

# Package ‘plyinteractions’

October 17, 2024

**Title** Extending tidy verbs to genomic interactions

**Description** Operate on `GInteractions` objects as tabular data using `dplyr`-like verbs. The functions and methods in `plyinteractions` provide a grammatical approach to manipulate `GInteractions`, to facilitate their integration in genomic analysis workflows.

**Version** 1.2.0

**Date** 2023-08-21

**License** Artistic-2.0

**URL** <https://github.com/js2264/plyinteractions>

**BugReports** <https://github.com/js2264/plyinteractions/issues>

**biocViews** Software, Infrastructure

**Encoding** UTF-8

**Roxygen** list(markdown = TRUE)

**Imports** InteractionSet, GenomeInfoDb, BiocGenerics, GenomicRanges, plyranges, IRanges, S4Vectors, rlang, dplyr, tibble, tidyselect, methods, utils

**Suggests** tidyverse, BSgenome.Mmusculus.UCSC.mm10, Biostrings, BiocParallel, scales, HiContactsData, rtracklayer, BiocStyle, covr, knitr, rmarkdown, sessioninfo, testthat (>= 3.0.0), RefManageR

**Config/testthat.edition** 3

**VignetteBuilder** knitr

**RoxygenNote** 7.2.3

**Collate** 'AllGenerics.R' 'AllClasses.R' 'anchor.R' 'annotate.R'  
'arrange.R' 'count-overlaps.R' 'count.R' 'data.R'  
'filter-overlaps.R' 'filter.R' 'find-overlaps.R' 'flank.R'  
'ginteractions-construct.R' 'ginteractions-env.R'  
'ginteractions-getters.R' 'ginteractions-scoping.R'  
'ginteractions-setters.R' 'tbl\_vars.R' 'group\_data.R'  
'group\_by.R' 'internals.R' 'join-overlap-left.R'  
'methods-AnchoredPinnedGInteractions.R'

```
'methods-DelegatingGInteractions.R'
'methods-GroupedGInteractions.R'
'methods-PinnedGInteractions.R' 'methods-show.R' 'mutate.R'
'pin.R' 'plyinteractions.R' 'reexports-dplyr.R'
'reexports-plyranges.R' 'reexports.R' 'rename.R'
'replace-anchors.R' 'select.R' 'shift.R' 'slice.R' 'stretch.R'
'summarize.R'

LazyData false
Depends R (>= 4.3.0)
git_url https://git.bioconductor.org/packages/plyinteractions
git_branch RELEASE_3_19
git_last_commit f23ac2b
git_last_commit_date 2024-04-30
Repository Bioconductor 3.19
Date/Publication 2024-10-16
Author Jacques Serizay [aut, cre]
Maintainer Jacques Serizay <jacquesserizay@gmail.com>
```

## Contents

plyinteractions-package . . . . .	3
anchors1 . . . . .	4
annotate . . . . .	6
as_ginteractions . . . . .	8
ce10_ARCC . . . . .	10
ce10_REs . . . . .	11
delegating-ginteractions-methods . . . . .	11
dplyr-arrange . . . . .	13
dplyr-count . . . . .	14
dplyr-filter . . . . .	15
dplyr-group_by . . . . .	16
dplyr-mutate . . . . .	18
dplyr-rename . . . . .	20
dplyr-select . . . . .	21
dplyr-slice . . . . .	22
dplyr-summarize . . . . .	23
ginteractions-anchor . . . . .	24
ginteractions-count-overlaps . . . . .	26
ginteractions-filter-overlaps . . . . .	27
ginteractions-find-overlaps . . . . .	29
ginteractions-join-overlap-left . . . . .	31
GM12878_HiCCUPS . . . . .	33
group-group_data . . . . .	34
pin . . . . .	35
plyranges-flank . . . . .	37

plyranges-shift . . . . .	39
plyranges-stretch . . . . .	41
reexports . . . . .	42
replace_anchors . . . . .	43
set_seqnames1 . . . . .	44
<b>Index</b>	<b>47</b>

---

## plyinteractions-package

*plyinteractions: a grammar of data manipulation for genomic interactions*

---

### Description

plyinteractions is a dplyr-like API to the GInteractions infrastructure in Bioconductor.

### Details

plyinteractions provides a consistent interface for importing and wrangling genomic interactions from a variety of sources. The package defines a grammar of genomic interactions manipulation through a set of verbs. These verbs can be used to construct human-readable analysis pipelines based on GInteractions.

- Group genomic interactions with `group_by`;
- Summarize grouped genomic interactions with `summarize`;
- Tally/count grouped genomic interactions with `tally` and `count`;
- Modify genomic interactions with `mutate`;
- Subset genomic interactions with `filter` using `<data-masking>` and logical expressions;
- Pick out any columns from the associated metadata with `select` using `<tidy-select>` arguments;
- Subset using indices with `slice`;
- Order genomic interactions with `arrange` using categorical/numerical variables.  
For more details on the features of plyinteractions, read the vignette: `browseVignettes(package = "plyinteractions")`

### Author(s)

**Maintainer:** Jacques Serizay <jacquesserizay@gmail.com>

### See Also

Useful links:

- <https://github.com/js2264/plyinteractions>
- Report bugs at <https://github.com/js2264/plyinteractions/issues>

---

anchors1                    *Enhanced GInteractions getters*

---

## Description

Enhanced GInteractions getters

## Usage

```
anchors1(x)

anchors2(x)

seqnames1(x)

seqnames2(x)

start1(x)

start2(x)

end1(x)

end2(x)

width1(x)

width2(x)

strand1(x)

strand2(x)

ranges1(x)

ranges2(x)

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

## S4 method for signature 'GInteractions'
anchors1(x)

## S4 method for signature 'GInteractions'
anchors2(x)

## S4 method for signature 'GInteractions'
```

```
seqnames1(x)

## S4 method for signature 'GInteractions'
seqnames2(x)

## S4 method for signature 'GInteractions'
start1(x)

## S4 method for signature 'GInteractions'
start2(x)

## S4 method for signature 'GInteractions'
end1(x)

## S4 method for signature 'GInteractions'
end2(x)

## S4 method for signature 'GInteractions'
width1(x)

## S4 method for signature 'GInteractions'
width2(x)

## S4 method for signature 'GInteractions'
strand1(x)

## S4 method for signature 'GInteractions'
strand2(x)

## S4 method for signature 'GInteractions'
ranges1(x)

## S4 method for signature 'GInteractions'
ranges2(x)
```

### Arguments

x	a GInteractions object
name	The pattern or name of a column stored in the GInteractions metadata (mcols).

### Value

One of the core GInteractions fields (e.g. seqnames1, start1, ...) or one of the metadata columns when using \$. Note that auto-completion works with \$.

### Examples

```
gi <- data.frame(
  seqnames1 = 'chr1', start1 = 1, end1 = 10,
```

```

seqnames2 = 'chr1', start2 = 2, end2 = 20
) |> as_ginteractions() |> mutate(type = 'cis')
anchors1(gi)
anchors2(gi)
seqnames1(gi)
seqnames2(gi)
start1(gi)
start2(gi)
end1(gi)
end2(gi)
width1(gi)
width2(gi)
ranges1(gi)
ranges2(gi)
strand1(gi)
strand2(gi)
gi$type

```

**annotate***Annotate both anchors of a GInteractions***Description**

For each interaction in a `GInteractions` object, `annotate` returns the pairs of annotations from the `GRanges` object it overlaps with.

**Usage**

```

annotate(x, y, by)

annotate_directed(x, y, by)

## S4 method for signature 'GInteractions,GRanges,character'
annotate(x, y, by)

## S4 method for signature 'GInteractions,GRanges,character'
annotate_directed(x, y, by)

```

**Arguments**

- x a `GInteractions` object
- y a `GRanges` object to extract annotations from
- by Column name from y to use to extract annotations

**Value**

a `GInteractions` object with two extra metadata columns named `by.1` and `by.2`.

## Examples

```
#####
# 1. Basic example
#####

gi <- read.table(text =
  chr1 11 20 - chr1 21 30 +
  chr1 21 30 + chr2 51 60 "+",
  col.names = c(
    "seqnames1", "start1", "end1", "strand1",
    "seqnames2", "start2", "end2", "strand2"
  )
) |> as_ginteractions()

gr <- GenomicRanges::GRanges(c("chr1:20-30:+", "chr2:55-65:+")) |>
  plyranges::mutate(id = 1:2)

annotate(gi, gr, by = 'id')

annotate_directed(gi, gr, by = 'id')

#####
# 2. Match loops with tiled genomic bins
#####

data(GM12878_HiCCUPS)
loops <- GM12878_HiCCUPS |>
  pin_by('first') |>
  anchor_center() |>
  mutate(width1 = 500) |>
  pin_by('second') |>
  anchor_center() |>
  mutate(width2 = 500)

genomic_bins <- GenomeInfoDb::getChromInfoFromUCSC(
  'hg19', assembled.molecules.only = TRUE, as.Seqinfo = TRUE
) |>
  GenomicRanges::tileGenome(tilewidth = 10000) |>
  unlist() |>
  plyranges::mutate(binID = seq_len(plyranges::n()))

annotate(loops, genomic_bins, by = 'binID') |>
  select(starts_with('binID'))

#####
# 3. Annotate interactions by a set of regulatory elements
#####

data(ce10_ARCC)
data(ce10_REs)
annotate(ce10_ARCC, ce10_REs, by = 'annot') |>
  count(annot.1, annot.2) |>
```

```
as.data.frame() |>
dplyr::arrange(desc(n))
```

as_ginteractions	<i>Construct a GInteractions object from a tibble, DataFrame or data.frame</i>
------------------	--

## Description

The `as_ginteractions` function looks for column names in `.data` called `seqnames{1,2}`, `start{1,2}`, `end{1,2}`, and `strand{1,2}` in order to construct a `GInteractions` object. By default other columns in `.data` are placed into the `mcols` (metadata columns) slot of the returned object.

## Usage

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

## Arguments

<code>.data</code>	A <code>data.frame()</code> , <code>DataFrame()</code> or <code>tibble()</code> to construct a <code>GInteractions</code> object from.
<code>...</code>	Optional named arguments specifying which the columns in <code>.data</code> containin the core components a <code>GInteractions</code> object.
<code>keep.extra.columns</code>	TRUE or FALSE (the default). If TRUE, the columns in df that are not used to form the genomic ranges of the returned <code>GRanges</code> object are then returned as metadata columns on the object. Otherwise, they are ignored.
<code>starts.in.df.are.0based</code>	TRUE or FALSE (the default). If TRUE, then the start positions of the genomic ranges in df are considered to be 0-based and are converted to 1-based in the returned <code>GRanges</code> object.

## Value

a `GInteractions` object.

## See Also

[InteractionSet::GInteractions\(\)](#)

## Examples

```
#####
# 1. GInteractions from bedpe files imported into a data.frame
#####

bedpe <- read.table(text =
chr1 100 200 chr1 5000 5100 bedpe_example1 30 + -
chr1 1000 5000 chr1 3000 3800 bedpe_example2 100 + -",
col.names = c(
  "chrom1", "start1", "end1",
  "chrom2", "start2", "end2", "name", "score", "strand1", "strand2"))
bedpe |>
  as_ginteractions(seqnames1 = chrom1, seqnames2 = chrom2)

#####
# 2. GInteractions from standard pairs files imported into a data.frame
#####

# Note how the pairs are 0-based and no "end" field is provided
# (the standard pairs file format does not have "end" fields)
# We can provide width1 and width2 to fix this problem.

pairs <- read.table(text =
pair1 chr1 10000 chr1 20000 + +
pair2 chr1 50000 chr1 70000 + +
pair3 chr1 60000 chr2 10000 + +
pair4 chr1 30000 chr3 40000 + -",
col.names = c(
  "pairID", "chr1", "pos1", "chr2", "pos2", "strand1", "strand2")
)
pairs |>
  as_ginteractions(
    seqnames1 = chr1, start1 = pos1, width1 = 1000,
    seqnames2 = chr2, start2 = pos2, width2 = 1000,
    starts.in.df.are.0based = TRUE
  )

#####
# 3. GInteractions from data.frame with extra fields
#####

df <- read.table(text =
chr1 100 200 chr1 5000 5100
chr1 1000 5000 chr1 3000 3800",
col.names = c("chr1", "start1", "end1", "chr2", "start2", "end2"))
df |>
  as_ginteractions(seqnames1 = chr1, seqnames2 = chr2)

df <- read.table(text =
chr1 100 200 chr1 5000 5100
chr1 1000 5000 chr1 3000 3800",
col.names = c("chr1", "start1", "end1", "chr2", "start2", "end2"))
```

```

df |>
  as_ginteractions(
    seqnames1 = chr1, seqnames2 = chr2, strand1 = '+', strand2 = '-'
  )

data.frame(type = "cis", count = 3) |>
  as_ginteractions(
    seqnames1 = 'chr1', start1 = 1, end1 = 10,
    seqnames2 = 'chr1', start2 = 40, end2 = 50
  )

#####
# 4. GInteractions from a real like pairs files
#####

pairsf <- system.file('extdata', 'pairs.gz', package = 'plyinteractions')
pairs <- read.table(pairsf, comment.char = '#', header = FALSE)
head(pairs)
pairs |>
  as_ginteractions(
    seqnames1 = V2, start1 = V3, width1 = 1, strand1 = V6,
    seqnames2 = V4, start2 = V5, width2 = 1, strand2 = V7,
    starts.in.df.are.0based = TRUE
  )

```

ce10\_ARCC

*Interactions identified in L3 C. elegans by ARC-C*

## Description

Supplemental Table 2 obtained from Genome Biology online publication.

Huang N, Seow WQ, Appert A, Dong Y, Stempor P and Ahringer J Accessible Region Conformation Capture (ARC-C) gives high-resolution insights into genome architecture and regulation. *Genome Res* 2022 Feb;32(2):357-366. PMID: 34933938

## Usage

ce10\_ARCC

## Format

An object of class GInteractions of length 14992.

## Value

A GInteractions object

## Source

<https://genome.cshlp.org/content/early/2021/12/21/gr.275669.121>

**Description**

Figure 2 - Source data 1 obtained from eLife online publication.

Jänes J, Dong Y, Schoof M, Serizay J, Appert A, Cerrato C, Woodbury C, Chen R, Gemma C, Huang N, Kissiov D, Stempor P, Steward A, Zeiser E, Sauer S and Ahringer J Chromatin accessibility dynamics across *C. elegans* development and ageing. *Elife* 2018 Oct 26;7. PMID: 30362940

**Usage**

ce10\_REs

**Format**

An object of class GRanges of length 42245.

**Value**

A GRanges object

**Source**

<https://genome.cshlp.org/content/early/2021/12/21/gr.275669.121>

---

**Description**

Methods for DelegatingGInteractions objects

**Usage**

```
## S4 method for signature 'DelegatingGInteractions'  
anchors1(x)  
  
## S4 method for signature 'DelegatingGInteractions'  
ranges1(x)  
  
## S4 method for signature 'DelegatingGInteractions'  
seqnames1(x)
```

```
## S4 method for signature 'DelegatingGInteractions'
start1(x)

## S4 method for signature 'DelegatingGInteractions'
end1(x)

## S4 method for signature 'DelegatingGInteractions'
width1(x)

## S4 method for signature 'DelegatingGInteractions'
strand1(x)

## S4 method for signature 'DelegatingGInteractions'
anchors2(x)

## S4 method for signature 'DelegatingGInteractions'
ranges2(x)

## S4 method for signature 'DelegatingGInteractions'
seqnames2(x)

## S4 method for signature 'DelegatingGInteractions'
start2(x)

## S4 method for signature 'DelegatingGInteractions'
end2(x)

## S4 method for signature 'DelegatingGInteractions'
width2(x)

## S4 method for signature 'DelegatingGInteractions'
strand2(x)

## S4 method for signature 'DelegatingGInteractions'
anchors(x)

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

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

## S4 method for signature 'DelegatingGInteractions'
mcols(x)

## S4 method for signature 'DelegatingGInteractions'
show(object)
```

**Value**

One of the core GInteractions fields (e.g. seqnames1, start1, ...)

dplyr-arrange

Arrange a GInteractions by a column

**Description**

Arrange a GInteractions by a column

**Usage**

```
## S3 method for class 'GInteractions'
arrange(.data, ...)
```

**Arguments**

- .data            a GInteractions object
- ...              Variables, or functions of variables. Use dplyr::desc() to sort a variable in descending order.

**Value**

a GInteractions object.

**Examples**

```
gi <- read.table(text =
  chr1 1 10 chr1 1 10
  chr1 2 10 chr2 1 10
  chr3 3 10 chr3 1 10
  chr4 4 10 chr4 1 10
  chr5 5 10 chr5 1 10",
  col.names = c(
    "seqnames1", "start1", "end1",
    "seqnames2", "start2", "end2")
) |>
  as_ginteractions() |>
  mutate(cis = seqnames1 == seqnames2, score = runif(5)*100, gc = runif(5))
gi

#####
# 1. Arrange GInteractions by a numerical column
#####

gi |> arrange(gc)

#####
# 2. Arrange GInteractions by a logical column
#####
```

```
#####
gi |> arrange(cis)

#####
# 3. Arrange GInteractions by a factor
#####

gi |>
  mutate(rep = factor(c("rep1", "rep2", "rep1", "rep2", "rep1"))) |>
  arrange(rep)

#####
# 4. Combine sorting variables
#####

gi |>
  mutate(rep = factor(c("rep1", "rep2", "rep1", "rep2", "rep1"))) |>
  arrange(dplyr::desc(rep), score)
```

**dplyr-count***Count or tally GInteractions per group***Description**

Count or tally GInteractions per group

**Usage**

```
## S3 method for class 'GroupedGInteractions'
tally(x, wt = NULL, sort = FALSE, name = NULL)

## S3 method for class 'GroupedGInteractions'
count(x, ..., wt = NULL, sort = FALSE, name = NULL)

## S3 method for class 'GInteractions'
count(x, ..., wt = NULL, sort = FALSE, name = NULL)
```

**Arguments**

x	A grouped GInteractions object
wt	<data-masking> Frequency weights. Can be NULL or a variable: <ul style="list-style-type: none"> <li>If NULL (the default), counts the number of rows in each group.</li> <li>If a variable, computes sum(wt) for each group.</li> </ul>
sort	If TRUE, will show the largest groups at the top.
name	The name of the new column in the output.
...	<data-masking> Variables to group by.

**Value**

a S4Vectors::DataFrame() object, with an added column with the count/tblly per group.

**Examples**

```
gi <- read.table(text = "
chr1 11 20 chr1 21 30 + +
chr1 11 20 chr1 51 55 + +
chr1 11 30 chr1 51 55 - -
chr1 11 30 chr2 51 60 - -",
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2", "strand1", "strand2")
) |>
  as_ginteractions() |>
  mutate(score = runif(4), type = c('cis', 'cis', 'cis', 'trans'))

#####
# 1. Tally groups
#####

gi

gi |> group_by(strand1) |> tally()

gi |> group_by(type) |> tally()

gi |> group_by(type) |> tally(wt = score)

#####
# 2. Count per groups
#####

gi |> count(type)

gi |> group_by(type) |> count(strand1)

gi |> group_by(type, strand1) |> count(wt = score)
```

**Description**

Subset a GInteractions with tidyverse-like filter

**Usage**

```
## S3 method for class 'GInteractions'
filter(.data, ...)
```

**Arguments**

- .data a GInteractions object
- ... Expressions that return a logical value, and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.

**Value**

a GInteractions object.

**Examples**

```
gi <- read.table(text =
chr1 1 10 chr1 1 10
chr1 2 10 chr2 1 10
chr3 3 10 chr3 1 10
chr4 4 10 chr4 1 10
chr5 5 10 chr5 1 10",
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2")
) |>
  as_ginteractions() |>
  mutate(cis = seqnames1 == seqnames2, score = runif(5)*100, gc = runif(5))
gi

#####
# 1. Filter metadata columns from GInteractions by condition
#####

gi |> filter(gc > 0.1)
gi |> filter(gc > 0.1, score > 50)
gi |> filter(cis)

#####
# 2. On-the-fly calculations
#####

gi
gi |> filter(start1 >= start2 + 3)
gi |> filter(score * gc > score * 0.5)
```

**Description**

Group GInteractions by columns

**Usage**

```
## S3 method for class 'GInteractions'
group_by(.data, ..., .add = FALSE)

## S3 method for class 'DelegatingGInteractions'
group_by(.data, ..., .add = FALSE)

## S3 method for class 'GroupedGInteractions'
ungroup(x, ...)
```

**Arguments**

- .data, x            a (Grouped)GInteractions object
- ...                Column(s) to group by.
- .add               When FALSE, the default, group\_by() will override existing groups. To add to the existing groups, use .add = TRUE.

**Value**

a GroupedGInteractions object. When a (Anchored)PinnedGInteractions object is grouped, both anchoring and pinning are dropped.

**Examples**

```
gi <- read.table(text =
  chr1 11 20 chr1 21 30
  chr1 11 20 chr1 51 55
  chr1 11 30 chr1 51 55
  chr1 11 30 chr2 51 60",
  col.names = c(
    "seqnames1", "start1", "end1",
    "seqnames2", "start2", "end2")
) |>
  as_ginteractions() |>
  mutate(type = c('cis', 'cis', 'cis', 'trans'), score = runif(4))

#####
# 1. Group by core column
#####

gi |> group_by(end1)

gi |> group_by(end1, end2) |> group_data()

#####
# 2. Group by metadata column
#####

gi |> group_by(type) |> group_data()
```

```
#####
# 3. Combine core and metadata column grouping
#####

gi |> group_by(end1, type)
gi |> group_by(end1, type) |> group_data()

#####
# 4. Create a new column and group by this new variable
#####

gi |> group_by(class = c(1, 2, 1, 2))

#####
# 5. Replace or add groups to a GroupedGInteractions
#####

ggi <- gi |> group_by(class = c(1, 2, 1, 2))
ggi |> group_data()
ggi |> group_by(type) |> group_data()
ggi |> group_by(type, .add = TRUE) |> group_data()

#####
# 6. Ungroup GInteractions
#####

ggi <- gi |> group_by(type, class = c(1, 2, 1, 2))
ggi
ungroup(ggi, type)
ungroup(ggi, class)
```

**dplyr-mutate***Mutate columns from a GInteractions object***Description**

Mutate columns from a GInteractions object

**Usage**

```
## S3 method for class 'GInteractions'
mutate(.data, ...)
```

**Arguments**

- .data            a GInteractions object
- ...              Optional named arguments specifying which the columns in .data to create/modify.

**Value**

a GInteractions object.

**Examples**

```
gi <- read.table(text = "
chr1 10 20 chr1 50 51
chr1 10 50 chr2 30 40",
col.names = c("chr1", "start1", "end1", "chr2", "start2", "end2")) |>
  as_ginteractions(seqnames1 = chr1, seqnames2 = chr2)

#####
# 1. Add metadata columns to a GInteractions object
#####

gi |>
  mutate(type = c('cis', 'trans'), score = runif(2)) |>
  mutate(type2 = type)

#####
# 2. More complex, nested or inplace changes
#####

gi |>
  mutate(type = c('cis', 'trans'), score = runif(2)) |>
  mutate(type2 = type) |>
  mutate(count = c(1, 2), score = count * 2, new_col = paste0(type2, score))

#####
# 3. Core GInteractions columns can also be modified
#####

gi |>
  mutate(start1 = 1, end1 = 10, width2 = 30, strand2 = c('-', '+'))

# Note how the core columns are modified sequentially

gi |>
  mutate(start1 = 1, end1 = 10)

gi |>
  mutate(start1 = 1, end1 = 10, width1 = 50)

#####
# 4. Evaluating core GInteractions columns
#####

gi |>
  mutate(
    score = runif(2),
    cis = seqnames1 == seqnames2,
    distance = ifelse(cis, start2 - end1, NA)
```

)

**dplyr-rename***Rename columns from a GInteractions with tidyverse-like rename***Description**

Rename columns from a GInteractions with tidyverse-like `rename`

**Usage**

```
## S3 method for class 'GInteractions'
rename(.data, ...)
```

**Arguments**

.data	a GInteractions object
...	Use <code>new_name = old_name</code> to rename selected variables.

**Value**

a GInteractions object.

**Examples**

```
gi <- read.table(text =
chr1 10 20 chr1 50 51
chr1 10 50 chr2 30 40",
col.names = c("chr1", "start1", "end1", "chr2", "start2", "end2")) |>
as_ginteractions(seqnames1 = chr1, seqnames2 = chr2) |>
mutate(type = c('cis', 'trans'), score = runif(2))

#####
# 1. Rename metadata columns to a GInteractions object
#####

gi |> rename(interaction_type = type, GC = score)
```

---

dplyr-select	<i>Select columns within GInteractions metadata columns</i>
--------------	---

---

## Description

Select columns within GInteractions metadata columns

## Usage

```
## S3 method for class 'GInteractions'  
select(.data, ..., .drop_ranges = FALSE)
```

## Arguments

- .data            a GInteractions object
- ...              Integer indicating rows to keep.
- .drop\_ranges    if TRUE, returns a DataFrame object. In this case, it enables selection of any column including core GInteractions columns.

## Value

a GInteractions object.

## Examples

```
gi <- read.table(text = "  
chr1 1 10 chr1 1 10  
chr2 1 10 chr2 1 10  
chr3 1 10 chr3 1 10  
chr4 1 10 chr4 1 10  
chr5 1 10 chr5 1 10",  
col.names = c(  
    "seqnames1", "start1", "end1",  
    "seqnames2", "start2", "end2"))  
) |>  
as_ginteractions() |>  
mutate(score = runif(5)*100, cis = TRUE, gc = runif(5))  
  
#####  
# 1. Select metadata columns from GInteractions by index  
#####  
  
gi |> select(2, 1)  
gi |> select(-3)  
  
#####  
# 2. Select metadata columns from GInteractions by name  
#####
```

```

gi |> select(gc, score)

#####
# 3. Select metadata columns from GInteractions with <tidy-select>
#####

gi |> select(contains('s'))
gi |> select(matches('^s'))

#####
# 4. Select core and metadata columns with .drop_ranges = TRUE
#####

gi |> select(matches('^s'), .drop_ranges = TRUE)

```

**dplyr-slice***Slice a GInteractions rows by their index***Description**

Slice a GInteractions rows by their index

**Usage**

```
## S3 method for class 'GInteractions'
slice(.data, ...)
```

**Arguments**

.data	a GInteractions object
...	Integer indicating rows to keep.

**Value**

a GInteractions object.

**Examples**

```

gi <- read.table(text =
chr1 1 10 chr1 1 10
chr2 1 10 chr2 1 10
chr3 1 10 chr3 1 10
chr4 1 10 chr4 1 10
chr5 1 10 chr5 1 10",
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2")
) |>
```

```

as_ginteractions()

#####
# 1. Slice a GInteractions
#####

gi |> slice(1, 2, 3)
gi |> slice(-3)
gi |> slice(1:2, 5:4)

```

**dplyr-summarize***Summarize GInteractions per group***Description**

Summarize GInteractions per group

**Usage**

```

## S3 method for class 'GroupedGInteractions'
summarise(.data, ...)

## S3 method for class 'GroupedGInteractions'
summarize(.data, ...)

```

**Arguments**

.data	a (grouped) GInteractions object
...	Name-value pairs of summary functions. The name will be the name of the variable in the result.

**Value**

a S4Vectors::[DataFrame\(\)](#) object:

- The rows come from the underlying `group_keys()`.
- The columns are a combination of the grouping keys and the summary expressions that you provide.
- GInteractions class is **not** preserved, as a call to `summarize` fundamentally creates a new data frame

**Examples**

```

gi <- read.table(text =
chr1 11 20 chr1 21 30 + +
chr1 11 20 chr1 51 55 + +
chr1 11 30 chr1 51 55 - -
chr1 11 30 chr2 51 60 - -",

```

```

col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2", "strand1", "strand2")
) |>
  as_ginteractions() |>
  mutate(score = runif(4), type = c('cis', 'cis', 'cis', 'trans'))

#####
# 1. Summarize a single column
#####

gi

gi |> group_by(type) |> summarize(m = mean(score))

gi |> group_by(strand1) |> summarize(m = mean(score))

df <- gi |>
  group_by(strand1) |>
  summarize(m = mean(score), n = table(seqnames2))
df

df$n

#####
# 2. Summarize by multiple columns
#####

gi |>
  group_by(strand1, seqnames2) |>
  summarise(m = mean(score), n = table(type))

```

ginteractions-anchor *Manage GInteractions anchors with plyranges*

## Description

Manage GInteractions anchors with plyranges

## Usage

```

## S3 method for class 'AnchoredPinnedGInteractions'
anchor(x)

## S3 method for class 'AnchoredPinnedGInteractions'
unanchor(x)

## S3 method for class 'PinnedGInteractions'
anchor_start(x)

```

```

## S3 method for class 'PinnedGInteractions'
anchor_end(x)

## S3 method for class 'PinnedGInteractions'
anchor_center(x)

## S3 method for class 'PinnedGInteractions'
anchor_3p(x)

## S3 method for class 'PinnedGInteractions'
anchor_5p(x)

## S3 method for class 'AnchoredPinnedGInteractions'
anchor_start(x)

## S3 method for class 'AnchoredPinnedGInteractions'
anchor_end(x)

## S3 method for class 'AnchoredPinnedGInteractions'
anchor_center(x)

## S3 method for class 'AnchoredPinnedGInteractions'
anchor_3p(x)

## S3 method for class 'AnchoredPinnedGInteractions'
anchor_5p(x)

```

## Arguments

x A PinnedGInteractions object

## Value

- anchor\_\* functions return an AnchoredPinnedGInteractions object.
- anchor returns a character string indicating where the pinned anchors are anchored at.
- unanchor removes the anchoring for a AnchoredPinnedGInteractions object.

## Examples

```

gi <- read.table(text =
chr1 11 20 chr1 21 30 + +
chr1 11 20 chr1 51 55 + +
chr1 11 30 chr1 51 55 - -
chr1 11 30 chr2 51 60 - --
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2", "strand1", "strand2")
) |>
  as_ginteractions() |>

```

```

  mutate(score = runif(4), type = c('cis', 'cis', 'cis', 'trans'))

#####
# 1. Anchoring pinned genomic interactions with plyranges
#####

gi |> pin_by("second") |> anchor_end()

```

**ginteractions-count-overlaps***Count overlaps between a query GInteractions and a GRanges***Description**

Count overlaps between a query GInteractions and a GRanges

**Usage**

```

## S3 method for class 'PinnedGInteractions'
count_overlaps(x, y, maxgap = -1L, minoverlap = 0L)

## S3 method for class 'GInteractions'
count_overlaps(x, y, maxgap = -1L, minoverlap = 0L)

## S3 method for class 'PinnedGInteractions'
count_overlaps_directed(x, y, maxgap = -1L, minoverlap = 0L)

## S3 method for class 'GInteractions'
count_overlaps_directed(x, y, maxgap = -1L, minoverlap = 0L)

```

**Arguments**

x	A (Pinned)GInteractions object
y	A GRanges object
maxgap, minoverlap	See <a href="#">?countOverlaps</a> in the <b>GenomicRanges</b> package for a description of these arguments

**Value**

An integer vector of same length as x.

**Pinned GInteractions**

When using `count_overlaps()` with a PinnedGInteractions object, only the pinned anchors are used to check for overlap with y. This is equivalent to specifying `use.region="both"` in `InteractionSet::findOverlaps()`.

## Examples

```

gi <- read.table(text =
  chr1 11 20 - chr1 21 30 +
  chr1 11 20 - chr1 51 55 +
  chr1 21 30 - chr1 51 55 +
  chr1 21 30 - chr2 51 60 +",
  col.names = c(
    "seqnames1", "start1", "end1", "strand1",
    "seqnames2", "start2", "end2", "strand2"
  )
) |> as_ginteractions() |> mutate(id = 1:4, type = 'gi')

gr <- GenomicRanges::GRanges(
  c("chr1:20-30:+", "chr2:55-65:-")
) |> plyranges::mutate(id = 1:2, type = 'gr')

gi
gr

#####
# 1. Count overlaps between GInteractions and a subject GRanges
#####

count_overlaps(gi, gr)

count_overlaps_directed(gi, gr)

#####
# 2. Count overlaps between PinnedGInteractions and a subject GRanges
#####

gi |> pin_by("first") |> count_overlaps(gr)

gi |> pin_by("second") |> count_overlaps(gr)

gi |> pin_by("first") |> count_overlaps_directed(gr)

gi |> pin_by("second") |> count_overlaps_directed(gr)

```

## Description

Filter GInteractions overlapping with a GRanges

## Usage

```
## S3 method for class 'PinnedGInteractions'
filter_by_overlaps(x, y, maxgap = -1L, minoverlap = 0L)

## S3 method for class 'GInteractions'
filter_by_overlaps(x, y, maxgap = -1L, minoverlap = 0L)

## S3 method for class 'PinnedGInteractions'
filter_by_non_overlaps(x, y, maxgap = -1L, minoverlap = 0L)

## S3 method for class 'GInteractions'
filter_by_non_overlaps(x, y, maxgap = -1L, minoverlap = 0L)
```

## Arguments

x A (Pinned)GInteractions object  
y A GRanges object  
maxgap, minoverlap  
See [?countOverlaps](#) in the **GenomicRanges** package for a description of these arguments

## Value

An integer vector of same length as x.

## Pinned GInteractions

When using `filter_by_overlaps()` with a PinnedGInteractions object, only the pinned anchors are used to check for overlap with y. This is equivalent to specifying `use.region="both"` in `InteractionSet::findOverlaps()`.

## Examples

```
gi <- read.table(text =
  chr1 11 20 - chr1 21 30 +
  chr1 11 20 - chr1 51 55 +
  chr1 21 30 - chr1 51 55 +
  chr1 21 30 - chr2 51 60 "+",
  col.names = c(
    "seqnames1", "start1", "end1", "strand1",
    "seqnames2", "start2", "end2", "strand2")
) |> as_ginteractions() |> mutate(id = 1:4, type = 'gi')

gr <- GenomicRanges::GRanges(
  c("chr1:20-30:+", "chr2:55-65:-")
) |> plyranges::mutate(id = 1:2, type = 'gr')

gi
gr
```

```
#####
# 1. Filter GInteractions overlapping with a subject GRanges
#####

filter_by_overlaps(gi, gr)

filter_by_non_overlaps(gi, gr)

#####
# 2. Filter PinnedGInteractions overlapping with a subject GRanges
#####

gi |> pin_by("first") |> filter_by_overlaps(gr)

gi |> pin_by("first") |> filter_by_non_overlaps(gr)

gi |> pin_by("second") |> filter_by_overlaps(gr)

gi |> pin_by("second") |> filter_by_non_overlaps(gr)
```

**ginteractions-find-overlaps***Find overlaps between a query GInteractions and a GRanges***Description**

Find overlaps between a query GInteractions and a GRanges

**Usage**

```
## S3 method for class 'PinnedGInteractions'
find_overlaps(x, y, maxgap = -1L, minoverlap = 0L, suffix = c(".x", ".y"))

## S3 method for class 'GInteractions'
find_overlaps(x, y, maxgap = -1L, minoverlap = 0L, suffix = c(".x", ".y"))

## S3 method for class 'PinnedGInteractions'
find_overlaps_directed(
  x,
  y,
  maxgap = -1L,
  minoverlap = 0L,
  suffix = c(".x", ".y")
)

## S3 method for class 'GInteractions'
find_overlaps_directed(
```

```

  x,
  y,
  maxgap = -1L,
  minoverlap = 0L,
  suffix = c(".x", ".y")
)

```

## Arguments

x	A (Pinned)GInteractions object
y	GRanges object
maxgap, minoverlap	See <a href="#">?findOverlaps</a> in the <b>GenomicRanges</b> package for a description of these arguments
suffix	Suffix to add to metadata columns (character vector of length 2, default to c(".x", ".y")).

## Value

a GInteractions object with rows corresponding to the GInteractions in x that overlap y.

## Rationale

`find_overlaps()` will search for any overlap between GInteractions in x and GRanges in y. It will return a GInteractions object of length equal to the number of times x overlaps y. This GInteractions will have additional metadata columns corresponding to the metadata from y. `find_overlaps_directed()` takes the strandness of each object into account.

## Pinned GInteractions

When using `find_overlaps()` with a PinnedGInteractions object, only the pinned anchors are used to check for overlap with y. This is equivalent to specifying `use.region="both"` in `InteractionSet::findOverlaps()`.

## Examples

```

gi <- read.table(text =
  chr1 11 20 - chr1 21 30 +
  chr1 11 20 - chr1 51 55 +
  chr1 21 30 - chr1 51 55 +
  chr1 21 30 - chr2 51 60 "+",
  col.names = c(
    "seqnames1", "start1", "end1", "strand1",
    "seqnames2", "start2", "end2", "strand2"
  )
) |> as_ginteractions() |> mutate(id = 1:4, type = 'gi')

gr <- GenomicRanges::GRanges(
  c("chr1:20-30:+", "chr2:55-65:-")
) |> plyranges::mutate(id = 1:2, type = 'gr')

```

```

gi

gr

#####
# 1. Find overlaps between GInteractions and a subject GRanges
#####

find_overlaps(gi, gr)

find_overlaps_directed(gi, gr)

#####
# 2. Find overlaps between PinnedGInteractions and a subject GRanges
#####

gi |> pin_by("first") |> find_overlaps(gr)

gi |> pin_by("second") |> find_overlaps(gr)

gi |> pin_by("first") |> find_overlaps_directed(gr)

gi |> pin_by("second") |> find_overlaps_directed(gr)

```

**ginteractions-join-overlap-left***Join overlaps between a query GInteractions and a GRanges***Description**

Join overlaps between a query GInteractions and a GRanges

**Usage**

```

## S3 method for class 'PinnedGInteractions'
join_overlap_left(x, y, maxgap = -1L, minoverlap = 0L, suffix = c(".x", ".y"))

## S3 method for class 'GInteractions'
join_overlap_left(x, y, maxgap = -1L, minoverlap = 0L, suffix = c(".x", ".y"))

## S3 method for class 'PinnedGInteractions'
join_overlap_left_directed(
  x,
  y,
  maxgap = -1L,
  minoverlap = 0L,
  suffix = c(".x", ".y")
)

```

```
## S3 method for class 'GInteractions'
join_overlap_left_directed(
  x,
  y,
  maxgap = -1L,
  minoverlap = 0L,
  suffix = c(".x", ".y")
)
```

## Arguments

x	A (Pinned)GInteractions object
y	A GRanges object
maxgap, minoverlap	See <a href="#">?countOverlaps</a> in the <b>GenomicRanges</b> package for a description of these arguments
suffix	Suffix to add to metadata columns (character vector of length 2, default to c(".x", ".y")).

## Value

An integer vector of same length as x.

## Examples

```
gi <- read.table(text =
  chr1 11 20 - chr1 21 30 +
  chr1 11 20 - chr1 51 55 +
  chr1 21 30 - chr1 51 55 +
  chr1 21 30 - chr2 51 60 +",
  col.names = c(
    "seqnames1", "start1", "end1", "strand1",
    "seqnames2", "start2", "end2", "strand2")
) |> as_ginteractions() |> mutate(id = 1:4, type = 'gi')

gr <- GenomicRanges::GRanges(
  c("chr1:20-30:+", "chr2:55-65:-")
) |> plyranges::mutate(id = 1:2, type = 'gr')

gi

gr

#####
# 1. Join overlaps between GInteractions and a subject GRanges
#####

join_overlap_left(gi, gr)
```

```
join_overlap_left_directed(gi, gr)

#####
# 2. Join overlaps between PinnedGInteractions and a subject GRanges
#####

gi |> pin_by("first") |> join_overlap_left(gr)

gi |> pin_by("first") |> join_overlap_left_directed(gr)

gi |> pin_by("second") |> join_overlap_left(gr)

gi |> pin_by("second") |> join_overlap_left_directed(gr)
```

---

GM12878\_HiCCUPS

*Loops identified in GM12878 with HiCCUPS*

---

## Description

File obtained from GEO entry GSE63525 (GSE63525\_GM12878\_primary+replicate\_HiCCUPS\_looplist.txt.gz).

Rao SS, Huntley MH, Durand NC, Stamenova EK et al. A 3D map of the human genome at kilobase resolution reveals principles of chromatin looping. Cell 2014 Dec 18;159(7):1665-80.  
PMID: 25497547

## Usage

GM12878\_HiCCUPS

## Format

An object of class GInteractions of length 9448.

## Value

A GInteractions object

## Source

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE63525>

`group-group_data`      *GInteractions grouping metadata*

## Description

`GInteractions` grouping metadata

## Usage

```
## S3 method for class 'GroupedGInteractions'
group_data(.data)

## S3 method for class 'GroupedGInteractions'
group_keys(.tbl, ...)

## S3 method for class 'GroupedGInteractions'
group_indices(.data, ...)

## S3 method for class 'GInteractions'
group_vars(x)

## S3 method for class 'GroupedGInteractions'
group_vars(x)

## S3 method for class 'GroupedGInteractions'
groups(x)

## S3 method for class 'GroupedGInteractions'
group_size(x)

## S3 method for class 'GroupedGInteractions'
n_groups(x)
```

## Arguments

.data, .tbl, x	a <code>GInteractions</code> object
...	Ignored.

## Value

a `GInteractions` object.

## Examples

```
gi <- read.table(text =
  chr1 11 20 chr1 21 30
  chr1 11 20 chr1 51 55
```

```
chr1 11 30 chr1 51 55
chr1 11 30 chr2 51 60",
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2")
) |>
  as_ginteractions() |>
  mutate(type = c('cis', 'cis', 'cis', 'trans'), score = runif(4))

ggi <- gi |> group_by(end1)
ggi
group_data(ggi)
group_keys(ggi)
group_rows(ggi)
group_indices(ggi)
group_vars(ggi)
groups(ggi)
group_size(ggi)
n_groups(ggi)
```

---

pin

*Pin GInteractions by anchors set (anchors1 or anchors2).*

---

## Description

Pin GInteractions by anchors set (anchors1 or anchors2).

## Usage

```
pin(x, anchors)

pin_by(x, anchors)

pinned_anchors(x)

unpin(x)

## S4 method for signature 'GroupedGInteractions,character'
pin(x, anchors)

## S4 method for signature 'GroupedGInteractions,numeric'
pin(x, anchors)

## S4 method for signature 'GInteractions,character'
pin(x, anchors)

## S4 method for signature 'GInteractions,numeric'
pin(x, anchors)
```

```

## S4 method for signature 'PinnedGInteractions,missing'
pin(x, anchors)

## S4 method for signature 'PinnedGInteractions,character'
pin(x, anchors)

## S4 method for signature 'PinnedGInteractions,numERIC'
pin(x, anchors)

## S4 method for signature 'AnchoredPinnedGInteractions,character'
pin(x, anchors)

## S4 method for signature 'AnchoredPinnedGInteractions,numERIC'
pin(x, anchors)

pin_first(x)

pin_second(x)

pin_anchors1(x)

pin_anchors2(x)

## S4 method for signature 'AnchoredPinnedGInteractions'
unpin(x)

## S4 method for signature 'PinnedGInteractions'
unpin(x)

## S4 method for signature 'GInteractions'
unpin(x)

## S4 method for signature 'PinnedGInteractions'
pinned_anchors(x)

## S4 method for signature 'AnchoredPinnedGInteractions'
pinned_anchors(x)

```

## Arguments

x	a GInteractions object
anchors	Anchors to pin on ("first" or "second")

## Value

- `pin_*` functions return a PinnedGInteractions object.
- `pin` returns a numerical value indicating which set of anchors is pinned.

- `unpin` removes the pinning of a `PinnedGInteractions` object.
- `pinned_anchors` returns an `(Anchored)GenomicRanges` object corresponding to the pinned anchors of a `PinnedGInteractions` object.

## Examples

```
gi <- read.table(text = "
chr1 11 20 chr1 21 30
chr1 11 20 chr1 51 55
chr1 11 30 chr1 51 55
chr1 11 30 chr2 51 60",
col.names = c(
    "seqnames1", "start1", "end1",
    "seqnames2", "start2", "end2")
) |>
  as_ginteractions() |>
  mutate(type = c('cis', 'cis', 'cis', 'trans'), score = runif(4))

#####
# 1. Pin by first anchors
#####

gi |> pin_by("first")

gi |> pin_first()

gi |> pin_anchors1()

#####
# 2. Pin by second anchors
#####

gi |> pin_by("second")

gi |> pin_second()

gi |> pin_anchors2()

#####
# 3. Unpin
#####

gi |> pin("second") |> unpin()
```

plyranges-flank

*Generate flanking regions from pinned anchors of a GInteractions object with plyranges*

## Description

Generate flanking regions from pinned anchors of a `GInteractions` object with `plyranges`

**Usage**

```

flank_downstream(x, width)

## S3 method for class 'Ranges'
flank_downstream(x, width)

## S3 method for class 'PinnedGInteractions'
flank_downstream(x, width)

flank_upstream(x, width)

## S3 method for class 'Ranges'
flank_upstream(x, width)

## S3 method for class 'PinnedGInteractions'
flank_upstream(x, width)

flank_right(x, width)

## S3 method for class 'Ranges'
flank_right(x, width)

## S3 method for class 'PinnedGInteractions'
flank_right(x, width)

flank_left(x, width)

## S3 method for class 'Ranges'
flank_left(x, width)

## S3 method for class 'PinnedGInteractions'
flank_left(x, width)

```

**Arguments**

- x** a PinnedGInteractions object
- width** The width of the flanking region relative to the ranges in x. Either an integer vector of length 1 or an integer vector the same length as x. The width can be negative in which case the flanking region is reversed.

**Value**

A PinnedGInteractions object

**Examples**

```

gi <- read.table(text =
chr1 11 20 chr1 21 30 + +

```

```

chr1 11 20 chr1 51 55 + +
chr1 11 30 chr1 51 55 - -
chr1 11 30 chr2 51 60 - -
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2", "strand1", "strand2")
) |>
  as_ginteractions() |>
  mutate(score = runif(4), type = c('cis', 'cis', 'cis', 'trans'))

#####
# 1. Simple flanking
#####

gi

gi |> pin_by("first") |> flank_left(-2)

gi |> pin_by("second") |> flank_upstream(4)

#####
# 2. Chained flanking of each set of anchors
#####

gi |>
  pin_by("first") |> flank_left(2) |>
  pin_by("second") |> flank_right(2)

```

**plyranges-shift***Shift pinned anchors of a GInteractions object with plyranges***Description**

Shift pinned anchors of a GInteractions object with plyranges

**Usage**

```

shift_downstream(x, shift)

## S3 method for class 'Ranges'
shift_downstream(x, shift)

## S3 method for class 'PinnedGInteractions'
shift_downstream(x, shift)

shift_upstream(x, shift)

## S3 method for class 'Ranges'
shift_upstream(x, shift)

```

```

## S3 method for class 'PinnedGInteractions'
shift_upstream(x, shift)

shift_right(x, shift)

## S3 method for class 'Ranges'
shift_right(x, shift)

## S3 method for class 'PinnedGInteractions'
shift_right(x, shift)

shift_left(x, shift)

## S3 method for class 'Ranges'
shift_left(x, shift)

## S3 method for class 'PinnedGInteractions'
shift_left(x, shift)

```

### Arguments

- x            a PinnedGInteractions object  
 shift        The amount to move the genomic interval in the Ranges object by. Either a non-negative integer vector of length 1 or an integer vector the same length as x.

### Value

A PinnedGInteractions object

### Examples

```

gi <- read.table(text = "
chr1 11 20 chr1 21 30 + +
chr1 11 20 chr1 51 55 + +
chr1 11 30 chr1 51 55 - -
chr1 11 30 chr2 51 60 - -",
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2", "strand1", "strand2")
) |>
  as_ginteractions() |>
  mutate(score = runif(4), type = c('cis', 'cis', 'cis', 'trans'))

#####
# 1. Simple shifting
#####

gi

```

```

gi |> pin_by("first") |> shift_left(15)

gi |> pin_by("second") |> shift_downstream(10)

#####
# 2. Chained shifting of each set of anchors
#####

gi |>
  pin_by("first") |> shift_downstream(20) |>
  pin_by("second") |> shift_upstream(20)

```

**plyranges-stretch***Stretch pinned anchors of a GInteractions object with plyranges***Description**

Stretch pinned anchors of a GInteractions object with plyranges

**Usage**

```

## S3 method for class 'AnchoredPinnedGInteractions'
stretch(x, extend)

## S3 method for class 'PinnedGInteractions'
stretch(x, extend)

```

**Arguments**

<code>x</code>	a PinnedGInteractions object
<code>extend</code>	The amount to alter the width of a Ranges object by. Either an integer vector of length 1 or an integer vector the same length as <code>x</code> .

**Value**

A PinnedGInteractions object

**Examples**

```

gi <- read.table(text =
  chr1 11 20 chr1 21 30 + +
  chr1 11 20 chr1 51 55 + +
  chr1 11 30 chr1 51 55 - -
  chr1 11 30 chr2 51 60 - -",
  col.names = c(
    "seqnames1", "start1", "end1",
    "seqnames2", "start2", "end2", "strand1", "strand2")
) |>
  as_ginteractions() |>

```

```

  mutate(score = runif(4), type = c('cis', 'cis', 'cis', 'trans'))

#####
# 1. Simple stretching
#####

gi

gi |> pin_by("first") |> anchor_start() |> stretch(15)

gi |> pin_by("second") |> anchor_center() |> stretch(10)

gi |> pin_by("second") |> anchor_3p() |> stretch(20)

#####
# 2. Chained stretching of each set of anchors
#####

gi |>
  pin_by("first") |> anchor_start() |> stretch(20) |>
  pin_by("second") |> stretch(20)

```

## reexports

*Objects exported from other packages***Description**

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

**dplyr** [arrange](#), [count](#), [filter](#), [group\\_by](#), [group\\_data](#), [group\\_indices](#), [group\\_keys](#), [group\\_rows](#), [group\\_size](#), [group\\_vars](#), [groups](#), [mutate](#), [n\\_groups](#), [rename](#), [select](#), [slice](#), [summarise](#), [summarize](#), [tally](#), [ungroup](#)  
**plyranges** [anchor](#), [anchor\\_3p](#), [anchor\\_5p](#), [anchor\\_center](#), [anchor\\_end](#), [anchor\\_start](#), [count\\_overlaps](#), [count\\_overlaps\\_directed](#), [filter\\_by\\_non\\_overlaps](#), [filter\\_by\\_overlaps](#), [find\\_overlaps](#), [find\\_overlaps\\_directed](#), [join\\_overlap\\_left](#), [join\\_overlap\\_left\\_directed](#), [stretch](#), [unanchor](#)

**Value**

Depending on the re-exported function

**Examples**

```
1 + 1
```

---

`replace_anchors`

*Replace anchors of a GInteractions*

---

## Description

Replace anchors of a GInteractions

## Usage

```
replace_anchors(x, id, value)

## S4 method for signature 'GInteractions,character,GenomicRanges'
replace_anchors(x, id, value)

## S4 method for signature 'GInteractions,numeric,GenomicRanges'
replace_anchors(x, id, value)

## S4 method for signature 'PinnedGInteractions,missing,GenomicRanges'
replace_anchors(x, id, value)

## S4 method for signature 'AnchoredPinnedGInteractions,missing,GRanges'
replace_anchors(x, id, value)

## S4 method for signature 'AnchoredPinnedGInteractions,numeric,GRanges'
replace_anchors(x, id, value)
```

## Arguments

x	a (Pinned)GInteractions object
id	Which anchors to replace ("first" or "second"). Ignored if the GInteractions is already pinned to a specific set of anchors.
value	A GRanges object vector the same length as x.

## Value

a (Pinned)GInteractions object.

## Examples

```
gi <- read.table(text = "
chr1 11 20 chr1 21 30
chr1 11 20 chr1 51 55
chr1 11 30 chr1 51 55
chr1 11 30 chr2 51 60",
col.names = c(
  "seqnames1", "start1", "end1",
  "seqnames2", "start2", "end2")
```

```
) |>
  as_ginteractions() |>
  mutate(type = c('cis', 'cis', 'cis', 'trans'), score = runif(4))

#####
# 1. Replace anchors of a GInteractions object
#####

gi |> replace_anchors(2, value = anchors1(gi))

gi |> replace_anchors(1, value = anchors2(gi))

gi |> replace_anchors(1, value = GenomicRanges::GRanges(c(
  "chr1:1-2", "chr1:2-3", "chr1:3-4", "chr1:4-5"
)))

#####
# 2. Replace anchors of a pinned GInteractions object
#####

gi |> pin_by(1) |> replace_anchors(value = anchors1(gi))

gi |> replace_anchors(1, value = anchors2(gi))

gi |>
  pin_by(1) |>
  replace_anchors(value = GenomicRanges::GRanges(c(
    "chr1:1-2", "chr1:2-3", "chr1:3-4", "chr1:4-5"
  ))) |>
  pin_by(2) |>
  replace_anchors(value = GenomicRanges::GRanges(c(
    "chr2:1-2", "chr2:2-3", "chr2:3-4", "chr2:4-5"
  )))
```

`set_seqnames1`      *Internal GInteractions setters*

## Description

Internal GInteractions setters

## Usage

```
set_seqnames1(x, value)

set_seqnames2(x, value)

set_start1(x, value)
```

```
set_start2(x, value)

set_end1(x, value)

set_end2(x, value)

set_width1(x, value)

set_width2(x, value)

set_strand1(x, value)

set_strand2(x, value)

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

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

## S4 method for signature 'GInteractions,factor'
set_seqnames1(x, value)

## S4 method for signature 'GInteractions,factor'
set_seqnames2(x, value)

## S4 method for signature 'GInteractions,numeric'
set_start1(x, value)

## S4 method for signature 'GInteractions,numeric'
set_start2(x, value)

## S4 method for signature 'GInteractions,numeric'
set_end1(x, value)

## S4 method for signature 'GInteractions,numeric'
set_end2(x, value)

## S4 method for signature 'GInteractions,numeric'
set_width1(x, value)

## S4 method for signature 'GInteractions,numeric'
set_width2(x, value)

## S4 method for signature 'AnchoredPinnedGInteractions,numeric'
set_width1(x, value)

## S4 method for signature 'AnchoredPinnedGInteractions,numeric'
```

```
set_width2(x, value)

## S4 method for signature 'GInteractions,character'
set_strand1(x, value)

## S4 method for signature 'GInteractions,character'
set_strand2(x, value)
```

**Arguments**

x	a GInteractions object
value	a value passed to the corresponding field

**Value**

A modified GInteractions

# Index

\* **internal**  
  ce10\_ARCC, 10  
  ce10\_REs, 11  
  delegating-ginteractions-methods,  
    11  
  GM12878\_HiCCUPS, 33  
  reexports, 42  
  set\_seqnames1, 44  
\$,GInteractions-method (anchors1), 4  
  
anchor, 42  
anchor (reexports), 42  
anchor.AnchoredPinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_3p, 42  
anchor\_3p(reexports), 42  
anchor\_3p.AnchoredPinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_3p.PinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_5p, 42  
anchor\_5p(reexports), 42  
anchor\_5p.AnchoredPinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_5p.PinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_center, 42  
anchor\_center(reexports), 42  
anchor\_center.AnchoredPinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_center.PinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_end, 42  
anchor\_end(reexports), 42  
anchor\_end.AnchoredPinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_end.PinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_start, 42  
anchor\_start(reexports), 42  
  
anchor\_start.AnchoredPinnedGInteractions  
  (ginteractions-anchor), 24  
anchor\_start.PinnedGInteractions  
  (ginteractions-anchor), 24  
anchors,DelegatingGInteractions-method  
  (delegating-ginteractions-methods),  
    11  
anchors1, 4  
anchors1,DelegatingGInteractions-method  
  (delegating-ginteractions-methods),  
    11  
anchors1,GInteractions-method  
  (anchors1), 4  
anchors2(anchors1), 4  
anchors2,DelegatingGInteractions-method  
  (delegating-ginteractions-methods),  
    11  
anchors2,GInteractions-method  
  (anchors1), 4  
annotate, 6  
annotate,GInteractions,GRanges,character-method  
  (annotation), 6  
annotate\_directed(annotation), 6  
annotate\_directed,GInteractions,GRanges,character-method  
  (annotation), 6  
arrange, 3, 42  
arrange(reexports), 42  
arrange.GInteractions(dplyr-arrange),  
  13  
as\_ginteractions, 8  
  
ce10\_ARCC, 10  
ce10\_REs, 11  
count, 3, 42  
count(reexports), 42  
count.GInteractions(dplyr-count), 14  
count.GroupedGInteractions  
  (dplyr-count), 14  
count\_overlaps, 42  
count\_overlaps(reexports), 42

count\_overlaps.GInteractions  
     (ginteractions-count-overlaps),  
     26  
 count\_overlaps.PinnedGInteractions  
     (ginteractions-count-overlaps),  
     26  
 count\_overlaps\_directed, 42  
 count\_overlaps\_directed(reexports), 42  
 count\_overlaps\_directed.GInteractions  
     (ginteractions-count-overlaps),  
     26  
 count\_overlaps\_directed.PinnedGInteractions  
     (ginteractions-count-overlaps),  
     26  
 countOverlaps, 26, 28, 32

data.frame(), 8  
 DataFrame(), 8, 15, 23  
 delegating-ginteractions-methods, 11  
 dplyr-arrange, 13  
 dplyr-count, 14  
 dplyr-filter, 15  
 dplyr-group\_by, 16  
 dplyr-mutate, 18  
 dplyr-rename, 20  
 dplyr-select, 21  
 dplyr-slice, 22  
 dplyr-summarise (dplyr-summarize), 23  
 dplyr-summarize, 23

end1 (anchors1), 4  
 end1, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 end1, GInteractions-method (anchors1), 4  
 end2 (anchors1), 4  
 end2, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 end2, GInteractions-method (anchors1), 4

filter, 3, 42  
 filter (reexports), 42  
 filter.GInteractions (dplyr-filter), 15  
 filter\_by\_non\_overlaps, 42  
 filter\_by\_non\_overlaps (reexports), 42  
 filter\_by\_non\_overlaps.GInteractions  
     (ginteractions-filter-overlaps),  
     27

filter\_by\_non\_overlaps.PinnedGInteractions  
     (ginteractions-filter-overlaps),  
     27  
 filter\_by\_overlaps, 42  
 filter\_by\_overlaps (reexports), 42  
 filter\_by\_overlaps.GInteractions  
     (ginteractions-filter-overlaps),  
     27  
 filter\_by\_overlaps.PinnedGInteractions  
     (ginteractions-filter-overlaps),  
     27  
 find\_overlaps, 42  
 find\_overlaps (reexports), 42  
 find\_overlaps.GInteractions  
     (ginteractions-find-overlaps),  
     29  
 find\_overlaps.PinnedGInteractions  
     (ginteractions-find-overlaps),  
     29  
 find\_overlaps\_directed, 42  
 find\_overlaps\_directed (reexports), 42  
 find\_overlaps\_directed.GInteractions  
     (ginteractions-find-overlaps),  
     29  
 find\_overlaps\_directed.PinnedGInteractions  
     (ginteractions-find-overlaps),  
     29  
 findOverlaps, 30  
 first<-, GInteractions-method  
     (set\_seqnames1), 44  
 flank\_downstream (plyranges-flank), 37  
 flank\_left (plyranges-flank), 37  
 flank\_right (plyranges-flank), 37  
 flank\_upstream (plyranges-flank), 37

ginteractions-anchor, 24  
 ginteractions-annotate (annotate), 6  
 ginteractions-count-overlaps, 26  
 ginteractions-filter-overlaps, 27  
 ginteractions-find-overlaps, 29  
 ginteractions-getters (anchors1), 4  
 ginteractions-join-overlap-left, 31  
 ginteractions-pin (pin), 35  
 ginteractions-setters (set\_seqnames1),  
     44  
 GM12878\_HiCCUPS, 33  
 group-group\_data, 34  
 group\_by, 3, 42  
 group\_by (reexports), 42

group\_by.DelegatingGInteractions  
  (dplyr-group\_by), 16  
group\_by.GInteractions  
  (dplyr-group\_by), 16  
group\_data, 42  
group\_data (reexports), 42  
group\_data.GroupedGInteractions  
  (group-group\_data), 34  
group\_indices, 42  
group\_indices (reexports), 42  
group\_indices.GroupedGInteractions  
  (group-group\_data), 34  
group\_keys, 42  
group\_keys (reexports), 42  
group\_keys.GroupedGInteractions  
  (group-group\_data), 34  
group\_rows, 42  
group\_rows (reexports), 42  
group\_size, 42  
group\_size (reexports), 42  
group\_size.GroupedGInteractions  
  (group-group\_data), 34  
group\_vars, 42  
group\_vars (reexports), 42  
group\_vars.GInteractions  
  (group-group\_data), 34  
group\_vars.GroupedGInteractions  
  (group-group\_data), 34  
groups, 42  
groups (reexports), 42  
groups.GroupedGInteractions  
  (group-group\_data), 34

InteractionSet::GInteractions(), 8

join\_overlap\_left, 42  
join\_overlap\_left (reexports), 42  
join\_overlap\_left.GInteractions  
  (ginteractions-join-overlap-left),  
  31  
join\_overlap\_left.PinnedGInteractions  
  (ginteractions-join-overlap-left),  
  31  
join\_overlap\_left\_directed, 42  
join\_overlap\_left\_directed (reexports),  
  42  
join\_overlap\_left\_directed.GInteractions  
  (ginteractions-join-overlap-left),  
  31

join\_overlap\_left\_directed.PinnedGInteractions  
  (ginteractions-join-overlap-left),  
  31

mcols,DelegatingGInteractions-method  
  (delegating-ginteractions-methods),  
  11  
mutate, 3, 42  
mutate (reexports), 42  
mutate.GInteractions (dplyr-mutate), 18

n\_groups, 42  
n\_groups (reexports), 42  
n\_groups.GroupedGInteractions  
  (group-group\_data), 34

pin, 35  
pin,AnchoredPinnedGInteractions,character-method  
  (pin), 35  
pin,AnchoredPinnedGInteractions,numeric-method  
  (pin), 35  
pin,GInteractions,character-method  
  (pin), 35  
pin,GInteractions,numeric-method (pin),  
  35  
pin,GroupedGInteractions,character-method  
  (pin), 35  
pin,GroupedGInteractions,numeric-method  
  (pin), 35  
pin,PinnedGInteractions,character-method  
  (pin), 35  
pin,PinnedGInteractions,missing-method  
  (pin), 35  
pin,PinnedGInteractions,numeric-method  
  (pin), 35  
pin\_anchors1 (pin), 35  
pin\_anchors2 (pin), 35  
pin\_by (pin), 35  
pin\_first (pin), 35  
pin\_second (pin), 35  
pinned\_anchors (pin), 35  
pinned\_anchors,AnchoredPinnedGInteractions-method  
  (pin), 35  
pinned\_anchors,PinnedGInteractions-method  
  (pin), 35  
plyinteractions  
  (plyinteractions-package), 3  
plyinteractions-package, 3  
plyranges-flank, 37

plyranges-shift, 39  
 plyranges-stretch, 41  
  
 ranges1 (anchors1), 4  
 ranges1, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 ranges1, GInteractions-method  
     (anchors1), 4  
 ranges2 (anchors1), 4  
 ranges2, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 ranges2, GInteractions-method  
     (anchors1), 4  
 reexports, 42  
 regions, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 rename, 42  
 rename (reexports), 42  
 rename.GInteractions (dplyr-rename), 20  
 replace\_anchors, 43  
 replace\_anchors, AnchoredPinnedGInteractions, missing, GRanges-method  
     (replace\_anchors), 43  
 replace\_anchors, AnchoredPinnedGInteractions, numeric, GRanges-method  
     (replace\_anchors), 43  
 replace\_anchors, GInteractions, character, GenomicRanges-method  
     (replace\_anchors), 43  
 replace\_anchors, GInteractions, numeric, GenomicRanges-method  
     (replace\_anchors), 43  
 replace\_anchors, PinnedGInteractions, missing, GenomicRanges-method  
     (replace\_anchors), 43  
  
 second<-, GInteractions-method  
     (set\_seqnames1), 44  
 select, 3, 42  
 select (reexports), 42  
 select.GInteractions (dplyr-select), 21  
 seqinfo, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 seqnames1 (anchors1), 4  
 seqnames1, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 seqnames1, GInteractions-method  
     (anchors1), 4  
 seqnames2 (anchors1), 4  
  
 seqnames2, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 seqnames2, GInteractions-method  
     (anchors1), 4  
 set\_end1 (set\_seqnames1), 44  
 set\_end1, GInteractions, numeric-method  
     (set\_seqnames1), 44  
 set\_end2 (set\_seqnames1), 44  
 set\_end2, GInteractions, numeric-method  
     (set\_seqnames1), 44  
 set\_seqnames1, 44  
 set\_seqnames1, GInteractions, factor-method  
     (set\_seqnames1), 44  
 set\_seqnames2 (set\_seqnames1), 44  
 set\_seqnames2, GInteractions, factor-method  
     (set\_seqnames1), 44  
 set\_start1 (set\_seqnames1), 44  
 set\_start1, GInteractions, numeric-method  
     (set\_seqnames1), 44  
 set\_start2 (set\_seqnames1), 44  
 set\_start2, GInteractions, numeric-method  
     (set\_seqnames1), 44  
 set\_strand1 (set\_seqnames1), 44  
     set\_strand1, GInteractions, character-method  
         (set\_seqnames1), 44  
     set\_strand2 (set\_seqnames1), 44  
         set\_strand2, GInteractions, character-method  
             (set\_seqnames1), 44  
     set\_width1 (set\_seqnames1), 44  
         set\_width1, AnchoredPinnedGInteractions, numeric-method  
             (set\_seqnames1), 44  
     set\_width1, GInteractions, numeric-method  
         (set\_seqnames1), 44  
 set\_width2 (set\_seqnames1), 44  
 set\_width2, AnchoredPinnedGInteractions, numeric-method  
     (set\_seqnames1), 44  
 set\_width2, GInteractions, numeric-method  
     (set\_seqnames1), 44  
 shift\_downstream (plyranges-shift), 39  
 shift\_left (plyranges-shift), 39  
 shift\_right (plyranges-shift), 39  
 shift\_upstream (plyranges-shift), 39  
 show, DelegatingGInteractions-method  
     (delegating-ginteractions-methods),  
     11  
 slice, 3, 42  
 slice (reexports), 42

slice.GInteractions (dplyr-slice), 22  
start1 (anchors1), 4  
start1,DelegatingGInteractions-method  
(delegating-ginteractions-methods),  
11  
start1,GInteractions-method (anchors1),  
4  
start2 (anchors1), 4  
start2,DelegatingGInteractions-method  
(delegating-ginteractions-methods),  
11  
start2,GInteractions-method (anchors1),  
4  
strand1 (anchors1), 4  
strand1,DelegatingGInteractions-method  
(delegating-ginteractions-methods),  
11  
strand1,GInteractions-method  
(anchors1), 4  
strand2 (anchors1), 4  
strand2,DelegatingGInteractions-method  
(delegating-ginteractions-methods),  
11  
strand2,GInteractions-method  
(anchors1), 4  
stretch, 42  
stretch (reexports), 42  
stretch.ArcheredPinnedGInteractions  
(plyranges-stretch), 41  
stretch.PinnedGInteractions  
(plyranges-stretch), 41  
summarise, 42  
summarise (reexports), 42  
summarise.GroupedGInteractions  
(dplyr-summarize), 23  
summarize, 3, 42  
summarize (reexports), 42  
summarize.GroupedGInteractions  
(dplyr-summarize), 23  
  
tally, 3, 42  
tally (reexports), 42  
tally.GroupedGInteractions  
(dplyr-count), 14  
  
unanchor, 42  
unanchor (reexports), 42  
unanchor.ArcheredPinnedGInteractions  
(ginteractions-anchor), 24