

# Package ‘InteractionSet’

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**Title** Base Classes for Storing Genomic Interaction Data

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**Description** Provides the GInteractions, InteractionSet and ContactMatrix objects and associated methods for storing and manipulating genomic interaction data from Hi-C and ChIA-PET experiments.

**License** GPL-3

**NeedsCompilation** yes

**VignetteBuilder** knitr

**Collate** ContactMatrix-class.R GInteractions-class.R  
InteractionSet-class.R getset.R matching.R swapAnchors.R  
GRanges-methods.R pairs.R conversion.R linearize.R  
boundingBox.R distances.R overlaps.R linkOverlaps.R

## R topics documented:

boundingBox . . . . .	2
ContactMatrix accessors . . . . .	3
ContactMatrix class . . . . .	6
ContactMatrix distances . . . . .	8
ContactMatrix overlaps . . . . .	10
ContactMatrix sorting . . . . .	11
ContactMatrix subsetting . . . . .	13
Convert classes . . . . .	15
GInteractions class . . . . .	18
GRanges methods . . . . .	20
Interaction accessors . . . . .	21

Interaction compare . . . . .	26
Interaction distances . . . . .	29
Interaction overlaps . . . . .	31
Interaction subsetting . . . . .	34
InteractionSet class . . . . .	37
Linearize interactions . . . . .	39
linkOverlaps . . . . .	40
pairs . . . . .	42

<b>Index</b>	<b>44</b>
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boundingBox	<i>Get the bounding box</i>
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---

## Description

Computing a minimum bounding box for groups of pairwise interactions.

## Usage

```
## S4 method for signature 'GInteractions'
boundingBox(x, f)
```

```
# Note, the same call is used for the InteractionSet method.
```

## Arguments

x	A GInteractions or InteractionSet object.
f	A factor or vector of length equal to that of x, indicating the group to which each pairwise interaction belongs.

## Details

For any group of pairwise interactions, the minimum bounding box is the smallest rectangle in the interaction space that contains all interactions in the group. Each side of the box has coordinates spanning the most extreme anchor regions on the corresponding chromosome. This is often useful for summarizing clusters of interactions.

Grouping of interactions is specified using `f`, where interactions in `x` with the same level of `f` are considered to be in the same group. If `f` is not specified, all interactions in `x` are assumed to be in a single group (named as “1”). An error will be raised if a group spans multiple chromosomes for either the first or second anchor regions.

The function returns a GInteractions object containing the coordinates of the bounding boxes for all groups. Each interaction represents a bounding box for a group, where the anchor regions represent the sides of the box. Entries are named according to the levels of `f`, in order to specify which bounding box corresponds to which group.

It is recommended to run [swapAnchors](#) prior to computing the bounding box for intra-chromosomal groups. If all `anchor1 >= anchor2` or all `anchor1 <= anchor2`, all interactions will lie on one side of the diagonal of the intra-chromosomal interaction space. This results in the smallest possible minimum bounding box, which will only increase in size if interactions are placed on the other side of the diagonal. Alternatively, users can specify a StrictGInteractions object as an input into `x`, in which `anchor1 <= anchor2` is enforced automatically.

**Value**

A GInteractions object containing the coordinates of each bounding box.

**Author(s)**

Aaron Lun

**See Also**

[GInteractions-class](#), [InteractionSet-class](#)

**Examples**

```
example(GInteractions, echo=FALSE)

# Making up a sensible grouping.
gi <- sort(gi)
all.chrs <- as.character(seqnames(regions(gi)))
f <- paste0(all.chrs[anchors(gi, type="first", id=TRUE)], ".",
            all.chrs[anchors(gi, type="second", id=TRUE)])

boundingBox(gi, f)
boundingBox(swapAnchors(gi), f)

# Fails for multiple chromosomes
try(out <- boundingBox(gi))
in.A <- f=="chrA.chrA"
out <- boundingBox(gi[in.A])
```

---

ContactMatrix accessors

*ContactMatrix accessors*

---

**Description**

Methods to get and set fields in an ContactMatrix object.

**Usage**

```
## S4 method for signature 'ContactMatrix'
anchors(x, type="both", id=FALSE)
## S4 replacement method for signature 'ContactMatrix'
anchors(x) <- value

## S4 method for signature 'ContactMatrix'
regions(x)
## S4 replacement method for signature 'ContactMatrix'
regions(x) <- value
## S4 replacement method for signature 'ContactMatrix'
replaceRegions(x) <- value
## S4 replacement method for signature 'ContactMatrix'
appendRegions(x) <- value
```

```

## S4 method for signature 'ContactMatrix'
reduceRegions(x)

## S4 method for signature 'ContactMatrix'
show(x)
## S4 method for signature 'ContactMatrix'
as.matrix(x)
## S4 replacement method for signature 'ContactMatrix'
as.matrix(x) <- value
## S4 method for signature 'ContactMatrix'
t(x)

## S4 method for signature 'ContactMatrix'
dim(x)
## S4 method for signature 'ContactMatrix'
dimnames(x)
## S4 replacement method for signature 'ContactMatrix'
dimnames(x) <- value
## S4 method for signature 'ContactMatrix'
length(x)

```

### Arguments

<code>x</code>	A <code>ContactMatrix</code> object.
<code>type</code>	a string specifying which anchors are to be extracted or replaced.
<code>id</code>	a scalar logical indicating whether indices or <code>GRanges</code> should be returned.
<code>value</code>	For <code>anchors&lt;-</code> and <code>type="row"</code> or <code>"column"</code> , an integer vector of length equal to the number of rows or columns in <code>x</code> , respectively. For <code>type="both"</code> , a list of two such vectors must be supplied (first and second for rows and columns, respectively). For <code>anchors&lt;-</code> , a list of two integer vectors when <code>type="both"</code> . The first and second vectors must have length equal to the number of rows and columns of <code>x</code> , respectively. For <code>type="row"</code> or <code>"column"</code> , only one vector needs to be supplied (rows or columns, respectively). For <code>regions&lt;-</code> , a <code>GRanges</code> object of length equal to that of <code>regions(x)</code> . For <code>newRegions&lt;-</code> , a <code>GRanges</code> object that is a superset of all entries in <code>regions(x)</code> involved in interactions. For <code>appendRegions&lt;-</code> , a <code>GRanges</code> of any length containing additional regions. For <code>as.matrix&lt;-</code> , a <code>Matrix</code> object of the same dimensions as that in the <code>matrix</code> slot. For <code>dimnames&lt;-</code> , a list of two character vectors corresponding to the row and column names, respectively. These can also be passed separately via <code>rownames&lt;-</code> and <code>colnames&lt;-</code> .

### Details

The return value of `anchors` varies depending on `type` and `id`. If `id=FALSE`, a `GRanges` object will be returned containing the regions specified by the `anchor1` or `anchor2` slots in `x`, for `type=="row"` or `"column"`, respectively. If `type="both"`, a `GRangesList` will be returned with two entries - `row` and `column`, containing regions specified by `anchor1` and `anchor2` respectively. However, if `id=TRUE`, the integer vectors in the `anchor1` or `anchor2` slots of `x` are returned directly (a list of length two is returned for `type="both"`, containing both of the vectors).

Replacement in `anchors<-` can only be performed using anchor indices. If `type="both"`, a list of two integer vectors is required in `value`, specifying the indices of the row- and column-wise interacting regions in `x`. If `type="row"` or `"column"`, an integer vector is required to replace the existing row- or column-wise indices in the `anchor1` or `anchor2` slot, respectively.

For regions, a `GRanges` is returned equal to the `regions` slot in `x`. For `regions<-`, the `GRanges` in `value` is used to replace the `regions` slot. Resorting of the replacement `GRanges` is performed automatically, along with re-indexing of the anchors. In addition, the input `GRanges` must be of the same length as the existing object in the `regions` slot. The `newRegions` replacement method can take variable length `GRanges`, but requires that the replacement contain (at least) all ranges contained in `anchors(x)`. The `appendRegions` replacement method appends extra intervals to the existing `regions` slot of `x`. The `reduceRegions` method removes unused entries in the `regions` slot, to save memory – see [reduceRegions, GInteractions-method](#) for more details.

The `show` method will print out various details of the object, such as the dimensions of the `matrix` slot and the length of the `regions` slot. The `as.matrix` method will return the value of the `matrix` slot, containing the Matrix of interaction data. Replacement with a Matrix of the same dimensions can be performed using the `as.matrix<-` function. The `t` method will transpose the matrix, i.e., switch the rows and columns (and switch the vectors in the `anchor1` and `anchor2` slots).

The `dim` method will return a vector of length 2, containing the dimensions of the `matrix` slot. The `dimnames` method will return a list of length 2, containing the row and column names of `matrix` (these can be modified with the `dimnames<-` method). The `length` method will return an integer scalar corresponding to the total number of entries in the `matrix` slot.

As the `ContactMatrix` class inherits from the [Annotated](#) class, additional metadata can be stored in the `metadata` slot. This can be accessed or modified with [metadata, Annotated-method](#).

## Value

For the getters, values in various slots of `x` are returned, while for the setters, the slots of `x` are modified accordingly – see [Details](#).

## Author(s)

Aaron Lun

## See Also

[ContactMatrix-class](#)

## Examples

```
example(ContactMatrix, echo=FALSE) # Generate a nice object.
show(x)

# Various matrix methods:
as.matrix(x)
t(x)

nrow(x)
ncol(x)
length(x)

# Accessing anchor ranges or indices:
anchors(x)
anchors(x, type="row")
```

```

anchors(x, id=TRUE)

anchors(x, id=TRUE, type="row")
anchors(x, id=TRUE, type="column")

# Modifying anchor indices:
nregs <- length(regions(x))
anchors(x) <- list(sample(nregs, nrow(x), replace=TRUE),
                  sample(nregs, ncol(x), replace=TRUE))
anchors(x, id=TRUE, type="row")
anchors(x, id=TRUE, type="column")

# Accessing or modifying regions:
regions(x)
regions(x)$score <- runif(length(regions(x)))

new.ranges <- c(regions(x), resize(regions(x), fix="center", width=50))
try(regions(x) <- new.ranges) # Fails
replaceRegions(x) <- new.ranges # Succeeds

length(regions(x))
appendRegions(x) <- GRanges("chrA", IRanges(5:10+1000, 1100+5:10), score=runif(6))
length(regions(x))

reduceRegions(x)

# Setting metadata
metadata(x)$name <- "I am a contact matrix"
metadata(x)

```

---

ContactMatrix class    *ContactMatrix class and constructors*

---

## Description

The ContactMatrix class contains a matrix where rows and columns represent genomic loci. Each entry of the matrix contains information about the interaction between the loci represented by the corresponding row/column, e.g., contact frequencies. Coordinates of the loci are also contained within this class.

## Usage

```

## S4 method for signature 'ANY,numeric,numeric,GRanges'
ContactMatrix(matrix, anchor1, anchor2, regions, metadata=list())

## S4 method for signature 'ANY,GRanges,GRanges,GenomicRangesORmissing'
ContactMatrix(matrix, anchor1, anchor2, regions, metadata=list())

## S4 method for signature 'missing,missing,missing,GenomicRangesORmissing'
ContactMatrix(matrix, anchor1, anchor2, regions, metadata=list())

```

## Arguments

matrix	Any object containing interaction data that can be coerced into a Matrix object.
anchor1, anchor2	Either a pair of numeric vectors containing indices to regions or a pair of GRanges objects. In both cases, anchor1 and anchor2 should have lengths equal to the number of rows and columns in matrix, respectively.
regions	A GRanges object containing the coordinates of the interacting regions. This argument is optional for InteractionSet, ANY, GRanges, GRanges-method.
metadata	A list containing experiment-wide metadata - see <a href="#">?Annotated</a> for more details.

## Details

The ContactMatrix class inherits from the [Annotated](#) class, with several additional slots:

matrix: A Matrix object.

anchor1: An integer vector specifying the index of the first interacting region.

anchor2: An integer vector specifying the index of the second interacting region.

regions: A sorted GRanges object containing the coordinates of all interacting regions.

Each entry of anchor1 corresponds to a row in matrix, while each entry of anchor2 corresponds to a column. Each entry of matrix represents an interaction between the corresponding entries in anchor1 and anchor2, i which point to the relevant coordinates in regions for each locus.

ContactMatrix objects can be constructed by specifying numeric vectors as anchor1 and anchor2 in the ContactMatrix function. These vectors will define the regions corresponding to the rows and columns of the matrix. Specifically, each value of the vector acts as an index to specify the relevant coordinates from regions. This means that the range of entries must lie within [1, length(regions)].

Alternatively, ContactMatrix objects can be constructed by directly supplying the GRanges of the interacting loci in ContactMatrix. If regions is not specified, this will be constructed automatically from the two sets of supplied GRanges. If regions is supplied, exact matching will be performed to identify the indices in regions corresponding to the regions in the supplied GRanges. Missing values are not tolerated and will cause an error to be raised.

Both methods will return a ContactMatrix object containing all of the specified information. Sorting of regions is also performed automatically, with re-indexing of all anchor indices to preserve the correct pairings between regions.

## Value

For the constructors, a ContactMatrix object is returned.

## Choosing between Matrix classes

The ContactMatrix class provides support for Matrix objects from the Matrix package. This provides several classes for storing sparse and symmetric matrices in a memory-efficient manner, e.g., via the sparseMatrix class. These classes may be preferable for handling very large matrices spanning substantial areas in the interaction space.

There are, however, some caveats to the use of the various Matrix subclasses. All numeric values are coerced to double-precision types, which may take up more memory than a direct integer representation. Character data is not supported, though this is unlikely to be relevant in most cases. Another issue is how missing values should be interpreted in the sparseMatrix – see [?inflate](#) for more details.

In general, the choice of class depends on the type of data and the intended application. A dense Matrix is often sufficient for most use cases where small or dense regions of the interaction space are being examined. Switching to sparseMatrix may be desirable in emptier parts of the space, where there can be substantial memory savings.

### Author(s)

Aaron Lun

### See Also

[ContactMatrix-access](#), [ContactMatrix-subset](#), [ContactMatrix-sort](#), [Annotated-class](#), [Matrix-class](#)

### Examples

```
set.seed(1000)
N <- 30
all.starts <- round(runif(N, 1, 100))
all.ends <- all.starts + round(runif(N, 5, 20))
all.regions <- GRanges(rep(c("chrA", "chrB"), c(N-10, 10)),
  IRanges(all.starts, all.ends))

Nr <- 10
Nc <- 20
all.anchor1 <- sample(N, Nr)
all.anchor2 <- sample(N, Nc)
counts <- matrix(rpois(Nr*Nc, lambda=10), Nr, Nc)
x <- ContactMatrix(counts, all.anchor1, all.anchor2, all.regions)

# Equivalent construction:
x <- ContactMatrix(counts, all.regions[all.anchor1],
  all.regions[all.anchor2])
x <- ContactMatrix(counts, all.regions[all.anchor1],
  all.regions[all.anchor2], all.regions)

# Also works directly with Matrix objects.
counts2 <- Matrix::Matrix(counts)
x2 <- ContactMatrix(counts2, all.anchor1, all.anchor2, all.regions)
counts2 <- as(counts2, "dgCMatrix")
x2 <- ContactMatrix(counts2, all.anchor1, all.anchor2, all.regions)
```

---

ContactMatrix distances

*Compute linear distances from ContactMatrix objects*

---

### Description

Methods to compute linear distances between pairs of interacting regions in a ContactMatrix object.

## Usage

```
## S4 method for signature 'ContactMatrix'  
pairdist(x, type="mid")
```

```
## S4 method for signature 'ContactMatrix'  
intrachr(x)
```

## Arguments

x	A ContactMatrix object.
type	A character string specifying the type of distance to compute. See <a href="#">?pairdist, InteractionSet-method</a> for possible values.

## Details

`pairdist, ContactMatrix-method` will return a matrix of integer (or, if `type="intra"`, logical) values. Each entry of this matrix specifies the distance between the interacting loci that are represented by the corresponding row and column. If `type="intra"`, each entry indicates whether the corresponding interaction is intra-chromosomal. Running `intrachr(x)` is equivalent to `pairdist(x, type="intra")` for any ContactMatrix object `x`. See [pairdist, InteractionSet-method](#) for more details on the type of distances that can be computed.

## Value

An integer or logical matrix of the same dimensions as `x`, containing the specified distances.

## Author(s)

Aaron Lun

## See Also

[ContactMatrix-class](#), [pairdist](#), [InteractionSet-method](#)

## Examples

```
example(ContactMatrix, echo=FALSE)  
  
pairdist(x)  
pairdist(x, type="gap")  
pairdist(x, type="span")  
pairdist(x, type="diag")  
  
intrachr(x)
```

---

ContactMatrix overlaps

*Find overlaps between GRanges and a ContactMatrix*

---

**Description**

Find overlaps between a set of linear intervals in a GRanges object, and the set of regions representing the rows or columns of a ContactMatrix.

**Usage**

```
## S4 method for signature 'ContactMatrix,GRanges'
overlapsAny(query, subject, maxgap=0L, minoverlap=1L,
            type=c("any", "start", "end", "within", "equal"),
            ignore.strand=TRUE)
```

```
## S4 method for signature 'ContactMatrix,GRangesList'
overlapsAny(query, subject, maxgap=0L, minoverlap=1L,
            type=c("any", "start", "end", "within", "equal"),
            ignore.strand=TRUE, use.region="both")
```

```
# The same call is used for the methods taking GRangesList, GInteractions and
# InteractionSet objects as 'subject'. For brevity, these will not be listed.
```

**Arguments**

query	A ContactMatrix object.
subject	A GRanges, GRangesList, GInteractions or InteractionSet object.
maxgap, minoverlap, type	See <a href="#">?findOverlaps</a> in the <b>GenomicRanges</b> package.
ignore.strand	See <a href="#">?findOverlaps</a> in the <b>GenomicRanges</b> package.
use.region	A string specifying how the interacting regions are to be matched to row/column regions.

**Details**

When subject is a GRanges, overlaps are identified between the row regions of query and the regions in subject. This is repeated for the column regions of query. A list of two logical vectors is returned, specifying the rows and columns in query that are overlapped by any region in subject. These vectors can be directly used to subset query. Alternatively, they can be used in outer to generate a logical matrix for masking – see Examples.

For all other classes of subject, two-dimensional overlaps are identified. A logical matrix is returned indicating which entries in the ContactMatrix have overlaps with the specified interactions. For any given entry, an overlap is only considered if the regions for the corresponding row and column both overlap anchor regions in the subject. See [?"findOverlaps,GInteractions,GInteractions-method"](#) for more details.

If use.region="both", overlaps between any row/column region and the first/second interacting region of subject are considered. If use.region="same", only overlaps between row regions and the first interacting regions, or between column regions and the second interacting regions are considered. If use.region="reverse", only overlaps between row regions and the second interacting regions, or between row regions and the first interacting regions are considered.

**Value**

For `overlapsAny`, `ContactMatrix`, `GRanges`-method, a named list of two logical vectors is returned specifying the rows of columns of query overlapped by subject.

For the other `overlapsAny` methods, a logical matrix is returned indicating which entries in query are overlapped by subject.

**Author(s)**

Aaron Lun

**See Also**

[ContactMatrix-class](#), [findOverlaps](#)

**Examples**

```
example(ContactMatrix, echo=FALSE)

of.interest <- resize(sample(regions(x), 2), width=1, fix="center")
olap <- overlapsAny(x, of.interest)
olap
x[olap$row,]
x[,olap$column]
x[olap$row,olap$column]

keep <- outer(olap$row, olap$column, "|") # OR mask
temp <- as.matrix(x)
temp[!keep] <- NA

keep <- outer(olap$row, olap$column, "&") # AND mask
temp <- as.matrix(x)
temp[!keep] <- NA

# Two dimensional overlaps.
pairing <- GRangesList(first=regions(x), second=regions(x))
olap <- overlapsAny(x, pairing)
olap
olap <- overlapsAny(sort(x), pairing) # A bit prettier
olap
```

---

ContactMatrix sorting *ContactMatrix sorting and ordering*

---

**Description**

Methods to sort and order `ContactMatrix` objects, based on the anchor indices. Also, methods to remove duplicate rows or columns in each `ContactMatrix`.

**Usage**

```
## S4 method for signature 'ContactMatrix'
order(..., na.last=TRUE, decreasing=FALSE)
## S4 method for signature 'ContactMatrix'
sort(x, decreasing=FALSE, ...)
## S4 method for signature 'ContactMatrix'
duplicated(x, incomparables=FALSE, fromLast=FALSE, ...)
## S4 method for signature 'ContactMatrix'
unique(x, incomparables=FALSE, fromLast=FALSE, ...)
```

**Arguments**

...	For <code>sort, ContactMatrix</code> -method, one or more <code>ContactMatrix</code> objects with the same dimensions. Otherwise, ignored in all other methods.
x	A <code>ContactMatrix</code> object.
na.last	A logical scalar indicating whether NA values should be ordered last. This should not be relevant as anchor indices should be finite.
decreasing	A logical scalar indicating whether sorting should be performed in decreasing order.
incomparables	A logical scalar, ignored.
fromLast	A logical scalar indicating whether the last entry of a repeated set in x should be considered as a non-duplicate.

**Details**

Sorting is performed based on the anchor indices of the `ContactMatrix` object. Rows are ordered for increasing values of the `anchor1` slot, while columns are ordered for increasing values of the `anchor2` slot. This equates to ordering by the coordinates directly, as the `GRanges` in the `regions` slot is always sorted. Based on this, `sort, ContactMatrix`-method will return a sorted copy of x with permuted rows/columns in increasing order. This can be set to decreasing order by specifying `decreasing=TRUE`.

`order, ContactMatrix`-method returns a list of 2 integer vectors. The first vector contains the permutation to rearrange the rows of x in increasing order, while the second vector does the same for the columns of x. If multiple objects are supplied to `order`, ordering will be computed using anchor indices from successive objects. In other words, ordering will be performed using anchor indices from the first object; any rows with the same `anchor1` or columns with the same `anchor2` will be split using the corresponding indices in the second object; and so on.

`duplicated, ContactMatrix`-method returns a list of two logical vectors. The first vector indicates whether rows are duplicated, based on identical values in the `anchor1` slot. The second does the same for columns based on the `anchor2` slot. For a set of duplicated rows or columns, the first occurrence of that row/column is marked as the non-duplicate if `fromLast=FALSE`, and the last entry otherwise.

`unique, ContactMatrix`-method returns a `ContactMatrix` object where all duplicate rows and columns have been removed from x. This is equivalent to subsetting based on the non-duplicate rows and columns defined in `duplicated, ContactMatrix`-method.

Note that sorting and duplicate identification only use the anchor indices. The values of the `matrix` slot are *not* used in distinguishing between rows or columns with the same index.

**Value**

For `sort` and `unique`, a `ContactMatrix` object is returned with sorted or unique rows/columns, respectively.

For `order`, a list of two integer vectors is returned containing row/column permutations.

For `duplicated`, a list of logical vectors is returned specifying which rows/columns are duplicated.

**Author(s)**

Aaron Lun

**See Also**

[ContactMatrix-class](#)

**Examples**

```
example(ContactMatrix, echo=FALSE)

anchors(x)
x2 <- sort(x)
x2
anchors(x2)

# Can also order them.
o <- order(x)
o
stopifnot(all.equal(x[o$row,o$column], x2))

# Checking duplication.
duplicated(x)
duplicated(rbind(x, x))
stopifnot(all.equal(x, unique(rbind(x, x))))
```

---

ContactMatrix subsetting

*ContactMatrix subsetting and combining*

---

**Description**

Methods to subset or combine `ContactMatrix` objects.

**Usage**

```
### Subsetting

## S4 method for signature 'ContactMatrix,ANY,ANY'
x[i, j, ..., drop=TRUE]
## S4 replacement method for signature 'ContactMatrix,ANY,ANY,ContactMatrix'
x[i, j] <- value
## S4 method for signature 'ContactMatrix'
subset(x, i, j)
```

```
### Combining

## S4 method for signature 'ContactMatrix'
cbind(..., deparse.level=1)
## S4 method for signature 'ContactMatrix'
rbind(..., deparse.level=1)
```

### Arguments

<code>x</code>	A ContactMatrix object.
<code>i, j</code>	A vector of subscripts, indicating the rows and columns to be subsetted for <code>i</code> and <code>j</code> , respectively.
<code>...</code>	For <code>cbind</code> , <code>rbind</code> and <code>c</code> , <code>...</code> contains ContactMatrix objects to be combined. Otherwise, this argument is ignored during subsetting.
<code>drop</code>	A logical scalar, ignored by <code>[, ContactMatrix, ANY, ANY-method</code> .
<code>value</code>	A ContactMatrix object with dimensions equal to the length of the two subscripts (or the corresponding dimensions of <code>x</code> , if either subscript is missing).
<code>deparse.level</code>	An integer scalar; see <code>?base::cbind</code> for a description of this argument.

### Details

Subsetting of ContactMatrix objects behaves like that for standard matrices. Rows and columns can be selected and rearranged, with concomitant changes to the `anchor1` and `anchor2` slots. All subsetting operations will return an ContactMatrix with the specified rows and columns. However, note that the value of regions will not be modified by subsetting.

`cbind` will combines objects with the same rows but different columns. Errors will occur if the row regions are not identical between objects (i.e., must have same values in the slots for regions and `anchor1`). Conversely, `rbind` will combines objects with the same columns but different rows. Again, errors will occur if the columns are not identical (i.e., must have same values in the slots for regions and `anchor2`).

If subsetted anchors are required, see `?interaction-subset` for why subsetting should be done before calling the `anchors` method.

### Value

A subsetted or combined ContactMatrix object.

### Author(s)

Aaron Lun

### See Also

[ContactMatrix-class](#)

### Examples

```
example(ContactMatrix, echo=FALSE)

# Subsetting:
x[1:5,]
x[,10:15]
```

```
x[1:5,10:15]

x2 <- x
x2[1:5,] <- x[6:10,]
as.matrix(x2[,1]) <- 20

# Combining
cbind(x, x)
rbind(x, x)
```

---

 Convert classes

 Convert between classes
 

---

## Description

Inflate a `GInteractions` or `InteractionSet` into a `ContactMatrix`, or deflate a `ContactMatrix` to an `InteractionSet`.

## Usage

```
## S4 method for signature 'GInteractions'
inflate(x, rows, columns, fill, swap=TRUE, sparse=FALSE, ...)

## S4 method for signature 'InteractionSet'
inflate(x, rows, columns, assay=1L, sample=1L, fill=NULL, swap=TRUE, sparse=FALSE, ...)

## S4 method for signature 'ContactMatrix'
deflate(x, collapse=TRUE, extract, use.zero, use.na, ...)
```

## Arguments

<code>x</code>	A <code>GInteractions</code> or <code>InteractionSet</code> object for <code>inflate</code> , or a <code>ContactMatrix</code> object for <code>deflate</code> .
<code>rows, columns</code>	An integer, logical or character vector, a <code>GRanges</code> object or <code>NULL</code> , indicating the regions of interest to be used as the rows or columns of the <code>ContactMatrix</code> .
<code>assay</code>	A numeric scalar indicating the assay of the <code>InteractionSet</code> object, from which values are extracted to fill the <code>ContactMatrix</code> .
<code>sample</code>	A numeric scalar indicating the sample (i.e., column) of the assay to extract values to fill the <code>ContactMatrix</code> .
<code>fill</code>	A numeric vector of length equal to <code>nrow(x)</code> , containing values with which to fill the <code>ContactMatrix</code> . If specified, this overrides extraction of assay values for <code>inflate, InteractionSet-method</code> .
<code>swap</code>	A logical scalar indicating whether filling should also be performed after swapping anchor indices.
<code>sparse</code>	A logical scalar indicating whether the inflated matrix should use a <code>sparseMatrix</code> representation.
<code>collapse</code>	A logical scalar indicating whether duplicated interactions should be removed from <code>x</code> prior to deflation.
<code>extract</code>	A logical vector or matrix indicating which entries of <code>x</code> to convert into pairwise interactions.

`use.zero`, `use.na`  
 A logical scalar indicating whether to convert zero or NA entries into pairwise interactions.

...  
 For `inflate`, additional arguments to pass to `overlapsAny` when rows or columns is a `GRanges`.  
 For `deflate`, additional arguments to pass to the `InteractionSet` constructor.

### Value

For `inflate`, a `ContactMatrix` is returned.

For `deflate`, an `InteractionSet` object is returned.

### Inflating to a ContactMatrix

The `inflate` method will return a `ContactMatrix` where the rows and columns correspond to specified regions of interest in rows and columns. Regions can be specified by supplying an object of various types:

- If it is an integer vector, it is assumed to refer to intervals in the `regions` slot of the input object `x`. Values of the vector need not be sorted or unique, but must lie within `[1, regions(x)]`.
- If it is a logical vector, it will subset to retain intervals in `regions(x)` that are `TRUE`.
- If it is a character vector, it is assumed to contain the names of the reference sequences of interest (i.e., chromosome names).
- If it is a `GRanges` object, `overlapsAny` will be called to identify the overlapping intervals of `regions(x)`.
- If it is `NULL`, all regions in `regions(x)` will be used to construct that dimension of the `ContactMatrix`.

For the `GInteractions` method, values in the matrix are filled based on user-supplied values in `fill`. Each entry of `fill` corresponds to an interaction in `x` is used to set the matrix value at the matching row/column. Some entries of the matrix will correspond to pairwise interactions that are not present in `x` - these are filled with NA values.

By default, filling is repeated after swapping the anchor indices. This means that the value of the matrix at (1, 2) will be the same as that at (2, 1), i.e., the matrix is symmetric around the diagonal of the interaction space. However, if `swap=FALSE`, filling is performed so that the first and second anchor indices correspond strictly to rows and columns, respectively. This may be preferable if the order of the anchors contains some relevant information. In all cases, if duplicated interactions are present in `x` (and redundant permutations, when `swap=TRUE`), one will be arbitrarily chosen to fill the matrix.

For the `InteractionSet` `inflate` method, entries in the matrix are filled in based on the values in the first sample of the first assay by default. For more complex `x`, values from different assays and samples can be extracted using the `assay` and `sample` arguments. Note that if `fill` is specified, it will override any extraction of values from the assays.

If `sparse=TRUE`, `inflate` will return a `ContactMatrix` containing a `sparseMatrix` in the `matrix` slot. Here, entries without a corresponding interaction in `x` are set to zero, not NA.

### Deflating from a ContactMatrix

The `deflate` method will return an `InteractionSet` where each relevant entry in the `ContactMatrix` is converted into a pairwise interaction. Relevant entries are defined as those that are non-zero, if `use.zero` is `FALSE`; and non-NA, if `use.na` is `FALSE`. If `x` contains a `sparseMatrix` representation,

the former is set to FALSE while the latter is set to TRUE, if either are not specified. For all other matrices, use `.zero=TRUE` and use `.na=FALSE` by default.

If `extract` is specified, this overrides all values of `use.zero` and `use.na`. A typical application would be to deflate a number of `ContactMatrix` objects with the same `extract` matrix. This ensures that the resulting `InteractionSet` objects can be easily combined with `cbind`, as the interactions are guaranteed to be the same. Otherwise, different interactions may be extracted depending on the presence of zero or NA values.

The values of all matrix entries are stored as a one-sample assay, with each value corresponding to its pairwise interaction after conversion. Duplicate interactions are removed by default, along with redundant permutations of the anchor indices. These can be included in the returned object by setting `collapse=FALSE`. Additional arguments can be used to specify the `colData` and `metadata`, which are stored in the `ContactMatrix` itself.

### Interpreting zeroes in a sparse matrix

Storing data as a `sparseMatrix` may be helpful as it is more memory-efficient for sparse areas of the interaction space. However, users should keep in mind that the zero values in the `sparseMatrix` may not represent zeroes in `fill`. The majority of these values are likely to be zero just because there was no corresponding interaction in `x` to set it to a non-zero value.

Whether or not this is a problem depends on the application. For example, if `fill` represents count data and only interactions with non-zero counts are stored in `x`, then setting all other entries to zero is sensible. However, in other cases, it is not appropriate to fill entries corresponding to missing interactions with zero. If `fill` represents, e.g., log-fold changes, then setting missing entries to a value of zero will be misleading. One could simply ignore zeroes altogether, though this will also discard entries that are genuinely zero.

These problems are largely avoided with the default `denseMatrix` classes, where missing entries are simply set to NA.

### Author(s)

Aaron Lun

### See Also

[InteractionSet-class](#), [GInteractions-class](#), [ContactMatrix-class](#), [sparseMatrix-class](#)

### Examples

```
example(InteractionSet, echo=FALSE)

inflate(iset, 1:10, 1:10)
inflate(iset, 1:10, 1:10, sparse=TRUE)
inflate(iset, 1:10, 1:5+10)
inflate(iset, "chrA", 1:5+10)
inflate(iset, "chrA", "chrB")
inflate(iset, "chrA", GRanges("chrB", IRanges(1, 10)))

y <- inflate(iset, 1:10, 1:10)
iset2 <- deflate(y)
iset2
assay(iset2)

y <- inflate(iset, 1:10, 1:10, swap=FALSE)
```

```

iset2 <- deflate(y)
iset2
assay(iset2)

# Testing with different fillings:
y <- inflate(iset, 1:10, 1:10, sample=2)
iset2 <- deflate(y)
assay(iset2)

y <- inflate(iset, 1:10, 1:10, fill=rowSums(assay(iset)))
iset2 <- deflate(y)
assay(iset2)

y2 <- inflate(interactions(iset), 1:10, 1:10, rowSums(assay(iset)))
identical(y, y2) # should be TRUE

# Effect of 'collapse'
y <- inflate(iset, c(8, 1:10), 1:10)
deflate(y)
deflate(y, collapse=FALSE)

```

---

GInteractions class    *GInteractions class and constructors*

---

## Description

The GInteractions class stores pairwise genomic interactions, and is intended for use in data analysis from Hi-C or ChIA-PET experiments. Each row of the GInteractions corresponds to a pairwise interaction between two loci, with indexing to improve computational efficiency.

## Usage

```

## S4 method for signature 'numeric,numeric,GRanges'
GInteractions(anchor1, anchor2, regions, metadata=list(), mode="normal", ...)

## S4 method for signature 'GRanges,GRanges,GenomicRangesORmissing'
GInteractions(anchor1, anchor2, regions, metadata=list(), mode="normal", ...)

## S4 method for signature 'missing,missing,GenomicRangesORmissing'
GInteractions(anchor1, anchor2, regions, metadata=list(), mode="normal", ...)

```

## Arguments

anchor1, anchor2	Either a pair of numeric vectors containing indices to regions, or a pair of GRanges objects specifying the interacting loci. Lengths of both anchor1 and anchor2 must be equal.
regions	A GRanges object containing the coordinates of the interacting regions. This is only mandatory if anchor1 and anchor2 are numeric vectors.
metadata	An optional list of arbitrary content describing the overall experiment.
mode	A string indicating what type of GInteractions object should be constructed.
...	Optional metadata columns.

**Value**

For the constructors, a GInteractions (or StrictGInteractions, or ReverseStrictGInteractions) object is returned.

**Description of the class**

The GInteractions class inherits from the Vector class and has access to all of its data members and methods (e.g, metadata and elementMetadata - see [Vector-class](#) for more details). It also contains several additional slots:

anchor1: An integer vector specifying the index of the first interacting region.

anchor2: An integer vector specifying the index of the second interacting region.

regions: A sorted GRanges object containing the coordinates of all interacting regions.

Each interaction is defined by the corresponding entries in the anchor1 and anchor2 slots, which point to the relevant coordinates in regions for each locus.

The StrictGInteractions class inherits from the GInteractions class and has the same slots. The only difference is that, for each interaction, anchor1 must be less than or equal to anchor2. This means that the first interacting region has a start position that is "lower" than the second interacting region. This condition is useful for comparing interactions within and between objects, as it ensures that redundant permutations of the same interaction are not being overlooked. However, it is not used by default as there may conceivably be instances where the order of interactions is informative. The ReverseStrictGInteractions class has the opposite behaviour, where anchor1 must be greater than or equal to anchor2.

**Class construction**

GInteractions objects can be constructed by specifying integer vectors to define the pairwise interactions in the GInteractions call. For entry  $x$ , the corresponding interaction is defined between regions[anchor1[x]] and regions[anchor2[x]]. Obviously, coordinates of all of the interacting loci must be specified in the regions argument. Any metadata in regions will be preserved. Note that regions will be resorted in the returned object, so the anchor indices may not be equal to the input anchor1 and anchor2.

Alternatively, GInteractions objects can be constructed by directly supplying the GRanges of the interacting loci to the GInteractions function. If regions is not specified, this will be constructed automatically from the two sets of supplied GRanges. If regions is supplied, exact matching will be performed to identify the indices in regions corresponding to the regions in the supplied anchor GRanges. Missing values are not tolerated and will cause an error to be raised. In both cases, any metadata in the input GRanges will be transferred to the mcols of the output GInteractions object.

All constructors will return a GInteractions object containing all of the specified information. Sorting of regions is performed automatically, with re-indexing of all anchor indices to preserve the correct pairings between regions. If mode="strict", a StrictGInteractions object is returned with anchor indices swapped such that anchor1 <= anchor2 for all interactions. If mode="reverse", a ReverseStrictGInteractions object is returned with anchor indices swapped such that anchor1 >= anchor2. If both anchors are missing, the constructor will return an empty GInteractions object.

**Author(s)**

Aaron Lun, with contributions from Malcolm Perry and Liz Ing-Simmons.

**See Also**

[interaction-access](#), [interaction-subset](#), [interaction-compare](#), [Vector-class](#)

**Examples**

```

set.seed(1000)
N <- 30
all.starts <- round(runif(N, 1, 100))
all.ends <- all.starts + round(runif(N, 5, 20))
all.regions <- GRanges(rep(c("chrA", "chrB"), c(N-10, 10)),
  IRanges(all.starts, all.ends))

Np <- 20
all.anchor1 <- sample(N, Np)
all.anchor2 <- sample(N, Np)
gi <- GInteractions(all.anchor1, all.anchor2, all.regions)

# Equivalent construction:
gi <- GInteractions(all.regions[all.anchor1], all.regions[all.anchor2])
gi <- GInteractions(all.regions[all.anchor1], all.regions[all.anchor2],
  all.regions)

# Putting in metadata, elementMetadata
temp.gi <- gi
metadata(temp.gi)$name <- "My GI object"
mcols(temp.gi)$score <- runif(Np)

# Strict construction
sgi <- GInteractions(all.regions[all.anchor1], all.regions[all.anchor2],
  all.regions, mode="strict")
rsgi <- GInteractions(all.regions[all.anchor1], all.regions[all.anchor2],
  all.regions, mode="reverse")

```

---

GRanges methods

*Equivalents of GRanges methods*


---

**Description**

Methods for GInteractions, InteractionSet and ContactMatrix that operate on the internal GenomicRanges.

**Usage**

```

## S4 method for signature 'GInteractions'
trim(x, use.names=FALSE)
## S4 method for signature 'GInteractions'
width(x)

# Equivalent calls for InteractionSet and ContactMatrix.

```

**Arguments**

x                    A GInteractions, InteractionSet or ContactMatrix object.

use.names           See `?trim, GenomicRanges-method`.

## Details

The `trim` method will return an object of the same class as `x` where out-of-bound ranges in `regions(x)` are truncated to fit. This is followed by resorting of the regions and removal of redundant regions. As such, that the length of `regions(x)` and the anchor indices for each interaction may change before and after trimming. However, the number and order of the interactions will not change.

For `GInteractions` and `InteractionSet` objects, the `width` method will return a `DataFrame` with integer fields `anchor1` and `anchor2`. Each row contains the width of the two anchor regions corresponding to each interaction. For `ContactMatrix` objects, the method will return a list with the same fields (a different format is necessary as the lengths of `anchor1` and `anchor2` may be different).

## Value

Depending on the method, an object of the same class as `x`, or information regarding the genomic regions – see `Details`.

## Author(s)

Aaron Lun, based on suggestions from Liz Ing-Simmons.

## See Also

[trim](#), [width](#)

## Examples

```
example(GInteractions, echo=FALSE)

trim(gi)
width(gi)

example(ContactMatrix, echo=FALSE)
trim(x)
width(x)
```

---

Interaction accessors *Interaction accessors*

---

## Description

Methods to get and set fields in an `InteractionSet` or `GInteractions` object.

## Usage

```
## S4 method for signature 'GInteractions'
anchors(x, type="both", id=FALSE)
## S4 method for signature 'GInteractions'
regions(x)
## S4 method for signature 'GInteractions'
seqinfo(x)
## S4 method for signature 'GInteractions'
show(x)
```

```

## S4 replacement method for signature 'GInteractions'
anchors(x, type="both") <- value
## S4 replacement method for signature 'GInteractions'
regions(x) <- value
## S4 replacement method for signature 'GInteractions'
seqinfo(x) <- value

## S4 replacement method for signature 'GInteractions'
replaceRegions(x) <- value
## S4 replacement method for signature 'GInteractions'
appendRegions(x) <- value
## S4 method for signature 'GInteractions'
reduceRegions(x)

## S4 method for signature 'GInteractions'
names(x)
## S4 replacement method for signature 'GInteractions'
names(x) <- value

## Each above method has an identical equivalent for
## InteractionSet objects (not shown here for brevity).

## S4 replacement method for signature 'StrictGInteractions'
anchors(x, type="both") <- value
## S4 replacement method for signature 'ReverseStrictGInteractions'
anchors(x, type="both") <- value

## S4 method for signature 'GInteractions'
length(x)
## S4 method for signature 'GInteractions'
as.data.frame(x, row.names=NULL, optional=FALSE, ...)
## S4 method for signature 'GInteractions'
x$name
## S4 replacement method for signature 'GInteractions'
x$name <- value

## S4 method for signature 'InteractionSet'
interactions(x)
## S4 replacement method for signature 'InteractionSet'
interactions(x) <- value
## S4 method for signature 'InteractionSet'
mcols(x, use.names=FALSE)
## S4 replacement method for signature 'InteractionSet'
mcols(x) <- value

```

### Arguments

<code>x</code>	An <code>InteractionSet</code> or <code>GInteractions</code> object.
<code>type</code>	a string specifying which anchors are to be extracted or replaced.
<code>id</code>	a scalar logical indicating whether indices or <code>GRanges</code> should be returned.
<code>name</code>	a string indicating the field of <code>mcols</code> to be accessed or modified for a <code>GInteractions</code> object.

value	<p>For <code>anchors&lt;-</code> and <code>type="first"</code> or <code>"second"</code>, an integer vector of length equal to the number of rows in <code>x</code>. For <code>type="both"</code>, a list of two such vectors must be supplied.</p> <p>For <code>regions&lt;-</code>, a <code>GRanges</code> object of length equal to that of <code>regions(x)</code>. For <code>replaceRegions&lt;-</code>, a <code>GRanges</code> object that is a superset of all entries in <code>regions(x)</code> involved in interactions. For <code>appendRegions&lt;-</code>, a <code>GRanges</code> of any length containing additional regions.</p> <p>For <code>seqinfo&lt;-</code>, a <code>SeqInfo</code> object containing chromosome length data for all regions. For <code>interactions&lt;-</code>, a <code>GInteractions</code> object of length equal to that of <code>interactions(x)</code>. For <code>mcols&lt;-</code>, a <code>DataFrame</code> with number of rows equal to the length of <code>x</code>. For <code>names&lt;-</code>, a character vector of length equal to that of <code>x</code>.</p>
row.names, optional, ...	Additional arguments, see <code>?as.data.frame</code> for more details.
use.names	A logical scalar, see <code>?mcols</code> for more details.

### Value

For the getters, values in various slots of `x` are returned, while for the setters, the slots of `x` are modified accordingly – see below for more details.

### Anchor manipulations for GInteractions

The return value of `anchors` varies depending on `type` and `id`. If `id=FALSE`, a `GRanges` object will be returned containing the regions specified by the `anchor1` or `anchor2` slots in `x`, for `type="first"` or `"second"`, respectively. If `type="both"`, a `GRangesList` will be returned with two entries - `first` and `second`, containing regions specified by `anchor1` and `anchor2` respectively.

If `id=TRUE`, the integer vectors in the `anchor1` or `anchor2` slots of `x` are returned directly. A list of length two is returned for `type="both"`, containing both of the vectors. Otherwise, each vector is returned corresponding to the requested value of `type`.

Replacement in `anchors<-` can only be performed using anchor indices. If `type="both"`, a list of two integer vectors is required in `value`, specifying the indices of the interacting regions in `regions(x)`. If `type="first"` or `"second"`, an integer vector is required to replace the existing values in the `anchor1` or `anchor2` slot, respectively. If the object is a `StrictGInteractions`, indices are automatically swapped so that `anchor1 >= anchor2` for each interaction. The opposite applies if the object is a `ReverseStrictGInteractions`.

### Region manipulations for GInteractions

For `regions`, a `GRanges` is returned equal to the `regions` slot in `x`. For `regions<-`, the `GRanges` in `value` is used to replace the `regions` slot. Resorting of the replacement `GRanges` is performed automatically, along with re-indexing of the anchors. In addition, the replacement must be of the same length as the existing object in the `regions` slot.

The `replaceRegions` replacement method can take variable length `GRanges`, but requires that the replacement contain (at least) all ranges contained in `anchors(x)`. The `appendRegions` replacement method appends extra intervals to the existing `regions` slot of `x`. This is more efficient than calling `replaceRegions` on a concatenated object with `regions(x)` and the extra intervals.

The `reduceRegions` method will return a `GInteractions` object where the `regions` slot is reduced to only those entries used in `anchors(x)`. This may save some memory in each object by removing unused regions. However, this is not recommended for large workflows with many `GInteractions` objects. R uses a copy-on-write memory management scheme, so only one copy of the `GRanges` in `regions` should be stored so long as it is not modified in different objects.

### Other methods for GInteractions

For access and setting of all other slots, see [Vector-class](#) for details on the appropriate methods. This includes `mcols` or `metadata`, to store interactions-specific or experiment-wide metadata, respectively. The `length` method will return the number of interactions stored in `x`.

The `show` method will print out the class, the number of pairwise interactions, and the total number of regions in the `GInteractions` object. The number and names of the various metadata fields will also be printed. The `as.data.frame` method will return a `data.frame` object containing the coordinates for the two anchor regions as well as any element-wise metadata.

The `seqinfo` method will return the sequence information of the `GRanges` in the `regions` slot. This can be replaced with the `seqinfo<-` method – see [?Seqinfo](#) for more details.

### Details for InteractionSet

Almost all `InteractionSet` methods operate by calling the equivalent method for the `GInteractions` object, and returning the resulting value. The only exception is `interactions`, which returns the `GInteractions` object in the `interactions` slot of the `InteractionSet` object. This slot can also be set by supplying a valid `GInteractions` object in `interactions<-`.

The `show` method will print information equivalent to that done for a `SummarizedExperiment` object. An additional line is added indicating the number of regions in the `regions` slot of the object.

For access and setting of all other slots, see [SummarizedExperiment-class](#) for details on the appropriate methods. This includes `assays`, `colData`, `mcols` or `metadata`, which can all be applied to `InteractionSet` objects.

### Handling different metadata

Note that there are several options for metadata - experiment-wide metadata, which goes into `metadata(x)<-`; region-specific metadata (e.g., adjacent genes, promoter/enhancer identity, GC content), which goes into `mcols(regions(x))<-`; and interaction-specific metadata (e.g., interaction distance, interaction type), which goes directly into `mcols(x)<-`. This is applicable to both `GInteractions` and `InteractionSet` objects. In addition, library-specific data (e.g., library size) should be placed into `colData(x)<-` for `InteractionSet` objects.

Users should take care with the differences in the `$` and `$<-` operators between these two classes. For `GInteractions` objects, this will access or modify fields in the `mcols` slot, i.e., for interaction-specific metadata. For `InteractionSet` objects, this will access or modify fields in the `colData` slot, i.e., for library-specific data. The difference in behaviour is due to the concept of libraries in the `InteractionSet`, which is lacking in the `GInteractions` class.

For `InteractionSet` objects, the `mcols` getter and setter functions operate on the `GInteractions` object stored in `interactions` slot, rather than accessing the `elementMetadata` slot of the `SummarizedExperiment` base class. This makes no difference for practical usage in the vast majority of cases, except that any metadata stored in this manner is carried over when the `GInteractions` object is extracted with `interactions(x)`. Similarly, the `names` getter and setter will operate the names of the internal `GInteractions` object. However, the `metadata` getter and setter will operate on the `SummarizedExperiment` base class, *not* on the internal `GInteractions` object.

### Author(s)

Aaron Lun

### See Also

[GInteractions-class](#), [InteractionSet-class](#), [Vector-class](#), [SummarizedExperiment-class](#)

**Examples**

```

example(GInteractions, echo=FALSE) # Generate a nice object.
show(gi)

# Accessing anchor ranges or indices:
anchors(gi)
anchors(gi, type="first")
anchors(gi, id=TRUE)

anchors(gi, id=TRUE, type="first")
anchors(gi, id=TRUE, type="second")

# Modifying anchor indices:
nregs <- length(regions(gi))
mod <- list(sample(nregs, length(gi), replace=TRUE),
            sample(nregs, length(gi), replace=TRUE))
anchors(gi) <- mod
anchors(gi, id=TRUE, type="first")
anchors(gi, id=TRUE, type="second")

anchors(gi, type="both") <- mod
anchors(gi, type="first") <- mod[[1]]
anchors(gi, type="first") <- mod[[2]]

# Accessing or modifying regions:
regions(gi)
reduceRegions(gi)
regions(gi)$score <- runif(length(regions(gi)))

new.ranges <- c(regions(gi), resize(regions(gi), fix="center", width=50))
try(regions(gi) <- new.ranges) # Fails
replaceRegions(gi) <- new.ranges # Succeeds

length(regions(gi))
appendRegions(gi) <- GRanges("chrA", IRanges(5:10+1000, 1100+5:10), score=runif(6))
length(regions(gi))

seqinfo(gi)
seqinfo(gi) <- Seqinfo(seqnames=c("chrA", "chrB"), seqlengths=c(1000, 2000))

# Accessing or modifying metadata.
gi$score <- runif(length(gi))
mcols(gi)
as.data.frame(gi)

#####
# Same can be done for an InteractionSet object:

example(InteractionSet, echo=FALSE)

anchors(iset)
regions(iset)
reduceRegions(iset)
regions(iset)$score <- regions(gi)$score
replaceRegions(iset) <- new.ranges

```

```

seqinfo(iset)
seqinfo(iset) <- Seqinfo(seqnames=c("chrA", "chrB"), seqlengths=c(1000, 2000))

# Standard SE methods also available:
colData(iset)
metadata(iset)
mcols(iset)

# Note the differences in metadata storage:
metadata(iset)$name <- "metadata for SE0"
metadata(interactions(iset))$name <- "metadata for GI"

iset$lib.size <- runif(ncol(iset))*1e6
colData(iset)
mcols(iset) # untouched by "$" operator

mcols(iset)$whee <- runif(nrow(iset))
mcols(iset)
mcols(interactions(iset)) # preserved

names(iset) <- paste0("X", seq_along(iset))
names(iset)
names(interactions(iset))

```

---

Interaction compare     *Interaction comparisons*

---

## Description

Methods to order, compare and de-duplicate GInteractions or InteractionSet objects, based on the anchor indices.

## Usage

```

## S4 method for signature 'GInteractions'
order(..., na.last=TRUE, decreasing=FALSE)

## S4 method for signature 'GInteractions'
sort(x, decreasing=FALSE, ...)

## S4 method for signature 'GInteractions'
duplicated(x, incomparables=FALSE, fromLast=FALSE, ...)

## S4 method for signature 'GInteractions'
unique(x, incomparables=FALSE, fromLast=FALSE, ...)

## S4 method for signature 'GInteractions'
swapAnchors(x, mode=c("order", "reverse", "all"))

## Each of the above methods has an identical equivalent for
## InteractionSet objects (not shown for brevity).

## S4 method for signature 'GInteractions,GInteractions'

```

```

match(x, table, nomatch=NA_integer_, incomparables=NULL, ...)

# The same call for 'match' is used for all possible combinations of
# GInteractions and InteractionSet objects in 'x' or 'table'.
# For brevity, these will not be listed here.

## S4 method for signature 'GInteractions,GInteractions'
pcompare(x, y)

```

### Arguments

...	For order, one or more InteractionSet or GInteractions objects with the same number of rows. Otherwise, ignored in all other methods.
x	An InteractionSet or GInteractions object. For pcompare, a GInteractions object only.
na.last	A logical scalar indicating whether NA values should be ordered last. This should not be relevant as anchor indices should be finite.
decreasing	A logical scalar indicating whether rows should be sorted in decreasing order.
incomparables	A logical scalar. See <code>?match</code> for a description of this in <code>match</code> . Otherwise, it is ignored.
fromLast	A logical scalar indicating whether the last entry of a repeated set in x should be considered as a non-duplicate.
mode	A string indicating what type of swapping should be performed in <code>swapAnchors</code> .
table	A GInteractions or InteractionSet object.
nomatch	An integer scalar indicating the value to use for unmatched entries.
y	A GInteractions object, of the same length as x.

### Value

For `sort` and `unique`, a GInteractions or InteractionSet object is returned, depending on the class of x.

For `order` and `duplicate`, an integer vector of permutations or a logical vector of duplicate specifications is returned, respectively.

### Sorting and ordering

Sorting is performed based on the anchor indices of the GInteraction object. Rows are ordered for increasing values of the anchor1 slot - if these are equal, ordering is performed with values of the anchor2 slot. This equates to ordering by the coordinates directly, as the GRanges in the regions slot is always sorted. Based on this, `sort` will return a sorted copy of x with permuted rows in increasing order.

The `order` method returns an integer vector indicating the permutation to rearrange x in increasing order. If multiple objects are supplied to `order`, ordering will be computed using anchor indices from successive objects. For example, ordering is first performed using anchor indices from the first object; any rows with the same anchor1 and anchor2 will be split using the second object; and so on.

For both of these methods, the sorting can be reversed by setting `decreasing=TRUE`. This will sort or order for decreasing values of anchor1 and anchor2, rather than for increasing values.

### Removing duplicates

The `duplicated` method returns a logical vector indicating whether the rows of `x` are duplicated. Duplicated rows are identified on the basis of identical entries in the `anchor1` and `anchor2` slots. Values in other slots (e.g., in `mcols`) are ignored. For a set of duplicated rows, the first occurrence in `x` is marked as the non-duplicate if `fromLast=FALSE`, and the last entry otherwise.

`unique` returns a `GInteractions` object where all duplicate rows have been removed from `x`. This is equivalent to `x[!duplicated(x),]`, with any additional arguments passed to `duplicated` as specified.

### Matching and comparing

The `match` function will return an integer vector of length equal to that of `x`. Each entry of the vector corresponds to an interaction in `x` and contains the index of the first interaction table with the same anchor regions. Interactions in `x` without any matches are assigned NA values by default. Note that the `regions` slot must be identical between `x` and `table`.

The `pcompare` function will return an integer vector of length equal to `x` and `y`. This performs parallel comparisons between corresponding entries in two `GInteractions` objects, based on the values of the anchor indices (`anchor1` first, and then `anchor2` if `anchor1` is tied). Negative, zero and positive values indicate that the corresponding interaction in `x` is 'lesser', equal or 'greater' than the corresponding interaction in `y`. Again, the `regions` must be identical between the two objects.

### Swapping anchors

For `GInteractions` objects, `swapAnchors` returns a `GInteractions` object where `anchor1` and `anchor2` values are swapped. If `mode="order"`, this is done so that all values in the `anchor2` slot are not less than values in `anchor1` in the returned object. If `mode="reverse"`, all values in `anchor1` are not less than all values in `anchor2`. If `mode="all"`, the anchor indices in `x` are directly swapped without consideration of ordering.

It is recommended to apply this method before running methods like `order` or `duplicated`. This ensures that redundant permutations are coerced into the same format for a valid comparison. In many applications, permutations of pairwise interactions are not of interest, i.e., an interaction between regions 1 and 2 is the same as that between 2 and 1. Application of `swapAnchors` with `mode="order"` ensures that all indices are arranged in a comparable manner. Alternatively, users can use a `(Reverse)StrictGInteractions` object which enforces a consistent arrangement of indices across interactions.

### Methods for InteractionSet objects

For all `InteractionSet` methods, the corresponding method is called on the `GInteractions` object in the `interactions` slot of the `InteractionSet` object. Return values for each `InteractionSet` method is the same as those for the corresponding `GInteractions` method - except for `sort` and `unique`, which return a row-permuted or row-subsetted `InteractionSet`, respectively, instead of a `GInteractions` object; and `swapAnchors`, which returns an `InteractionSet` object where the internal `GInteractions` has its anchor indices swapped around.

Note that no additional information from the `InteractionSet` (beyond that in `interactions`) is used for sorting or duplicate marking, i.e., the assay or metadata values for each interaction are *not* used in distinguishing rows with the same interaction. For this reason, the `pcompare` method is not implemented for `InteractionSet` objects. It makes little sense to do a parallel comparison in an `InteractionSet` without examining the data.

**Author(s)**

Aaron Lun

**See Also**[GInteractions-class](#), [InteractionSet-class](#), [match](#), [pcompare](#)**Examples**

```

example(GInteractions, echo=FALSE)

anchors(gi, id=TRUE)
anchors(swapAnchors(gi, mode="all"), id=TRUE)
gi <- swapAnchors(gi)

anchors(gi)
gi2 <- sort(gi)
gi2
anchors(gi2)

# Can also order them.
o <- order(gi)
o
stopifnot(all.equal(gi[o], gi2))

# Checking duplication.
summary(duplicated(gi))
summary(duplicated(rbind(gi, gi)))
stopifnot(all.equal(gi, unique(rbind(gi, gi))))

# Matching and comparing.
another.gi <- gi[sample(length(gi))]
match(gi, another.gi)
match(gi, another.gi[1:5])

pcompare(gi, another.gi)

#####
# Same can be done for an InteractionSet object:

example(InteractionSet, echo=FALSE)
iset <- swapAnchors(iset)

anchors(iset)
anchors(sort(iset))
order(iset)
summary(duplicated(iset))
unique(iset)

```

**Description**

Compute the distance between interacting regions on the linear genome, for each pairwise interaction contained in a `GInteractions` or `InteractionSet` object.

**Usage**

```
## S4 method for signature 'GInteractions'
pairdist(x, type="mid")
## S4 method for signature 'InteractionSet'
pairdist(x, type="mid")

## S4 method for signature 'GInteractions'
intrachr(x)
## S4 method for signature 'InteractionSet'
intrachr(x)
```

**Arguments**

<code>x</code>	A <code>GInteractions</code> or <code>InteractionSet</code> object.
<code>type</code>	A character string specifying the type of distance to compute. Can take values of "mid", "gap", "span", "diag" or "intra".

**Details**

For each interaction in `x`, the `pairdist` method computes the distance between the two interacting regions. An integer vector is returned, with values computed according to the specified value of `type`:

"mid": The distance between the midpoints of the two regions (rounded down to the nearest integer) is returned.

"gap": The length of the gap between the closest points of the two regions is computed - negative lengths are returned for overlapping regions, indicating the length of the overlap.

"span": The distance between the furthestmost points of the two regions is computed.

"diag": The difference between the anchor indices is returned. This corresponds to a diagonal on the interaction space when bins are used in the `regions` slot of `x`.

Interchromosomal interactions are marked with NA. Alternatively, if `type="intra"`, a logical vector is returned indicating whether the interaction occurs between two regions on the same chromosome. `intrachr(x)` is an alias for `pairdist(x, type="intra")`.

The return values of the assorted methods are the same for both `GInteractions` and `InteractionSet` objects. Methods for the latter operate on the `GInteractions` object in the `interactions` slot.

**Value**

An integer or logical vector of distances.

**Author(s)**

Aaron Lun

**See Also**

[GInteractions-class](#), [InteractionSet-class](#)

**Examples**

```

example(GInteractions, echo=FALSE)

pairedist(gi)
pairedist(gi, type="gap")
pairedist(gi, type="span")
pairedist(gi, type="diag")
intrachr(gi)

example(InteractionSet, echo=FALSE)

pairedist(iset)
pairedist(iset, type="gap")
pairedist(iset, type="span")
pairedist(iset, type="diag")
intrachr(iset)

```

---

Interaction overlaps *Find overlaps between interactions in one or two dimensions*

---

**Description**

Find overlaps between interactions and linear intervals, between interactions and pairs of intervals, and between interactions and other interactions in a GInteractions or InteractionSet object.

**Usage**

```

## S4 method for signature 'GInteractions,Vector'
findOverlaps(query, subject, maxgap=0L, minoverlap=1L,
  type=c("any", "start", "end", "within", "equal"),
  select=c("all", "first", "last", "arbitrary"),
  ignore.strand=FALSE, use.region="both")

## S4 method for signature 'GInteractions,Vector'
overlapsAny(query, subject, maxgap=0L, minoverlap=1L,
  type=c("any", "start", "end", "within", "equal"),
  ignore.strand=FALSE, use.region="both")

## S4 method for signature 'GInteractions,Vector'
countOverlaps(query, subject, maxgap=0L, minoverlap=1L,
  type=c("any", "start", "end", "within", "equal"),
  ignore.strand=FALSE, use.region="both")

## S4 method for signature 'GInteractions,Vector'
subsetByOverlaps(query, subject, maxgap=0L, minoverlap=1L,
  type=c("any", "start", "end", "within", "equal"),
  ignore.strand=FALSE, use.region="both")

# For brevity, only 'GInteractions,Vector-methods' are listed. Methods are
# available for all pairwise combinations of GInteractions, InteractionSet, and
# Vector objects, so long as at least one InteractionSet or GInteractions
# object is present. In all cases, function calls are identical. 'subject' can

```

```
# also be missing for all functions except for 'subsetByOverlaps', as long as
# 'query' is a GInteractions or an InteractionSet.
```

### Arguments

`query`, `subject` A Vector, GInteractions or InteractionSet object, depending on the specified method. At least one of these must be a GInteractions or InteractionSet object. Also, `subject` can be missing if `query` is a GInteractions or InteractionSet object.

`maxgap`, `minoverlap`, `type`  
See [?findOverlaps](#) in the **GenomicRanges** package.

`select`, `ignore.strand`  
See [?findOverlaps](#) in the **GenomicRanges** package.

`use.region` A string specifying the regions to be used to identify overlaps.

### Value

For `findOverlaps`, a Hits object is returned if `select="all"`, and an integer vector of subject indices otherwise.

For `countOverlaps` and `overlapsAny`, an integer or logical vector is returned, respectively.

For `subsetByOverlaps`, a subsetted object of the same class as `query` is returned.

### Overview of overlaps for GInteractions

For all methods taking a Vector and an GInteractions, the Vector is assumed to represent some region on the linear genome (e.g., GRanges) or set of such regions (GRangesList). An overlap will be defined between the interval and an GInteractions interaction if either anchor region of the latter overlaps the former. This is considered to be a one-dimensional overlap, i.e., on the linear genome.

For methods between two GInteractions objects, a two-dimensional overlap will be computed between the anchor regions of the two objects. An overlap is defined if each anchor region of the first object overlaps at least one anchor region of the second object, and each anchor region of the second object overlaps at least one anchor region of the first object, i.e., there are overlapping areas in the two-dimensional interaction space. If `subject` is missing, overlaps will be computed between interactions in `query`.

### Description of overlap methods

When `select="all"`, `findOverlaps` returns a Hits object containing overlapping pairs of queries and subjects (or more specifically, their indices in the supplied objects - see [?findOverlaps](#) for more details). For other values of `select`, an integer vector is returned with one entry for each element of `query`, which specifies the index of the chosen (first, last or arbitrary) overlapping feature in `subject` for that query. Queries with no overlaps at all are assigned NA values.

For the other methods, `countOverlaps` returns an integer vector indicating the number of elements in `subject` that were overlapped by each element in `query`. `overlapsAny` returns a logical vector indicating which elements in `query` were overlapped by at least one element in `subject`. `subsetByOverlaps` returns a subsetted `query` containing only those elements overlapped by at least one element in `subject`.

### Choice of regions to define overlaps

For one-dimensional overlaps, use `region="both"` by default such that overlaps with either interacting region are considered. If `region="first"`, overlaps are only considered between the interval and the first interacting region. Similarly, if `region="second"`, only the second interaction region is used.

For two-dimensional overlaps, use `region="both"` by default such that the order of first/second interacting/target regions is ignored. This means that an overlap will be considered between, e.g., the first interacting region and the second target region. If `region="same"`, overlaps are only considered between the set of first regions for the `GInteractions` object and the first regions for the target pairs, and similarly for the second regions. If `region="reverse"`, overlaps are only considered between the set of first regions for the `GInteractions` object and the second regions for the target pairs, and vice versa. These options tend only to be useful if the order of first/second regions is informative.

### Details for InteractionSet

The behaviour of each method for `InteractionSet` objects is largely the same as that described for `GInteractions` objects. For a given `InteractionSet` object `x`, the corresponding method is called on the `GInteractions` object in the `interactions` slot of `x`. The return value is identical to that from calling the method on `interactions(x)`, except for `subsetByOverlaps` for `InteractionSet` queries (which returns a subsetted `InteractionSet` object, containing only those rows/interactions overlapping the subject).

### Author(s)

Aaron Lun

### See Also

[InteractionSet-class](#), [findOverlaps](#), [linkOverlaps](#)

### Examples

```
example(GInteractions, echo=FALSE)

# Making a larger object, for more overlaps.
Np <- 100
N <- length(regions(gi))
all.anchor1 <- sample(N, Np, replace=TRUE)
all.anchor2 <- sample(N, Np, replace=TRUE)
gi <- GInteractions(all.anchor1, all.anchor2, regions(gi))

# GRanges overlaps:
of.interest <- resize(sample(regions(gi), 2), width=1, fix="center")
findOverlaps(of.interest, gi)
findOverlaps(gi, of.interest)
findOverlaps(gi, of.interest, select="first")
overlapsAny(gi, of.interest)
overlapsAny(of.interest, gi)
countOverlaps(gi, of.interest)
countOverlaps(of.interest, gi)
subsetByOverlaps(gi, of.interest)
subsetByOverlaps(of.interest, gi)
```

```

# GRangesList overlaps:
pairing <- GRangesList(first=regions(gi)[1:3], second=regions(gi)[4:6],
  third=regions(gi)[7:10], fourth=regions(gi)[15:17])
findOverlaps(pairing, gi)
findOverlaps(gi, pairing)
findOverlaps(gi, pairing, select="last")
overlapsAny(gi, pairing)
overlapsAny(pairing, gi)
countOverlaps(gi, pairing)
countOverlaps(pairing, gi)
subsetByOverlaps(gi, pairing)
subsetByOverlaps(pairing, gi)

# GInteractions overlaps (split into two):
first.half <- gi[1:(Np/2)]
second.half <- gi[Np/2+1:(Np/2)]
findOverlaps(first.half, second.half)
findOverlaps(first.half, second.half, select="arbitrary")
overlapsAny(first.half, second.half)
countOverlaps(first.half, second.half)
subsetByOverlaps(first.half, second.half)

findOverlaps(gi)
countOverlaps(gi)
overlapsAny(gi) # trivial result

#####
# Same can be done for an InteractionSet object:

Nlibs <- 4
counts <- matrix(rpois(Nlibs*Np, lambda=10), ncol=Nlibs)
colnames(counts) <- seq_len(Nlibs)
iset <- InteractionSet(counts, gi)

findOverlaps(of.interest, iset)
findOverlaps(iset, pairing)
findOverlaps(iset[1:(Np/2),], iset[Np/2+1:(Np/2),])

# Obviously returns InteractionSet objects instead
subsetByOverlaps(of.interest, iset)
subsetByOverlaps(iset, pairing)
subsetByOverlaps(iset[1:(Np/2),], iset[Np/2+1:(Np/2),])

# Self-overlaps
findOverlaps(iset)
countOverlaps(iset)
overlapsAny(iset) # trivial result

```

---

Interaction subsetting

*Interaction subsetting and combining*

---

## Description

Methods to subset or combine InteractionSet or GInteractions objects.

**Usage**

```
## S4 method for signature 'GInteractions,ANY,ANY'
x[i, j, ..., drop=TRUE]
## S4 replacement method for signature 'GInteractions,ANY,ANY,GInteractions'
x[i, j] <- value

## S4 method for signature 'GInteractions'
rbind(..., deparse.level=1)
## S4 method for signature 'GInteractions'
c(x, ..., recursive=FALSE)

## S4 method for signature 'GInteractions'
split(x, f, drop=FALSE)

## Each of the above methods has an identical equivalent for
## InteractionSet objects (not shown for brevity).

## S4 method for signature 'InteractionSet'
subset(x, i, j)
## S4 method for signature 'InteractionSet'
cbind(..., deparse.level=1)
```

**Arguments**

<code>x</code>	A <code>GInteractions</code> or <code>InteractionSet</code> object.
<code>i, j</code>	A vector of logical or integer subscripts. For <code>InteractionSet</code> objects, these indicate the rows and columns to be subsetted for <code>i</code> and <code>j</code> , respectively. Rows correspond to pairwise interactions while columns correspond to samples. For <code>GInteractions</code> objects, <code>i</code> indicates the genomic interactions to be retained. <code>j</code> is ignored as there is no concept of samples in this class.
<code>value</code>	A <code>GInteractions</code> or <code>InteractionSet</code> object with length or number of rows equal to length of <code>i</code> (or that of <code>x</code> , if <code>i</code> is not specified). For <code>InteractionSet</code> objects, the number of columns must be equal to the length of <code>j</code> (or number of columns in <code>x</code> , if <code>j</code> is not specified).
<code>...</code>	For <code>rbind</code> and <code>c</code> , <code>...</code> contains <code>GInteractions</code> or <code>InteractionSet</code> objects to be combined row-wise, depending on the method. All objects must be of the same class. For <code>c</code> , any objects are additional to that already specified in <code>x</code> . For <code>cbind</code> , <code>...</code> contains <code>InteractionSet</code> objects to be combined column-wise. For all other methods, this argument is ignored.
<code>f</code>	A vector-like object of length equal to <code>length(x)</code> (for <code>GInteractions</code> objects) or <code>nrow(x)</code> (for <code>InteractionSet</code> objects), indicating how entries/rows of <code>x</code> are to be split into separate objects. This is usually a factor.
<code>drop</code>	A logical scalar, indicating if levels of <code>f</code> that do not occur should be dropped in <code>split</code> , <code>InteractionSet</code> -method. This is ignored by the other methods.
<code>deparse.level</code>	An integer scalar; see <code>?base::cbind</code> for a description of this argument.
<code>recursive</code>	An integer scalar, ignored.

**Value**

A subsetted or combined object of the same class as `x`.

### Details for GInteractions

Subsetting operations will return a GInteractions object containing the specified interactions. Values of the anchor1 and anchor2 slots will be appropriately subsetting in the returned object, along with any metadata in mcols. However, note that the value of regions will not be modified by subsetting.

rbind and c are synonyms and will concatenate GInteractions objects. These methods will check whether the regions slot of all supplied objects are the same, in which case the regions and anchor indices are used directly. Otherwise, a new GRanges is constructed containing the (sorted) union of all regions across the input objects. Anchor indices in each object are refactored appropriately to refer to the relevant entries in this new object.

Note that the column names in mcols must be identical across all supplied objects in rbind and c. The column names of mcols for the regions slot must also be identical across objects. If GInteractions objects of different strictness (i.e., StrictGInteractions and ReverseGInteractions) are concatenated, the returned object will be of the same class as the first supplied object.

split produces a list of GInteractions objects, where each entry of the list corresponds to a level of f. Each GInteractions object contains all rows associated with the corresponding level of f.

For short index vectors, subsetting a GInteractions object prior to calling anchors may be much faster than the reverse procedure. This is because the anchors getter will construct a GRanges(List) containing the genomic loci for all pairwise interactions. Subsetting beforehand ensures that only loci for the desired interactions are included. This avoids constructing the entire object just to subset it later.

### Details for InteractionSet

Subsetting behaves in much the same way as that for the SummarizedExperiment class. Interactions are treated as rows and will be subsetting as such. All subsetting operations will return an InteractionSet with the specified interactions (rows) or samples (columns). Again, note that the value of regions will not be modified by subsetting.

cbind will combine objects with the same interactions but different samples. Errors will occur if the interactions are not identical between objects (i.e., must have same values in the interactions slots). Additional restrictions apply on the column and assay names - see [cbind, SummarizedExperiment-method](#) for details.

rbind and c will combine objects with the same samples but different interactions. In this case, the interactions need not be identical, and will be combined using the methods described above for GInteractions objects. Again, additional restrictions apply - see [rbind, SummarizedExperiment-method](#) for details.

split produces a list of InteractionSet objects, where each entry of the list corresponds to a level of f. Each InteractionSet object contains all rows associated with the corresponding level of f.

### Author(s)

Aaron Lun

### See Also

[InteractionSet-class](#)

### Examples

```
example(GInteractions, echo=FALSE)
```

```
# Subsetting:
```

```

gi[1,]
gi[1:2,]
gi[3]
gi[3:4]

temp.gi <- gi
temp.gi[3:4] <- gi[1:2]

# Combining:
rbind(gi, gi)
c(gi, gi)

new.gi <- gi
regions(new.gi) <- resize(regions(new.gi), width=20, fix="start")
rbind(gi, new.gi)

# Splitting:
f <- sample(4, length(gi), replace=TRUE)
out <- split(gi, f)
out[[1]]

#####
# Same can be done for an InteractionSet object:

example(InteractionSet, echo=FALSE)

# Subsetting:
iset[1,]
iset[1:2,]
iset[,1]
iset[,1:2]
iset[3,3]
iset[3:4,3:4]

# Combining:
cbind(iset, iset)
rbind(iset, iset)
c(iset, iset)

new.iset <- iset
regions(new.iset) <- resize(regions(new.iset), width=20, fix="start")
rbind(iset, new.iset)

# Splitting:
out <- split(iset, f)
out[[1]]

```

---

InteractionSet class    *InteractionSet class and constructors*

---

## Description

The InteractionSet class stores information about pairwise genomic interactions, and is intended for use in data analysis from Hi-C or ChIA-PET experiments. Each row of the InteractionSet corresponds to a pairwise interaction between two loci, as defined in the GInteractions object. Each

column corresponds to a library or sample. Each InteractionSet also contains one or more assays, intended to hold experimental data about interaction frequencies for each interaction in each sample.

### Usage

```
## S4 method for signature 'ANY,GInteractions'
InteractionSet(assays, interactions, ...)

## S4 method for signature 'missing,missing'
InteractionSet(assays, interactions, ...)
```

### Arguments

assays	A numeric matrix or a list or SimpleList of matrices, containing data for each interaction.
interactions	A GInteractions object of length equal to the number of rows in assays.
...	Other arguments to be passed to <a href="#">SummarizedExperiment</a> .

### Details

The InteractionSet class inherits from the SummarizedExperiment class and has access to all of its data members and methods (see [?SummarizedExperiment-class](#) for more details). It also contains an additional interactions slot which holds a GInteractions object (or an object from any derived classes, e.g., StrictGInteractions). Each row of the InteractionSet object corresponds to a pairwise interaction between two genomic loci in interactions.

The constructor will return an InteractionSet object containing all of the specified information - for InteractionSet,missing,missing-method, an empty InteractionSet object is returned. Note that any metadata arguments will be placed in the metadata of the internal SummarizedExperiment object, *not* the metadata of the internal GInteractions object. This is consistent with the behaviour of similar classes like RangedSummarizedExperiment.

### Value

For the constructors, an InteractionSet object is returned.

### Author(s)

Aaron Lun

### See Also

[SummarizedExperiment](#), [interaction-access](#), [interaction-subset](#), [interaction-compare](#), [SummarizedExperiment-class](#)

### Examples

```
example(GInteractions, echo=FALSE)
Nlibs <- 4
counts <- matrix(rpois(Np*Nlibs, lambda=10), ncol=Nlibs)
colnames(counts) <- seq_len(Nlibs)

iset <- InteractionSet(counts, gi)
iset <- InteractionSet(counts, gi, colData=DataFrame(lib.size=1:Nlibs*1000))
iset <- InteractionSet(counts, gi, metadata=list(name="My Hi-C data"))
```

```
# Note differences in metadata storage:
metadata(iset)
metadata(interactions(iset))
```

---

Linearize interactions

*Linearize 2D interactions into 1D ranges*

---

## Description

Convert interactions in two-dimensional space to one-dimensional ranges on the linear genome.

## Usage

```
## S4 method for signature 'GInteractions,numeric'
linearize(x, ref, internal=TRUE)

## S4 method for signature 'GInteractions,GRanges'
linearize(x, ref, ..., internal=TRUE)

# Equivalent calls for InteractionSet objects.
```

## Arguments

<code>x</code>	A <code>GInteractions</code> or <code>InteractionSet</code> object
<code>ref</code>	A numeric vector or a <code>GRanges</code> object, specifying the reference region(s) to use for linearization. If numeric, the entries should be indices pointing to a genomic interval in <code>regions(x)</code> .
<code>internal</code>	A logical scalar specifying whether interactions within <code>ref</code> should be reported.
<code>...</code>	Other arguments, passed to <code>overlapsAny</code> in the <code>GRanges</code> methods.

## Details

This method identifies all interactions with at least one anchor region overlapping the specified region(s) in `ref`. When `x` is a `GInteractions` object, the method returns a `GRanges` object with one entry per identified interaction, where the coordinates are defined as the *other* anchor region, i.e., the one that does *not* overlap the reference region.

If both of the anchor regions for an interaction overlap the reference regions, the genomic interval spanned by both anchor regions is returned. This is because it is not clear which region should be defined as the "other" anchor in such circumstances. Note that this will fail if the reference regions occur across multiple chromosomes. If `internal=FALSE`, interactions with both overlapping anchor regions are removed from the output.

When `x` is an `InteractionSet` object, a `RangedSummarizedExperiment` object is returned. Each entry corresponds to an identified interaction with the non-overlapping anchor region stored in the `rowRanges`. Experimental data associated with each identified interaction is stored in the various assays.

This method effectively converts two-dimensional interaction data into one-dimensional coverage across the linear genome. It is useful when a particular genomic region is of interest - this can be

used as ref, to examine the behaviour of all other regions relative to it. For example, Hi-C data in `x` can be converted into pseudo-4C contact frequencies after linearization.

Disjoint ranges across multiple chromosomes are supported when `ref` is a `GRanges` object. However, it usually only makes sense to use contiguous ranges as a single bait region. Similarly, if `ref` is numeric, it should refer to consecutive entries in `regions(x)` to specify the bait region.

### Value

A `GRanges` when `x` is a `GInteractions` object, and a `RangedSummarizedExperiment` when `x` is an `InteractionSet` object.

### Examples

```
example(InteractionSet, echo=FALSE)

# With integers
out <- linearize(iset, 1)
linearize(iset, 10)
linearize(iset, 20)

# With ranges
linearize(iset, regions(iset)[1], type="equal")
linearize(iset, regions(iset)[10], type="equal")
linearize(iset, regions(iset)[20], type="equal")
```

---

linkOverlaps

*Link overlapping regions*

---

### Description

Identify interactions that link two sets of regions by having anchor regions overlapping one entry in each set.

### Usage

```
## S4 method for signature 'GInteractions,GRanges,GRanges'
linkOverlaps(query, subject1, subject2, ..., use.region="both")
## S4 method for signature 'GInteractions,GRanges,missing'
linkOverlaps(query, subject1, subject2, ..., use.region="both")

# The same methods apply for InteractionSet objects as 'query'.
```

### Arguments

<code>query</code>	A <code>GInteractions</code> or <code>InteractionSet</code> object.
<code>subject1</code> , <code>subject2</code>	A <code>GRanges</code> object.
<code>...</code>	Additional arguments to be passed to <a href="#">findOverlaps</a> .
<code>use.region</code>	A string specifying which query regions should be used to overlap which subject. Ignored if <code>subject2</code> is missing.

## Details

This function identifies all interactions in query where one anchor overlaps an entry in subject1 and the other anchor overlaps an entry in subject2. It is designed to be used to identify regions that are linked by interactions in query. For example, one might specify genes as subject1 and enhancers as subject2, to identify all gene-enhancer contacts present in query. This is useful when the exact pairings between subject1 and subject2 are undefined.

The function returns a dataframe specifying the index of the interaction in query; the index of the overlapped region in subject1; and the index of the overlapped region in subject2. If multiple regions in subject1 and/or subject2 are overlapping the anchor regions of a particular interaction, all combinations of two overlapping regions (one from each subject\* set) are reported for that interaction.

By default, use `.region="both"` such that overlaps will be considered between any first/second interacting region in query and either subject1 or subject2. If use `.region="same"`, overlaps will only be considered between the first interacting region in query and entries in subject1, and between the second interacting region and subject2. The opposite applies with use `.region="reverse"`, where the first and second interacting regions are overlapped with subject2 and subject1 respectively.

If subject2 is not specified, links within subject1 are identified instead, i.e., subject2 is set to subject1. In such cases, the returned dataframe is such that the first subject index is always greater than the second subject index, to avoid redundant permutations.

## Value

A dataframe of integer indices indicating which elements of query link which elements of subject1 and subject2.

## Author(s)

Aaron Lun

## See Also

[findOverlaps](#), [GInteractions](#), [Vector-method](#)

## Examples

```
example(GInteractions, echo=FALSE)

all.genes <- GRanges("chrA", IRanges(0:9*10, 1:10*10))
all.enhancers <- GRanges("chrB", IRanges(0:9*10, 1:10*10))

out <- linkOverlaps(gi, all.genes, all.enhancers)
head(out)

out <- linkOverlaps(gi, all.genes)
head(out)

# Same methods apply for InteractionSet objects.

example(InteractionSet, echo=FALSE)
out <- linkOverlaps(iset, all.genes, all.enhancers)
out <- linkOverlaps(iset, all.genes)
```

---

pairs	<i>Extract paired ranges</i>
-------	------------------------------

---

### Description

Represent interactions in a GInteractions or Interaction object as a Pairs, SelfHits or GRangesList object.

### Usage

```
## S4 method for signature 'GInteractions'  
pairs(x, id=FALSE, as.grlist=FALSE)  
  
# Equivalent call for InteractionSet to above.  
  
makeGInteractionsFromGRangesPairs(x)
```

### Arguments

x	For pairs, a GInteractions or InteractionSet object. For makeGInteractionsFromGRangesPairs, a Pairs object containing two parallel GRanges.
id	A logical scalar specifying whether indices should be returned instead of regions.
as.grlist	A logical scalar specifying whether a GRangesList should be returned.

### Details

Recall that the GInteractions object stores anchor regions for each interaction in two parallel GRanges, where corresponding entries between the two GRanges constitute the pair of regions for one interaction. These parallel ranges can be extracted and stored as a Pairs object for further manipulation. This is similar to the GRangesList reported by [anchors](#) with type="both" and id=FALSE. The reverse conversion is performed using makeGInteractionsFromGRangesPairs.

An alternative representation involves storing the two anchors for each interaction in a single GRanges of length 2. Multiple interactions are then stored as a GRangesList, along with any meta-data and sequence information. This is returned if as.grlist=FALSE, may be more useful in some applications where the two interacting regions must be in the same GRanges. Finally, if id=TRUE, the anchor indices are extracted and returned as a SelfHits object. This may be useful for graph construction.

### Value

For pairs, if id=TRUE, a SelfHits object is returned. Otherwise, if as.grlist=TRUE, a GRangesList object is returned. Otherwise, a Pairs object is returned.

For makeGInteractionsFromGRangesPairs, a GInteractions object is returned.

### Author(s)

Aaron Lun

**See Also**

[GInteractions](#), [Pairs](#), [SelfHits](#), [GRangesList](#)

**Examples**

```
example(GInteractions, echo=FALSE)
y <- pairs(gi)
y
makeGInteractionsFromGRangesPairs(y)
```

```
pairs(gi, id=TRUE)
pairs(gi, as.grlist=TRUE)
```

```
example(InteractionSet, echo=FALSE)
pairs(iset)
pairs(iset, id=TRUE)
pairs(iset, as.grlist=TRUE)
```

# Index

- [,ContactMatrix,ANY,ANY,ANY-method  
(ContactMatrix subsetting), 13
- [,ContactMatrix,ANY,ANY-method  
(ContactMatrix subsetting), 13
- [,ContactMatrix,ANY-method  
(ContactMatrix subsetting), 13
- [,GInteractions,ANY,ANY-method  
(Interaction subsetting), 34
- [,GInteractions,ANY-method  
(Interaction subsetting), 34
- [,InteractionSet,ANY,ANY,ANY-method  
(Interaction subsetting), 34
- [,InteractionSet,ANY,ANY-method  
(Interaction subsetting), 34
- [,InteractionSet,ANY-method  
(Interaction subsetting), 34
- [<-,ContactMatrix,ANY,ANY,ContactMatrix-method  
(ContactMatrix subsetting), 13
- [<-,GInteractions,ANY,ANY,GInteractions-method  
(Interaction subsetting), 34
- [<-,InteractionSet,ANY,ANY,InteractionSet-method  
(Interaction subsetting), 34
- \$,GInteractions-method (Interaction  
accessors), 21
- \$<-,GInteractions-method (Interaction  
accessors), 21
  
- anchors, 14, 36, 42
- anchors (Interaction accessors), 21
- anchors,ContactMatrix-method  
(ContactMatrix accessors), 3
- anchors,GInteractions-method  
(Interaction accessors), 21
- anchors,InteractionSet-method  
(Interaction accessors), 21
- anchors<-(Interaction accessors), 21
- anchors<-,ContactMatrix-method  
(ContactMatrix accessors), 3
- anchors<-,GInteractions-method  
(Interaction accessors), 21
- anchors<-,InteractionSet-method  
(Interaction accessors), 21
- anchors<-,ReverseStrictGInteractions-method  
(Interaction accessors), 21
  
- anchors<-,StrictGInteractions-method  
(Interaction accessors), 21
- Annotated, 5, 7
- appendRegions<- (Interaction  
accessors), 21
- appendRegions<-,ContactMatrix-method  
(ContactMatrix accessors), 3
- appendRegions<-,GInteractions-method  
(Interaction accessors), 21
- appendRegions<-,InteractionSet-method  
(Interaction accessors), 21
- as.data.frame, 23
- as.data.frame,GInteractions-method  
(Interaction accessors), 21
- as.matrix,ContactMatrix-method  
(ContactMatrix accessors), 3
- as.matrix<- (ContactMatrix accessors), 3
- as.matrix<-,ContactMatrix-method  
(ContactMatrix accessors), 3
  
- boundingBox, 2
- boundingBox,GInteractions-method  
(boundingBox), 2
- boundingBox,InteractionSet-method  
(boundingBox), 2
  
- c,GInteractions-method (Interaction  
subsetting), 34
- c,InteractionSet-method (Interaction  
subsetting), 34
- cbind, 14, 35
- cbind,ContactMatrix-method  
(ContactMatrix subsetting), 13
- cbind,InteractionSet-method  
(Interaction subsetting), 34
- ContactMatrix (ContactMatrix class), 6
- ContactMatrix accessors, 3
- ContactMatrix class, 6
- ContactMatrix distances, 8
- ContactMatrix overlaps, 10
- ContactMatrix sorting, 11
- ContactMatrix subsetting, 13
- ContactMatrix,ANY,GRanges,GRanges,GenomicRangesORmissin  
(ContactMatrix class), 6

- ContactMatrix, ANY, numeric, numeric, GRanges-method (ContactMatrix class), 6
- ContactMatrix, missing, missing, missing, GenomicRangesORmissing-method (ContactMatrix class), 6
- ContactMatrix-access (ContactMatrix accessors), 3
- ContactMatrix-class (ContactMatrix class), 6
- ContactMatrix-dist (ContactMatrix distances), 8
- ContactMatrix-sort (ContactMatrix sorting), 11
- ContactMatrix-subset (ContactMatrix subsetting), 13
- Convert classes, 15
- countOverlaps (Interaction overlaps), 31
- countOverlaps, GInteractions, GInteractions-method (Interaction overlaps), 31
- countOverlaps, GInteractions, InteractionSet-method (Interaction overlaps), 31
- countOverlaps, GInteractions, missing-method (Interaction overlaps), 31
- countOverlaps, GInteractions, Vector-method (Interaction overlaps), 31
- countOverlaps, InteractionSet, GInteractions-method (Interaction overlaps), 31
- countOverlaps, InteractionSet, InteractionSet-method (Interaction overlaps), 31
- countOverlaps, InteractionSet, missing-method (Interaction overlaps), 31
- countOverlaps, InteractionSet, Vector-method (Interaction overlaps), 31
- countOverlaps, Vector, GInteractions-method (Interaction overlaps), 31
- countOverlaps, Vector, InteractionSet-method (Interaction overlaps), 31
- countOverlaps, GInteractions, 43
- GInteractions (GInteractions class), 18
- GInteractions class, 18
- GInteractions, GRanges, GRanges, GenomicRangesORmissing-method (GInteractions class), 18
- GInteractions, missing, missing, GenomicRangesORmissing-method (GInteractions class), 18
- GInteractions, numeric, numeric, GRanges-method (GInteractions class), 18
- GInteractions-class (GInteractions class), 18
- GRanges methods, 20
- GRangesList, 43
- inflate, 7
- inflate (Convert classes), 15
- inflate, GInteractions-method (Convert classes), 15
- inflate, InteractionSet-method (Convert classes), 15
- Interaction accessors, 21
- Interaction compare, 26
- Interaction distances, 29
- Interaction overlaps, 31
- Interaction subsetting, 34
- interaction-access (Interaction accessors), 21
- duplicated, InteractionSet-method (Interaction compare), 26
- findOverlaps, 10, 11, 32, 33, 40
- findOverlaps (Interaction overlaps), 31
- findOverlaps, GInteractions, GInteractions-method (Interaction overlaps), 31
- findOverlaps, GInteractions, InteractionSet-method (Interaction overlaps), 31
- findOverlaps, GInteractions, missing-method (Interaction overlaps), 31
- findOverlaps, GInteractions, Vector-method (Interaction overlaps), 31
- findOverlaps, InteractionSet, GInteractions-method (Interaction overlaps), 31
- findOverlaps, InteractionSet, InteractionSet-method (Interaction overlaps), 31
- findOverlaps, InteractionSet, missing-method (Interaction overlaps), 31
- findOverlaps, InteractionSet, Vector-method (Interaction overlaps), 31
- findOverlaps, Vector, GInteractions-method (Interaction overlaps), 31
- findOverlaps, Vector, InteractionSet-method (Interaction overlaps), 31

- interaction-compare (Interaction compare), 26
- Interaction-overlaps (Interaction overlaps), 31
- interaction-subset (Interaction subsetting), 34
- interactions (Interaction accessors), 21
- interactions, InteractionSet-method (Interaction accessors), 21
- interactions<- (Interaction accessors), 21
- interactions<-, InteractionSet-method (Interaction accessors), 21
- InteractionSet (InteractionSet class), 37
- InteractionSet class, 37
- InteractionSet, ANY, GInteractions-method (InteractionSet class), 37
- InteractionSet, missing, missing-method (InteractionSet class), 37
- InteractionSet-class (InteractionSet class), 37
- intrachr (Interaction distances), 29
- intrachr, ContactMatrix-method (ContactMatrix distances), 8
- intrachr, GInteractions-method (Interaction distances), 29
- intrachr, InteractionSet-method (Interaction distances), 29
- length, ContactMatrix-method (ContactMatrix accessors), 3
- length, GInteractions-method (Interaction accessors), 21
- linearize (Linearize interactions), 39
- Linearize interactions, 39
- linearize, GInteractions, GRanges-method (Linearize interactions), 39
- linearize, GInteractions, numeric-method (Linearize interactions), 39
- linearize, InteractionSet, GRanges-method (Linearize interactions), 39
- linearize, InteractionSet, numeric-method (Linearize interactions), 39
- linkOverlaps, 33, 40
- linkOverlaps, GInteractions, GRanges, GRanges-method (linkOverlaps), 40
- linkOverlaps, GInteractions, GRanges, missing-method (linkOverlaps), 40
- linkOverlaps, InteractionSet, GRanges, GRanges-method (linkOverlaps), 40
- linkOverlaps, InteractionSet, GRanges, missing-method (linkOverlaps), 40
- makeGInteractionsFromGRangesPairs (pairs), 42
- match, 27, 29
- match, GInteractions, GInteractions-method (Interaction compare), 26
- match, GInteractions, InteractionSet-method (Interaction compare), 26
- match, InteractionSet, GInteractions-method (Interaction compare), 26
- match, InteractionSet, InteractionSet-method (Interaction compare), 26
- mcols, 23
- mcols, InteractionSet-method (Interaction accessors), 21
- mcols<-, InteractionSet-method (Interaction accessors), 21
- names, GInteractions-method (Interaction accessors), 21
- names, InteractionSet-method (Interaction accessors), 21
- names<-, GInteractions-method (Interaction accessors), 21
- names<-, InteractionSet-method (Interaction accessors), 21
- order, ContactMatrix-method (ContactMatrix sorting), 11
- order, GInteractions-method (Interaction compare), 26
- order, InteractionSet-method (Interaction compare), 26
- overlapsAny, 16, 39
- overlapsAny (Interaction overlaps), 31
- overlapsAny, ContactMatrix, GInteractions-method (ContactMatrix overlaps), 10
- overlapsAny, ContactMatrix, GRanges-method (ContactMatrix overlaps), 10
- overlapsAny, ContactMatrix, GRangesList-method (ContactMatrix overlaps), 10
- overlapsAny, ContactMatrix, InteractionSet-method (ContactMatrix overlaps), 10
- overlapsAny, GInteractions, GInteractions-method (Interaction overlaps), 31
- overlapsAny, GInteractions, InteractionSet-method (Interaction overlaps), 31
- overlapsAny, GInteractions, missing-method (Interaction overlaps), 31
- overlapsAny, GInteractions, Vector-method (Interaction overlaps), 31
- overlapsAny, InteractionSet, GInteractions-method (Interaction overlaps), 31

- overlapsAny, InteractionSet, InteractionSet-method (Interaction overlaps), 31
- overlapsAny, InteractionSet, missing-method (Interaction overlaps), 31
- overlapsAny, InteractionSet, Vector-method (Interaction overlaps), 31
- overlapsAny, Vector, GInteractions-method (Interaction overlaps), 31
- overlapsAny, Vector, InteractionSet-method (Interaction overlaps), 31
  
- pairdist (Interaction distances), 29
- pairdist, ContactMatrix-method (ContactMatrix distances), 8
- pairdist, GInteractions-method (Interaction distances), 29
- pairdist, InteractionSet-method (Interaction distances), 29
- Pairs, 43
- pairs, 42
- pairs, GInteractions-method (pairs), 42
- pairs, InteractionSet-method (pairs), 42
- pcompare, 29
- pcompare, GInteractions, GInteractions-method (Interaction compare), 26
  
- rbind, ContactMatrix-method (ContactMatrix subsetting), 13
- rbind, GInteractions-method (Interaction subsetting), 34
- rbind, InteractionSet-method (Interaction subsetting), 34
- reduceRegions (Interaction accessors), 21
- reduceRegions, ContactMatrix-method (ContactMatrix accessors), 3
- reduceRegions, GInteractions-method (Interaction accessors), 21
- reduceRegions, InteractionSet-method (Interaction accessors), 21
- regions (Interaction accessors), 21
- regions, ContactMatrix-method (ContactMatrix accessors), 3
- regions, GInteractions-method (Interaction accessors), 21
- regions, InteractionSet-method (Interaction accessors), 21
- regions<- (Interaction accessors), 21
- regions<-, ContactMatrix-method (ContactMatrix accessors), 3
- regions<-, GInteractions-method (Interaction accessors), 21
- regions<-, InteractionSet-method (Interaction accessors), 21
- regions<-, InteractionSet-method (Interaction accessors), 21
- replaceRegions<- (Interaction accessors), 21
- replaceRegions<-, ContactMatrix-method (ContactMatrix accessors), 3
- replaceRegions<-, GInteractions-method (Interaction accessors), 21
- replaceRegions<-, InteractionSet-method (Interaction accessors), 21
- ReverseStrictGInteractions-class (GInteractions class), 18
  
- SelfHits, 43
- Seqinfo, 24
- seqinfo, GInteractions-method (Interaction accessors), 21
- seqinfo, InteractionSet-method (Interaction accessors), 21
- seqinfo<-, GInteractions-method (Interaction accessors), 21
- seqinfo<-, InteractionSet-method (Interaction accessors), 21
- show, ContactMatrix-method (ContactMatrix accessors), 3
- show, GInteractions-method (Interaction accessors), 21
- show, InteractionSet-method (Interaction accessors), 21
- sort, ContactMatrix-method (ContactMatrix sorting), 11
- sort, GInteractions-method (Interaction compare), 26
- sort, InteractionSet-method (Interaction compare), 26
- split, GInteractions-method (Interaction subsetting), 34
- split, InteractionSet-method (Interaction subsetting), 34
- StrictGInteractions-class (GInteractions class), 18
- subset, ContactMatrix-method (ContactMatrix subsetting), 13
- subset, GInteractions-method (Interaction subsetting), 34
- subset, InteractionSet-method (Interaction subsetting), 34
- subsetByOverlaps (Interaction overlaps), 31
- subsetByOverlaps, GInteractions, GInteractions-method (Interaction overlaps), 31
- subsetByOverlaps, GInteractions, InteractionSet-method (Interaction overlaps), 31

subsetByOverlaps,GInteractions,Vector-method  
(Interaction overlaps), 31

subsetByOverlaps,InteractionSet,GInteractions-method  
(Interaction overlaps), 31

subsetByOverlaps,InteractionSet,InteractionSet-method  
(Interaction overlaps), 31

subsetByOverlaps,InteractionSet,Vector-method  
(Interaction overlaps), 31

subsetByOverlaps,Vector,GInteractions-method  
(Interaction overlaps), 31

subsetByOverlaps,Vector,InteractionSet-method  
(Interaction overlaps), 31

SummarizedExperiment, 38

swapAnchors, 2

swapAnchors (Interaction compare), 26

swapAnchors,GInteractions-method  
(Interaction compare), 26

swapAnchors,InteractionSet-method  
(Interaction compare), 26

t,ContactMatrix-method (ContactMatrix  
accessors), 3

trim, 21

trim,ContactMatrix-method (GRanges  
methods), 20

trim,GInteractions-method (GRanges  
methods), 20

trim,InteractionSet-method (GRanges  
methods), 20

unique,ContactMatrix-method  
(ContactMatrix sorting), 11

unique,GInteractions-method  
(Interaction compare), 26

unique,InteractionSet-method  
(Interaction compare), 26

width, 21

width,ContactMatrix-method (GRanges  
methods), 20

width,GInteractions-method (GRanges  
methods), 20

width,InteractionSet-method (GRanges  
methods), 20