topologicalFeatures` Topological features associated with the dataset (e.g. loops (`\<GInteractions\>`), borders (`\<GRanges\>`), viewpoints (`\<GRanges\>`), etc...)

`pairsFile` Path to the .pairs file associated with the Hi-C contact file

`metadata` metadata associated with the Hi-C contact file.

See Also

[AggrHiCExperiment\(\)](#), [CoolFile\(\)](#), [HicFile\(\)](#), [HicproFile\(\)](#), [PairsFile\(\)](#)

Examples

```
#####
## Create a HiCExperiment object from a disk-stored contact matrix ##
#####

mcool_file <- HiContactsData::HiContactsData("yeast_wt", "mcool")
pairs_file <- HiContactsData::HiContactsData("yeast_wt", "pairs.gz")
contacts <- HiCExperiment(
  file = mcool_file,
  resolution = 8000L,
  pairsFile = pairs_file
)
contacts

#####
## ----- Manually create a HiCExperiment from GInteractions ----- ##
#####

gis <- interactions(contacts)[1:1000]
contacts2 <- makeHiCExperimentFromGInteractions(gis)
contacts2

#####
## ----- Slots present in an HiCExperiment object ----- ##
#####

fileName(contacts)
focus(contacts)
resolutions(contacts)
resolution(contacts)
interactions(contacts)
scores(contacts)
topologicalFeatures(contacts)
pairsFile(contacts)

#####
## ----- Slot getters ----- ##
#####

scores(contacts, 1) |> head()
scores(contacts, 'balanced') |> head()
topologicalFeatures(contacts, 1)

#####
## ----- Slot setters ----- ##
#####
```

```

scores(contacts, 'random') <- runif(length(contacts))
topologicalFeatures(contacts, 'loops') <- InteractionSet::GInteractions(
  GenomicRanges::GRanges('II:15324'),
  GenomicRanges::GRanges('II:24310')
)
pairsFile(contacts) <- HiContactsData('yeast_wt', 'pairs.gz')

#####
## ----- Subsetting functions -----
#####

contacts[1:100]
contacts['II']
contacts[c('II', 'III')]
contacts['II|III']
contacts['II:10001-30000|III:50001-90000']

#####
## ----- Utils functions -----
#####

## Adapted from other packages

seqinfo(contacts)
bins(contacts)
anchors(contacts)
regions(contacts)

#####
## ----- Coercing HiCExperiment objects -----
#####

as(contacts, 'GInteractions')
as(contacts, 'ContactMatrix')
as(contacts, 'matrix')[seq_len(10), seq_len(10)]
as(contacts, 'data.frame')[seq_len(10), seq_len(10)]

```

HiCExperiment utils *Utils functions*

Description

Utilities to facilitate parsing/handling of coordinates, GInteractions, Pairs, ... These functions are not exported.

Usage

```

splitCoords(coords)

coords2char(coords, big.mark = ",,")

char2coords(char)

sortPairs(pairs)

```

```

asGInteractions(df)

sdiag(A, k = 0)

.df2symmmat(diag, score)

distanceDecay(dump, threshold = NULL)

detrendingModel(file, resolution)

.fixRegions(gis, bins, coords)

```

Arguments

| | |
|------------|--|
| coords | A set of genomic coordinates (either as a GRanges object or as a character string) |
| big.mark | Separator for thousands when printing out genomic coordinates as character |
| char | char (e.g. "II:30001-50000" or "II:30001-50000 II:60001-80000") |
| pairs | Pairs object |
| df | a data.frame to turn into a GInteraction object. |
| A | Numerical matrix |
| k | secondary diagonal k |
| diag | vector of distances to diagonal |
| score | scores to parse into symmetrical matrix |
| dump | dumped contacts as GInteractions, e.g. from .dumpCool |
| threshold | maximum distance to compute distance decay for |
| file | path to a HiC contact matrix file |
| resolution | Resolution to use with the HiC contact matrix file |
| gis | GInteractions object |
| bins | Larger set of regions (usually bins from HiCEExperiment) |

Value

Reformatted coordinates or GInteractions.

HicFile-class

HicFile *S4 class*

Description

The HicFile class describes a BiocFile object, pointing to the location of a .hic file (usually created with juicer) and containing 3 additional slots:

1. resolution: at which resolution the associated .hic file should be parsed;
2. pairsFile: the path (in plain character) to an optional pairs file (stored as a PairsFile object);
3. metadata: a list metadata

HicFile methods.

Arguments

| | |
|------------|--|
| path | String; path to a .hic file |
| resolution | numeric; resolution to use with mcool file |
| pairsFile | String; path to a pairs file |
| metadata | list. |
| object | A <code>HicFile</code> object. |

See Also

[CoolFile\(\)](#), [HicproFile\(\)](#)

Examples

```
hicPath <- HiContactsData::HiContactsData('yeast_wt', 'hic')
pairsPath <- HiContactsData::HiContactsData('yeast_wt', 'pairs.gz')
hic <- HicFile(
  hicPath,
  resolution = 16000,
  pairsFile = pairsPath,
  metadata = list(type = 'example')
)
hic
resolution(hic)
pairsFile(hic)
metadata(hic)
```

Description

The `HicproFile` class describes a `BiocFile` object, pointing to the location of a HiC-Pro-generated matrix file and containing 4 additional slots:

1. bed: path to the matching .bed file generated by HiC-Pro;
2. resolution: at which resolution the associated mcool file should be parsed ;
3. pairsFile: the path (in plain character) to an optional pairs file (stored as a `PairsFile` object);
4. metadata: a list metadata

`HicproFile` methods.

Arguments

| | |
|-----------|---|
| path | String; path to the HiC-Pro output .matrix file (matrix file) |
| bed | String; path to the HiC-Pro output .bed file (regions file) |
| pairsFile | String; path to a pairs file |
| metadata | list. |
| object | A <code>HicproFile</code> object. |

Slots

`bed` Path to the matching .bed file generated by HiC-Pro

See Also

[CoolFile\(\)](#), [HicFile\(\)](#)

Examples

```
hicproMatrixPath <- HiContactsData::HiContactsData('yeast_wt', 'hicpro_matrix')
hicproBedPath <- HiContactsData::HiContactsData('yeast_wt', 'hicpro_bed')
pairsPath <- HiContactsData::HiContactsData('yeast_wt', 'pairs.gz')
hicpro <- HicproFile(
  hicproMatrixPath, bed = hicproBedPath, pairs = pairsPath ,
  metadata = list(type = 'example')
)
hicpro
resolution(hicpro)
pairsFile(hicpro)
metadata(hicpro)
```

Description

Import methods to parse Hi-C files (.(m)cool, .hic, HiC-Pro derived matrices, pairs files) into data structures implemented in the HiCExperiment package.

Usage

```
import(con, format, text, ...)

## S4 method for signature 'ANY'
availableResolutions(x, ...)

## S4 method for signature 'CoolFile'
availableResolutions(x)

## S4 method for signature 'HicFile'
availableResolutions(x)

## S4 method for signature 'HicproFile'
availableResolutions(x)

## S4 method for signature 'ANY'
availableChromosomes(x, ...)

## S4 method for signature 'CoolFile'
availableChromosomes(x)

## S4 method for signature 'HicFile'
```

```
availableChromosomes(x)

## S4 method for signature 'HicproFile'
availableChromosomes(x)
```

Arguments

| | |
|--------|---|
| ... | Extra parameters to pass to format-specific methods. A list of possible arguments is provided in the next section. |
| con, x | Path or connection to a cool, mcool, .hic or HiC-Pro derived files. Can also be a path to a pairs file. |
| format | The format of the output. If missing and 'con' is a filename, the format is derived from the file extension. This argument is unnecessary when files are directly provided as CoolFile, HicFile, HicproFile or PairsFile. |
| text | If 'con' is missing, this can be a character vector directly providing the string data to import. |

Value

A HiCExperiment or GInteractions object

import arguments for ContactFile class

ContactFile class gathers CoolFile, HicFile and HicproFile classes. When importing a ContactFile object in R, two main arguments can be provided besides the ContactFile itself:

- **resolution**: Resolutions available in the disk-stored contact matrix can be listed using `availableResolutions()`
- **focus**: A genomic locus (or pair of loci) provided as a string. It can be any of the following string structures:
 - "II" or "II:20001-30000": this will extract a symmetrical square HiCExperiment object, of an entire chromosome or an portion of it.
 - "II|III" or "II:20001-30000|III:40001-90000": this will extract a non-symmetrical HiCExperiment object, with an entire or portion of different chromosomes on each axis.

Examples

```
#####
## ----- Importing .(m)cool contact matrices ----- ##
#####

mcoolPath <- HiContactsData::HiContactsData('yeast_wt', 'mcool')
availableResolutions(mcoolPath)
availableChromosomes(mcoolPath)
import(mcoolPath, resolution = 16000, focus = 'XVI', format = 'cool')

#####
## ----- Importing .hic contact matrices ----- ##
#####

hicPath <- HiContactsData::HiContactsData('yeast_wt', 'hic')
availableResolutions(hicPath)
availableChromosomes(hicPath)
import(hicPath, resolution = 16000, focus = 'XVI', format = 'hic')
```

```
#####
## ----- Importing HiC-Pro derived contact matrices ----- ##
#####

hicproMatrixPath <- HiContactsData::HiContactsData('yeast_wt', 'hicpro_matrix')
hicproBedPath <- HiContactsData::HiContactsData('yeast_wt', 'hicpro_bed')
availableResolutions(hicproMatrixPath, hicproBedPath)
availableChromosomes(hicproMatrixPath, hicproBedPath)
import(hicproMatrixPath, bed = hicproBedPath, format = 'hicpro')
```

multi2Query*Querying multiple slices of a contact matrix*

Description

These functions are the workhorse internal functions used to extract counts from multiple genomic coordinates in a Hi-C contact matrix.

Usage

```
.multi2DQuery(
  file,
  resolution,
  pairs,
  maxDistance = NULL,
  bed = NULL,
  BPPARAM = BiocParallel::bpparam()
)
```

Arguments

| | |
|-------------|--|
| file | path to a Hi-C contact file (can be any format, (m)cool, .hic, or HiC-Pro-derived) |
| resolution | resolution to use to import matrix over specified targets |
| pairs | slices to read, provided as a Pairs object |
| maxDistance | Maximum distance to use when compiling distance decay |
| bed | associated bed file for HiC-Pro derived contact matrix. |
| BPPARAM | BiocParallel parameters |

Value

a GInteractions object with count, balanced, detrended and expected scores

PairsFile-class PairsFile *S4 class*

Description

The PairsFile class describes a BiocFile object, pointing to the location of pairs file, typically generated by HiCool::HiCool().

PairsFile methods

Arguments

x Path to a pairs file

See Also

[CoolFile\(\)](#), [HicFile\(\)](#), [HicproFile\(\)](#)

Examples

```
pairsPath <- HiContactsData::HiContactsData('yeast_wt', 'pairs.gz')
pf <- PairsFile(pairsPath)
pf
pairsFile(pf)
```

parse-cool *Parsing (m)cool files*

Description

These functions are the workhorse internal functions used to import a .(m)cool file as GInteractions (wrapped into a HiCExperiment object by HiCExperiment() function).

Usage

```
.getCoolAnchors(file, resolution = NULL, balanced = "cooler")

.getCountsFromPair(file, pair, anchors, resolution = NULL)

.getCounts(file, coords, anchors, resolution = NULL)

.fetchCool(file, path, resolution = NULL, idx = NULL, ...)

.dumpCool(file, resolution = NULL)

.lsCoolFiles(file, verbose = FALSE)

.lsCoolResolutions(file, verbose = FALSE)

.cool2seqinfo(file, resolution = NULL)

.cool2gi(file, coords = NULL, resolution = NULL)
```

Arguments

| | |
|------------|---|
| file | path to a Hi-C contact file (in (m)cool format) |
| resolution | resolution of the contact matrix |
| balanced | import balancing scores |
| pair | Genomic coordinates to extract contacts for, stored as a Pairs of GRanges (e.g. S4Vectors::Pairs(GRanges("II:200000-300000"), GRanges("II:70000-100000"))). |
| anchors | anchors from .getCoolAnchors() |
| coords | Genomic coordinates to extract contacts for, stored as a GRanges object |
| path | Internal path of the cool file to check |
| idx | Index to extract from the cool (HDF5) file |
| ... | Other arguments passed to .fetchCool |
| verbose | Print resolutions in the console |

Value

Silently, a numerical vector of resolutions stored in the cool file

parse-hic

*Parsing hic files***Description**

These functions are the workhorse internal functions used to import a .hic file as GInteractions (wrapped into a HiCExperiment object by HiCExperiment() function).

Usage

```
.hic2gi(file, coords = NULL, resolution = NULL)

.lsHicResolutions(file, verbose = FALSE)

.getHicAnchors(file, resolution = NULL)

.hic2seqinfo(file)

.dumpHic(file, resolution = NULL)
```

Arguments

| | |
|------------|--|
| file | path to a Hi-C contact file in .hic format |
| coords | NULL, character, or GRanges. Can also be a Pairs object of paired GRanges (length of 1). |
| resolution | resolution of the contact matrix to use |
| verbose | Print resolutions in the console |

Value

a GInteractions object

vector

| | |
|--------------|--|
| parse-hicpro | <i>Parsing hicpro files (matrix & bed)</i> |
|--------------|--|

Description

These functions are the workhorse internal functions used to import HiC-Pro .matrix and .bed files as GInteractions (wrapped into a HiCEExperiment object by HiCEExperiment() function).

Usage

```
.hicpro2gi(file, bed)  
.getHicproAnchors(bed)  
.hicpro2seqinfo(bed)  
.dumpHicpro(file, bed)
```

Arguments

| | |
|------|---|
| file | path to a matrix file from HiC-Pro |
| bed | path to the regions file generated by HiC-Pro |

Value

a GInteractions object

| | |
|-----------|---|
| reexports | <i>Objects exported from other packages</i> |
|-----------|---|

Description

These objects are imported from other packages. Follow the links below to see their documentation.

S4Vectors [metadata](#)

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