# Package 'plyranges'

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Type Package

Title A fluent interface for manipulating GenomicRanges

**Version** 1.27.0

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**Description** A dplyr-like interface for interacting with the common Bioconductor classes Ranges and GenomicRanges. By providing a grammatical and consistent way of manipulating these classes their accessibility for new Bioconductor users is hopefully increased.

**Depends** R (>= 3.5), BiocGenerics, IRanges (>= 2.12.0), GenomicRanges (>= 1.28.4)

**Imports** methods, dplyr, rlang (>= 0.2.0), magrittr, tidyselect (>= 1.0.0), rtracklayer, GenomicAlignments, GenomeInfoDb, Rsamtools, S4Vectors (>= 0.23.10), utils

biocViews Infrastructure, DataRepresentation, WorkflowStep, Coverage

BugReports https://github.com/tidyomics/plyranges

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Suggests knitr, BiocStyle, rmarkdown, testthat (>= 2.1.0), HelloRanges, HelloRangesData, BSgenome.Hsapiens.UCSC.hg19, pasillaBamSubset, covr, ggplot2

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# Description

plyranges is a dplyr like API to the Ranges/GenomicRanges infrastructure in Bioconductor.

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#### **Details**

plryanges provides a consistent interface for importing and wrangling genomics data from a variety of sources. The package defines a grammar of genomic data manipulation through a set of verbs. These verbs can be used to construct human readable analysis pipelines based on Ranges objects.

- Modify genomic regions with the set\_width() and stretch() functions.
- Modify genomic regions while fixing the start/end/center coordinates with the anchors() family of functions.
- Sort genomic ranges with arrange().
- Modify, subset, and aggregate genomic data with the mutate(), filter(), and summarise() functions.
- Any of the above operations can be performed on partitions of the data with group\_by().
- Find nearest neighbour genomic regions with the join\_nearest() family of functions.
- Find overlaps between ranges with the join\_overlap\_inner() family of functions.
- Merge all overlapping and adjacent genomic regions with reduce\_ranges().
- Merge the end points of all genomic regions with disjoin\_ranges().
- Import and write common genomic data formats with the read\_/write\_ family of functions.

For more details on the features of plryanges, read the vignette: browseVignettes(package = "plyranges")

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#### See Also

Useful links:

• Report bugs at https://github.com/sa-lee/plyranges

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add\_nearest\_distance Add distance to nearest neighbours between two Ranges objects

# **Description**

Appends distance to nearest subject range to query ranges similar to setting distance in join\_nearest\_. Distance is set to NA for features with no nearest feature by the selected nearest metric.

#### Usage

```
add_nearest_distance(x, y = x, name = "distance")
add_nearest_distance_left(x, y = x, name = "distance")
add_nearest_distance_right(x, y = x, name = "distance")
add_nearest_distance_upstream(x, y = x, name = "distance")
add_nearest_distance_downstream(x, y = x, name = "distance")
```

#### **Arguments**

x	The query ranges
У	the subject ranges within which the nearest ranges are found. If missing, query ranges are used as the subject.
name	column name to create containing distance values

### Details

By default add\_nearest\_distance will find arbitrary nearest neighbours in either direction and ignore any strand information. The add\_nearest\_distance\_left and add\_nearest\_distance\_right methods will find arbitrary nearest neighbour ranges on x that are left/right of those on y and ignore any strand information.

The add\_nearest\_distance\_upstream method will find arbitrary nearest neighbour ranges on x that are upstream of those on y. This takes into account strandedness of the ranges. On the positive strand nearest upstream will be on the left and on the negative strand nearest upstream will be on the right.

The add\_nearest\_distance\_downstream method will find arbitrary nearest neighbour ranges on x that are upstream of those on y. This takes into account strandedness of the ranges. On the positive strand nearest downstream will be on the right and on the negative strand nearest upstream will be on the left.

#### Value

ranges in x with additional column containing the distance to the nearest range in y.

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# See Also

```
join_nearest
```

# **Examples**

anchor

Anchored Ranges objects

# Description

The GRangesAnchored class and the IRangesAnchored class allow components of a GRanges or IRanges (start, end, center) to be held fixed.

# Usage

```
anchor(x)
unanchor(x)
anchor_start(x)
anchor_end(x)
anchor_center(x)
anchor_centre(x)
anchor_3p(x)
anchor_5p(x)
```

# Arguments

Χ

a Ranges object

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#### **Details**

Anchoring will fix a Ranges start, end, or center positions, so these positions will remain the same when performing arithmetic. For GRanges objects, the function (anchor\_3p()) will fix the start for the negative strand, while anchor\_5p() will fix the end for the positive strand. Anchoring modifies how arithmetic is performed, for example modifying the width of a range with set\_width() or stretching a range with stretch(). To remove anchoring use unanchor().

#### Value

a RangesAnchored object which has the same appearance as a regular Ranges object but with an additional slot displaying an anchor.

#### **Constructors**

Depending on how you want to fix the components of a Ranges, there are five ways to construct a RangesAnchored class. Here x is either an IRanges or GRanges object.

- anchor\_start(x)Fix the start coordinates
- anchor\_end(x)Fix the end coordinates
- anchor\_center(x)Fix the center coordinates
- anchor\_3p(x)On the negative strand fix the start coordinates, and for positive or unstranded ranges fix the end coordinates.
- anchor\_5p(x)On the positive or unstranded ranges fix the start coordinates, coordinates and for negative stranded ranges fix the end coordinates.

#### Accessors

To see what has been anchored use the function anchor. This will return a character vector containing a valid anchor. It will be set to one of c("start", "end", "center") for an IRanges object or one of c("start", "end", "center", "3p", "5p") for a GRanges object.

#### See Also

mutate, stretch

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arrange.Ranges

Sort a Ranges object

# **Description**

Sort a Ranges object

#### Usage

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

### **Arguments**

.data A Ranges object.

. . . Comma seperated list of variable names.

#### Value

A sorted Ranges object

# **Examples**

```
rng <- as_iranges(data.frame(start = 1:10, width = 10:1))
rng <- mutate(rng, score = runif(10))
arrange(rng, score)
# you can also use dplyr::desc to arrange by descending order</pre>
```

as\_iranges

Construct a I/GRanges object from a tibble or data.frame

# Description

The as\_i(g)ranges function looks for column names in .data called start, end, width, seqnames and strand in order to construct an IRanges or GRanges object. By default other columns in .data are placed into the mcols (metadata columns) slot of the returned object.

### Usage

```
as_iranges(.data, ..., keep_mcols = TRUE)
as_granges(.data, ..., keep_mcols = TRUE)
```

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# **Arguments**

.data a data.frame() or tibble() to construct a Ranges object from... optional named arguments specifying which the columns in .data containin the core components a Ranges object.

keep\_mcols place the remaining columns into the metadata columns slot (default=TRUE)

#### Value

a Ranges object.

#### See Also

```
IRanges::IRanges(), GenomicRanges::GRanges()
```

# **Examples**

```
df <- data.frame(start=c(2:-1, 13:15), width=c(0:3, 2:0))</pre>
as_iranges(df)
df <- data.frame(start=c(2:-1, 13:15), width=c(0:3, 2:0), strand = "+")
# will return an IRanges object
as_iranges(df)
df <- data.frame(start=c(2:-1, 13:15), width=c(0:3, 2:0),</pre>
strand = "+", seqnames = "chr1")
as_granges(df)
\# as_g/iranges understand alternate name specification
df <- data.frame(start=c(2:-1, 13:15), width=c(0:3, 2:0),</pre>
strand = "+", chr = "chr1")
as_granges(df, seqnames = chr)
# can also handle DFrame input
df <- methods::as(df, "DFrame")</pre>
df$y <- IRanges::IntegerList(c(1,2,3), NA, 5, 6, 8, 9, 10:12)
as_iranges(df)
as_granges(df, seqnames = chr)
```

as\_ranges

Coerce an Rle or RleList object to Ranges

# **Description**

Coerce an Rle or RleList object to Ranges

### Usage

```
as_ranges(.data)
```

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#### **Arguments**

```
.data a Rle() or an RleList() object.
```

#### **Details**

This function is behind compute\_coverage().

#### Value

```
an IRanges() object if the input is an Rle() object or a GRanges() object for an RleList() object.
```

#### See Also

```
S4Vectors::Rle(), IRanges::RleList()
```

# **Examples**

```
x <- S4Vectors::Rle(10:1, 1:10)
as_ranges(x)

# must have names set
y <- IRanges::RleList(chr1 = x)
as_ranges(y)</pre>
```

bind\_ranges

Combine Ranges by concatentating them together

# **Description**

Combine Ranges by concatentating them together

### Usage

```
bind_ranges(..., .id = NULL)
```

### **Arguments**

Ranges objects to combine. Each argument can be a Ranges object, or a list of Ranges objects.

. id Ranges object identifier. When .id is supplied a new column is created that links

each row to the original Range object. The contents of the column correspond to the named arguments or the names of the list supplied.

#### Value

a concatenated Ranges object

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#### Note

Currently GRangesList or IRangesList objects are not supported.

#### **Examples**

chop\_by\_introns

Group a GRanges object by introns or gaps

#### **Description**

Group a GRanges object by introns or gaps

#### Usage

```
chop_by_introns(x)
chop_by_gaps(x)
```

# Arguments

Χ

a GenomicRanges object with a cigar string column

#### **Details**

Creates a grouped Ranges object from a cigar string column, for chop\_by\_introns() will check for the presence of "N" in the cigar string and create a new column called intron where TRUE indicates the alignment has a skipped region from the reference. For chop\_by\_gaps() will check for the presence of "N" or "D" in the cigar string and create a new column called "gaps" where TRUE indicates the alignment has a deletion from the reference or has an intron.

#### Value

```
a GRanges object
```

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#### **Examples**

compute\_coverage

Compute coverage over a Ranges object

# **Description**

Compute coverage over a Ranges object

# Usage

```
compute_coverage(x, shift, width, weight, ...)
```

# Arguments

X	a Ranges object
shift	shift how much should each range in x be shifted by? (default = $0L$ )
width	width how long should the returned coverage score be? This must be either a positive integer or NULL (default = NULL)
weight	weight how much weight should be assigned to each range? Either an integer or numeric vector or a column in $x$ . (default = $1L$ )
	other optional parameters to pass to coverage

#### Value

An expanded Ranges object with a score column corresponding to the coverage value over that interval. Note that compute\_coverage drops metadata associated with the original ranges.

#### See Also

```
IRanges::coverage(), GenomicRanges::coverage()
```

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#### **Examples**

```
rng <- as_iranges(data.frame(start = 1:10, width = 5))
compute_coverage(rng)
compute_coverage(rng, shift = 14L)
compute_coverage(rng, width = 10L)</pre>
```

count\_overlaps

Count the number of overlaps between two Ranges objects

### **Description**

Count the number of overlaps between two Ranges objects

### Usage

```
count_overlaps(x, y, maxgap, minoverlap)
## S3 method for class 'IntegerRanges'
count_overlaps(x, y, maxgap = -1L, minoverlap = 0L)
## S3 method for class 'GenomicRanges'
count_overlaps(x, y, maxgap = -1L, minoverlap = 0L)
count_overlaps_within(x, y, maxgap, minoverlap)
## S3 method for class 'IntegerRanges'
count_overlaps_within(x, y, maxgap = 0L, minoverlap = 1L)
## S3 method for class 'GenomicRanges'
count_overlaps_within(x, y, maxgap = 0L, minoverlap = 1L)
count_overlaps_directed(x, y, maxgap, minoverlap)
## S3 method for class 'GenomicRanges'
count\_overlaps\_directed(x, y, maxgap = -1L, minoverlap = 0L)
count_overlaps_within_directed(x, y, maxgap, minoverlap)
## S3 method for class 'GenomicRanges'
count\_overlaps\_within\_directed(x, y, maxgap = -1L, minoverlap = 0L)
```

### **Arguments**

```
x, y Objects representing ranges maxgap, minoverlap
```

The maximimum gap between intervals as an integer greater than or equal to zero. The minimum amount of overlap between intervals as an integer greater than zero, accounting for the maximum gap.

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# Value

An integer vector of same length as x.

### **Examples**

DeferredGenomicRanges-class

DeferredGenomiRanges objects

# Description

Enables deferred reading of files (currently only BAM files) by caching results after a plyranges verb is called.

# **Slots**

```
delegate a GenomicRanges object to be cached ops A FileOperator object
```

### See Also

```
read_bam()
```

disjoin\_ranges

Disjoin then aggregate a Ranges object

# **Description**

Disjoin then aggregate a Ranges object

# Usage

```
disjoin_ranges(.data, ...)
disjoin_ranges_directed(.data, ...)
```

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# **Arguments**

```
.data a Ranges object to disjoin... Name-value pairs of summary functions.
```

#### Value

a Ranges object that is now disjoint (no bases overlap).

# **Examples**

```
df <- data.frame(start = 1:10, width = 5, seqnames = "seq1",
strand = sample(c("+", "-", "*"), 10, replace = TRUE), gc = runif(10))
rng <- as_granges(df)
rng %>% disjoin_ranges()
rng %>% disjoin_ranges(gc = mean(gc))
rng %>% disjoin_ranges_directed(gc = mean(gc))
```

expand\_ranges

Expand list-columns in a Ranges object

# **Description**

Expand list-columns in a Ranges object

# Usage

```
expand_ranges(
  data,
  ...,
  .drop = FALSE,
  .id = NULL,
  .keep_empty = FALSE,
  .recursive = FALSE
)
```

# **Arguments**

data	A Ranges object
	list-column names to expand then unlist
.drop	Should additional list columns be dropped (default = FALSE)? By default expand_ranges() will keep other list columns even if they are nested.
.id	A character vector of length equal to number of list columns. If supplied will create new $column(s)$ with name .id identifying the index of the list column (default = NULL).
.keep_empty	If a list-like column contains empty elements, should those elements be kept? (default = FALSE)

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.recursive

If there are multiple list-columns, should the columns be treated as parallel? If FALSE each column will be unnested recursively, otherwise they are treated as parallel, that is each list column has identical lengths. (deafualt = FALSE)

#### Value

a GRanges object with expanded list columns

### **Examples**

FileOperator-class

An abstract class to represent operations performed over a file

# Description

An abstract class to represent operations performed over a file

#### **Details**

This class is used internally by DeferredGenomicRanges objects. Currently, this class is only implemented for bam files (as a BamFileOperator) but will eventually be extended to the other available readers.

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object

filter-ranges	Subset a Ranges
TITUEL Langes	Subset a Nanges

# **Description**

Subset a Ranges object

# Usage

```
## S3 method for class 'Ranges'
filter(.data, ..., .preserve = FALSE)
```

# **Arguments**

.data A Ranges object
... valid logical predictates to subset .data by. These are determined by variables in .data. If more than one condition is supplied, the conditions are combined with &. Only rows where the condition evaluates to TRUE are kept.

. preserve when FALSE (the default) grouping structure is recalculated, TRUE is currently

not implemented.

#### **Details**

For any Ranges objects filter can act on all core components of the class including start, end, width (for IRanges) or seqnames and strand (for GRanges) in addition to metadata columns. If the Ranges object is grouped, filter will act seperately on each parition of the data.

#### Value

a Ranges object

#### See Also

```
dplyr::filter()
```

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```
# multiple criteria
filter(rng, strand == "+" | start > 5)
filter(rng, strand == "+" & start > 5)

# multiple conditions are the same as and
filter(rng, strand == "+", start > 5)

# grouping acts on each subset of the data
rng %>%
    group_by(strand) %>%
    filter(gc > 0.5)
```

filter\_by\_overlaps

Filter by overlapping/non-overlapping ranges

### Description

Filter by overlapping/non-overlapping ranges

### Usage

```
filter_by_overlaps(x, y, maxgap = -1L, minoverlap = 0L)
filter_by_non_overlaps(x, y, maxgap, minoverlap)
filter_by_overlaps_directed(x, y, maxgap = -1L, minoverlap = 0L)
filter_by_non_overlaps_directed(x, y, maxgap, minoverlap)
```

#### Arguments

x, y Objects representing ranges

maxgap The maximimum gap between intervals as a single integer greater than or equal

to -1. If you modify this argument, minoverlap must be held fixed.

minoverlap The minimum amount of overlap between intervals as a single integer greater

than 0. If you modify this argument, maxgap must be held fixed.

#### **Details**

By default, filter\_by\_overlaps and filter\_by\_non\_overlaps ignore strandedness for GRanges() objects. To perform stranded operations use filter\_by\_overlaps\_directed and filter\_by\_non\_overlaps\_directed. The argument maxgap is the maximum number of positions between two ranges for them to be considered overlapping. Here the default is set to be -1 as that is the the gap between two ranges that has its start or end strictly inside the other. The argument minoverlap refers to the minimum number of positions overlapping between ranges, to consider there to be overlap.

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#### Value

a Ranges object

#### See Also

IRanges::subsetByOverlaps()

#### **Examples**

```
df <- data.frame(seqnames = c("chr1", rep("chr2", 2),</pre>
                               rep("chr3", 3), rep("chr4", 4)),
                 start = 1:10,
                 width = 10:1,
                 strand = c("-", "+", "+", "*", "*", "+", "+", "+", "-", "-")
                 name = letters[1:10])
query <- as_granges(df)
df2 <- data.frame(seqnames = c(rep("chr2", 2), rep("chr1", 3), "chr2"),</pre>
                  start = c(4,3,7,13,1,4),
                  width = c(6,6,3,3,3,9),
                  strand = c(rep("+", 3), rep("-", 3)))
subject <- as_granges(df2)</pre>
filter_by_overlaps(query, subject)
filter_by_overlaps_directed(query, subject)
filter_by_non_overlaps(query, subject)
filter_by_non_overlaps_directed(query, subject)
```

find\_overlaps

Find overlap between two Ranges

# Description

Find overlap between two Ranges

# Usage

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

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```
find_overlaps_within(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))
## S3 method for class 'IntegerRanges'
find_overlaps_within(
  х,
 у,
 maxgap = -1L,
 minoverlap = 0L,
  suffix = c(".x", ".y")
)
## S3 method for class 'GenomicRanges'
find_overlaps_within(
  Х,
  у,
  maxgap = -1L,
 minoverlap = 0L,
  suffix = c(".x", ".y")
)
find_overlaps_directed(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))
## S3 method for class 'GenomicRanges'
find_overlaps_directed(
 х,
 у,
 maxgap = -1L,
 minoverlap = 0L,
  suffix = c(".x", ".y")
)
find_overlaps_within_directed(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))
## S3 method for class 'GenomicRanges'
find_overlaps_within_directed(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))
group_by_overlaps(x, y, maxgap, minoverlap)
## S3 method for class 'IntegerRanges'
group_by_overlaps(x, y, maxgap = -1L, minoverlap = 0L)
## S3 method for class 'GenomicRanges'
group_by_overlaps(x, y, maxgap = -1L, minoverlap = 0L)
```

### Arguments

```
x, y Objects representing ranges maxgap, minoverlap
```

The maximimum gap between intervals as an integer greater than or equal to

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negative one. The minimum amount of overlap between intervals as an integer greater than zero, accounting for the maximum gap.

suffix

A character vector of length two used to identify metadata columns coming from x and y.

#### **Details**

find\_overlaps() will search for any overlaps between ranges x and y and return a Ranges object of length equal to the number of times x overlaps y. This Ranges object will have additional metadata columns corresponding to the metadata columns in y. find\_overlaps\_within() is the same but will only search for overlaps within y. For GRanges objects strand is ignored, unless find\_overlaps\_directed() is used. If the Ranges objects have no metadata, one could use group\_by\_overlaps() to be able to identify the index of the input Range x that overlaps a Range in y. Alternatively, pair\_overlaps() could be used to place the x ranges next to the range in y they overlap.

#### Value

A Ranges object with rows corresponding to the ranges in x that overlap y. In the case of group\_by\_overlaps(), returns a GroupedRanges object, grouped by the number of overlaps of ranges in x that overlap y (stored in a column called query).

#### See Also

```
IRanges::findOverlaps(), GenomicRanges::findOverlaps()
```

```
query <- data.frame(start = c(5,10, 15,20), width = 5, gc = runif(4)) %>%
             as_iranges()
subject <- data.frame(start = 2:6, width = 3:7, label = letters[1:5]) %>%
             as_iranges()
find_overlaps(query, subject)
find_overlaps(query, subject, minoverlap = 5)
find_overlaps_within(query, subject) # same result as minoverlap
find_overlaps(query, subject, maxgap = 1)
# -- GRanges objects, strand is ignored by default
query <- data.frame(seqnames = "chr1",</pre>
               start = c(11, 101),
               end = c(21, 200),
               name = c("a1", "a2"),
               strand = c("+", "-"),
               score = c(1,2)) %>%
           as_granges()
subject <- data.frame(seqnames = "chr1",</pre>
                      strand = c("+", "-", "+", "-"),
                      start = c(21, 91, 101, 201),
                      end = c(30,101,110,210),
                      name = paste0("b", 1:4),
```

22 flank\_left

flank\_left

Generate flanking regions

# Description

Find flanking regions to the left or right or upstream or downstream of a Ranges object.

# Usage

```
flank_left(x, width = 0L)
flank_right(x, width = 0L)
flank_upstream(x, width = 0L)
flank_downstream(x, width = 0L)
```

# **Arguments**

x a Ranges 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.

#### **Details**

The function flank\_left will create the flanking region to the left of starting coordinates in x, while flank\_right will create the flanking region to the right of the starting coordinates in x. The function flank\_upstream will flank\_left if the strand of rows in x is not negative and will flank\_right if the strand of rows in x is negative. The function flank\_downstream will flank\_right if the strand of rows in x is not negative and will flank\_left if the strand of rows in x is negative.

By default flank\_left and flank\_right will ignore strandedness of any ranges, while flank\_upstream and flank\_downstream will take into account the strand of x.

#### Value

A Ranges object of same length as x.

#### See Also

```
IRanges::flank(), GenomicRanges::flank()
```

# **Examples**

GroupedGenomicRanges-class

Group a Ranges by one or more variables

# Description

The function group\_by takes a Ranges object and defines groups by one or more variables. Operations are then performed on the Ranges by their "group". ungroup() removes grouping.

# Usage

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

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

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

## S3 method for class 'GroupedIntegerRanges'
groups(x)
```

### **Arguments**

.data	a Ranges object.
	Variable names to group by. These can be either metadata columns or the core variables of a Ranges.
add	if .data is already a GroupedRanges object, when add = FALSE the (default), group_by() will override existing groups. If add = TRUE, additional groups will be added.
Х	a GroupedRanges object.

#### **Details**

group\_by() creates a new object of class GroupedGenomicRanges if the input is a GRanges object or an object of class GroupedIntegerRanges if the input is a IRanges object. Both of these classes contain a slot called groups corresponding to the names of grouping variables. They also inherit from their parent classes, Ranges and GenomicRanges respectively. ungroup() removes the grouping and will return either a GRanges or IRanges object.

#### Value

The group\_by() function will return a GroupedRanges object. These have the same appearance as a regular Ranges object but with an additional groups slot.

#### Accessors

To return grouping variables on a grouped Ranges use either

- groups(x)Returns a list of symbols
- group\_vars(x)Returns a character vector

```
set.seed(100)
df <- data.frame(start = 1:10,</pre>
                 width = 5,
                 gc = runif(10),
                 cat = sample(letters[1:2], 10, replace = TRUE))
rng <- as_iranges(df)</pre>
rng_by_cat <- rng %>% group_by(cat)
# grouping does not change appearance or shape of Ranges
rng_by_cat
# a list of symbols
groups(rng_by_cat)
# ungroup removes any grouping
ungroup(rng_by_cat)
# group_by works best with other verbs
grng <- as_granges(df,</pre>
                   seqnames = "chr1",
                   strand = sample(c("+", "-"), size = 10, replace = TRUE))
grng_by_strand <- grng %>% group_by(strand)
grng_by_strand
# grouping with other verbs
grng_by_strand %>% summarise(gc = mean(gc))
grng_by_strand %>% filter(gc == min(gc))
grng_by_strand %>%
 ungroup() %>%
 summarise(gc = mean(gc))
```

intersect\_ranges 25

 $intersect\_ranges$ 

Vector-wise Range set-operations

# Description

Vector-wise Range set-operations

# Usage

```
intersect_ranges(x, y)
intersect_ranges_directed(x, y)
union_ranges(x, y)
union_ranges_directed(x, y)
setdiff_ranges(x, y)
setdiff_ranges_directed(x, y)
complement_ranges(x)
```

# Arguments

x, y

Two Ranges objects to compare.

#### **Details**

These are usual set-operations that act on the sets of the ranges represented in x and y. By default these operations will ignore any strand information. The directed versions of these functions will take into account strand for GRanges objects.

# Value

A Ranges object

26 interweave

```
union_ranges(gr1, gr2)
union_ranges_directed(gr1, gr2)
intersect_ranges(gr1, gr2)
intersect_ranges_directed(gr1, gr2)
setdiff_ranges(gr1, gr2)
setdiff_ranges_directed(gr1, gr2)
# taking the complement of a ranges requires annotation information
gr1 <- set_genome_info(gr1, seqlengths = 100)
complement_ranges(gr1)</pre>
```

interweave

Interweave a pair of Ranges objects together

### Description

Interweave a pair of Ranges objects together

#### Usage

```
interweave(left, right, .id = NULL)
```

#### **Arguments**

left, right Ranges objects.

. id When supplied a new column that represents the origin column and is linked to

each row of the resulting Ranges object.

#### **Details**

The output of interweave() takes pairs of Ranges objects and combines them into a single Ranges object. If an .id argument is supplied, an origin column with name .id is created indicated which side the resulting Range comes from (eit)

### Value

a Ranges object

join\_follow 27

join\_follow

Find following Ranges

#### **Description**

Find following Ranges

# Usage

```
join_follow(x, y, suffix = c(".x", ".y"))
join_follow_left(x, y, suffix = c(".x", ".y"))
join_follow_upstream(x, y, suffix = c(".x", ".y"))
```

# **Arguments**

x, y Ranges objects, which ranges in x follow those in y.
 suffix A character vector of length two used to identify metadata columns coming from x and y.

#### **Details**

By default join\_follow will find abritrary ranges in y that are followed by ranges in x and ignore any strand information. On the other hand join\_follow\_left will find all ranges in y that are on the left-hand side of the ranges in x ignoring any strand information. Finally, join\_follow\_upstream will find all ranges in x that are that are upstream of the ranges in y. On the positive strand this will result in ranges in y that are left of those in x and on the negative strand it will result in ranges in y that are right of those in x.

#### Value

A Ranges object corresponding to the ranges in x ` that are followed by the ranges in y, all metadata is copied of

join\_nearest

join\_nearest

Find nearest neighbours between two Ranges objects

# Description

Find nearest neighbours between two Ranges objects

#### Usage

```
join\_nearest(x, y, suffix = c(".x", ".y"), distance = FALSE)
join\_nearest\_left(x, y, suffix = c(".x", ".y"), distance = FALSE)
join\_nearest\_right(x, y, suffix = c(".x", ".y"), distance = FALSE)
join\_nearest\_upstream(x, y, suffix = c(".x", ".y"), distance = FALSE)
join\_nearest\_downstream(x, y, suffix = c(".x", ".y"), distance = FALSE)
```

### Arguments

x, y	Ranges objects, add the nearest neighbours of ranges in x to those in y.
suffix	A character vector of length two used to identify metadata columns
distance	logical vector whether to add a column named "distance" containing the distance to the nearest region. If set to a character vector of length 1, will use that as distance column name.

#### **Details**

By default join\_nearest will find arbitrary nearest neighbours in either direction and ignore any strand information. The join\_nearest\_left and join\_nearest\_right methods will find arbitrary nearest neighbour ranges on x that are left/right of those on y and ignore any strand information.

The join\_nearest\_upstream method will find arbitrary nearest neighbour ranges on x that are upstream of those on y. This takes into account strandedness of the ranges. On the positive strand nearest upstream will be on the left and on the negative strand nearest upstream will be on the right.

The join\_nearest\_downstream method will find arbitrary nearest neighbour ranges on x that are upstream of those on y. This takes into account strandedness of the ranges. On the positive strand nearest downstream will be on the right and on the negative strand nearest upstream will be on the left.

#### Value

A Ranges object corresponding to the nearest ranges, all metadata is copied over from the right-hand side ranges y.

#### **Examples**

```
query <- data.frame(start = c(5,10, 15,20),
                   width = 5,
                   gc = runif(4)) %>%
             as_iranges()
subject <- data.frame(start = c(2:6, 24),</pre>
                      width = 3:8,
                      label = letters[1:6]) %>%
             as_iranges()
join_nearest(query, subject)
join_nearest_left(query, subject)
join_nearest_right(query, subject)
subject <- data.frame(seqnames = "chr1",</pre>
               start = c(11, 101),
               end = c(21, 200),
               name = c("a1", "a2"),
               strand = c("+", "-"),
               score = c(1,2)) \% > \%
           as_granges()
query <- data.frame(seqnames = "chr1",</pre>
                      strand = c("+", "-", "+", "-"),
                       start = c(21,91,101,201),
                       end = c(30,101,110,210),
                      name = paste0("b", 1:4),
                       score = 1:4) %>%
                   as_granges()
join_nearest_upstream(query, subject)
join_nearest_downstream(query, subject)
```

join\_overlap\_intersect

Join by overlapping Ranges

#### **Description**

Join by overlapping Ranges

#### Usage

```
join\_overlap\_intersect(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))
join\_overlap\_intersect\_within(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))
join_overlap_intersect_directed(
 Х,
 у,
 maxgap,
 minoverlap,
 suffix = c(".x", ".y")
)
join_overlap_intersect_within_directed(
 х,
 у,
 maxgap,
 minoverlap,
 suffix = c(".x", ".y")
join\_overlap\_inner(x, y, maxgap = -1L, minoverlap = 0L, suffix = c(".x", ".y"))
join_overlap_inner_within(
 х,
 у,
 maxgap = -1L,
 minoverlap = 0L,
 suffix = c(".x", ".y")
)
join_overlap_inner_directed(
 х,
 у,
 maxgap = -1L,
 minoverlap = 0L,
 suffix = c(".x", ".y")
)
join_overlap_inner_within_directed(
 у,
 maxgap = -1L,
 minoverlap = 0L,
```

```
suffix = c(".x", ".y")
)

join_overlap_left(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))

join_overlap_left_within(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))

join_overlap_left_directed(x, y, maxgap, minoverlap, suffix = c(".x", ".y"))

join_overlap_left_within_directed(
    x,
    y,
    maxgap,
    minoverlap,
    suffix = c(".x", ".y")
)
```

### **Arguments**

x, y Objects representing ranges maxgap, minoverlap

The maximimum gap between intervals as an integer greater than or equal to zero. The minimum amount of overlap between intervals as an integer greater than zero, accounting for the maximum gap.

suffix Character to vectors to append to common columns in x and y (default = c(".x", ".y")).

#### Details

The function join\_overlap\_intersect() finds the genomic intervals that are the overlapping ranges between x and y and returns a new ranges object with metadata columns from x and y.

The function join\_overlap\_inner() is equivalent to find\_overlaps().

The function <code>join\_overlap\_left()</code> performs a left outer join between x and y. It returns all ranges in x that overlap or do not overlap ranges in y plus metadata columns common to both. If there is no overlapping range the metadata column will contain a missing value.

The function join\_overlap\_self() find all overlaps between a ranges object x and itself.

All of these functions have two suffixes that modify their behavior. The within suffix, returns only ranges in x that are completely overlapped within in y. The directed suffix accounts for the strandedness of the ranges when performing overlaps.

#### Value

a GRanges object

#### See Also

```
join_overlap_self(), join_overlap_left(), find_overlaps()
```

join\_overlap\_self

#### **Examples**

join\_overlap\_self

Find overlaps within a Ranges object

#### **Description**

Find overlaps within a Ranges object

#### Usage

```
join_overlap_self(x, maxgap, minoverlap)
join_overlap_self_within(x, maxgap, minoverlap)
join_overlap_self_directed(x, maxgap, minoverlap)
join_overlap_self_within_directed(x, maxgap, minoverlap)
```

### **Arguments**

x A Ranges object

maxgap, minoverlap

The maximimum gap between intervals as an integer greater than or equal to zero. The minimum amount of overlap between intervals as an integer greater than zero, accounting for the maximum gap.

join\_precede 33

#### **Details**

Self overlaps find any overlaps (or overlaps within or overlaps directed) between a ranges object and itself.

#### Value

a Ranges object

#### See Also

```
find_overlaps(), join_overlap_inner()
```

#### **Examples**

```
query <- data.frame(start = c(5,10, 15,20), width = 5, gc = runif(4)) %>%
             as_iranges()
join_overlap_self(query)
# -- GRanges objects, strand is ignored by default
query <- data.frame(seqnames = "chr1",</pre>
               start = c(11, 101),
               end = c(21, 200),
               name = c("a1", "a2"),
               strand = c("+", "-"),
               score = c(1,2)) \%
           as_granges()
# ignores strandedness
join_overlap_self(query)
join_overlap_self_within(query)
# adding directed prefix includes strand
join_overlap_self_directed(query)
```

join\_precede

Find preceding Ranges

# **Description**

Find preceding Ranges

# Usage

```
join_precede(x, y, suffix = c(".x", ".y"))
join_precede_right(x, y, suffix = c(".x", ".y"))
join_precede_downstream(x, y, suffix = c(".x", ".y"))
```

join\_precede

# Arguments

x, yRanges objects, which ranges in x precede those in y.suffixA character vector of length two used to identify metadata columns coming from x and y.

#### **Details**

By default join\_precede will return the ranges in x that come before the ranges in y and ignore any strand information. The function join\_precede\_right will find all ranges in y that are on the right-hand side of the ranges in x ignoring any strand information. Finally, join\_precede\_downstream will find all ranges in y that are that are downstream of the ranges in x. On the positive strand this will result in ranges in y that are right of those in x and on the negative strand it will result in ranges in y that are left of those in x.

#### Value

A Ranges object corresponding to the ranges in y that are preceded by the ranges in x, all metadata is copied over from the right-hand side ranges y.

```
subject <- data.frame(start = c(5,10, 15,20), width = 5, gc = runif(4)) %>%
             as_iranges()
query <- data.frame(start = 2:6, width = 3:7, label = letters[1:5]) %>%
             as_iranges()
join_precede(query, subject)
query <- data.frame(seqnames = "chr1",</pre>
               start = c(11, 101),
               end = c(21, 200),
               name = c("a1", "a2"),
               strand = c("+", "-"),
               score = c(1,2)) \% > \%
           as_granges()
subject <- data.frame(seqnames = "chr1",</pre>
                      strand = c("+", "-", "+", "-"),
                      start = c(21,91,101,201),
                      end = c(30,101,110,210),
                      name = paste0("b", 1:4),
                      score = 1:4) %>%
                   as_granges()
join_precede(query, subject)
join_precede_right(query, subject)
join_precede_downstream(query, subject)
```

mutate.Ranges 35

mutate.Ranges

Modify a Ranges object

#### **Description**

Modify a Ranges object

# Usage

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

# **Arguments**

.data a Ranges object

Pairs of name-value expressions. The name-value pairs can either create new metadata columns or modify existing ones.

#### Value

a Ranges object

```
df <- data.frame(start = 1:10,</pre>
                 width = 5,
                 seqnames = "seq1",
                 strand = sample(c("+", "-", "*"), 10, replace = TRUE),
                 gc = runif(10)
rng <- as_granges(df)</pre>
# mutate adds new columns
rng %>%
   mutate(avg_gc = mean(gc), row_id = 1:n())
# can also compute on newly created columns
rng %>%
    mutate(score = gc * width, score2 = score + 1)
# group by partitions the data and computes within each group
rng %>%
    group_by(strand) %>%
   mutate(avg_gc = mean(gc), row_id = 1:n())
# mutate can be used in conjuction with anchoring to resize ranges
    mutate(width = 10)
# by default width modfication fixes by start
rng %>%
   anchor_start() %>%
   mutate(width = 10)
```

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```
# fix by end or midpoint
rng %>%
    anchor_end() %>%
    mutate(width = width + 1)
rng %>%
    anchor_center() %>%
    mutate(width = width + 1)
# anchoring by strand
rng %>%
    anchor_3p() %>%
    mutate(width = width * 2)
rng %>%
    anchor_5p() %>%
    mutate(width = width * 2)
```

Compute the number of ranges in each group.

# Description

This function should only be used within summarise(), mutate() and filter().

# Usage

n()

# Value

n() will only be evaluated inside a function call, where it returns an integer.

# Examples

n

n\_distinct 37

n\_distinct

Compute the number of distinct unique values in a vector or List

# Description

This is a wrapper to length(unique(x)) or lengths(unique(x)) if x is a List object

# Usage

```
n_distinct(var)
```

# **Arguments**

var

a vector of values

#### Value

an integer vector

# **Examples**

```
x <- CharacterList(c("a", "b", "c", "a"), "d")
n_distinct(x)
n_distinct(unlist(x))</pre>
```

overscope\_ranges

Create an overscoped environment from a Ranges object

# Description

Create an overscoped environment from a Ranges object

# Usage

```
overscope_ranges(x, envir = parent.frame())
```

# **Arguments**

```
x a Ranges object
```

envir the environment to place the Ranges in (default = parent.frame())

## **Details**

This is the backend for non-standard evaluation in plyranges.

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## Value

an environment

#### See Also

```
rlang::new_data_mask(), rlang::eval_tidy()
```

pair\_overlaps

Pair together two ranges objects

# Description

Pair together two ranges objects

# Usage

```
pair_overlaps(x, y, maxgap, minoverlap, suffix)
pair_nearest(x, y, suffix)
pair_precede(x, y, suffix)
pair_follow(x, y, suffix)
```

# **Arguments**

x, y Ranges objects to pair together.

maxgap, minoverlap

The maximimum gap between intervals as an integer greater than or equal to negative one. The minimum amount of overlap between intervals as an integer greater than zero, accounting for the maximum gap.

suffix

A character vector of length two used to identify metadata columns coming from x and y.

# **Details**

These functions return a DataFrame object, and is one way of representing paired alignments with plyranges.

# Value

a DataFrame with two ranges columns and the corresponding metadata columns.

## See Also

```
[join_nearest()][join_overlap_inner()][join_precede()][join_follow()]
```

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## **Examples**

```
query <- data.frame(start = c(5,10, 15,20), width = 5, gc = runif(4)) %>%
subject <- data.frame(start = 2:6, width = 3:7, label = letters[1:5]) %>%
             as_iranges()
pair_overlaps(query, subject)
pair_overlaps(query, subject, minoverlap = 5)
pair_nearest(query, subject)
query <- data.frame(segnames = "chr1",</pre>
               start = c(11,101),
               end = c(21, 200),
               name = c("a1", "a2"),
strand = c("+", "-"),
                score = c(1,2)) %>%
           as_granges()
subject <- data.frame(seqnames = "chr1",</pre>
                       strand = c("+", "-", "+", "-"),
                       start = c(21,91,101,201),
                       end = c(30, 101, 110, 210),
                       name = paste0("b", 1:4),
                       score = 1:4) %>%
                    as_granges()
# ignores strandedness
pair_overlaps(query, subject, suffix = c(".query", ".subject"))
pair_follow(query, subject, suffix = c(".query", ".subject"))
pair_precede(query, subject, suffix = c(".query", ".subject"))
pair_precede(query, subject, suffix = c(".query", ".subject"))
```

ranges-info

Construct annotation information

## **Description**

To construct annotations by supplying annotation information use genome\_info. To add annotations to an existing Ranges object use set\_genome\_info. To retrieve an annotation as a Ranges object use get\_genome\_info.

# Usage

```
genome_info(
  genome = NULL,
  seqnames = NULL,
  seqlengths = NULL,
  is_circular = NULL)
```

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```
set_genome_info(
    .data,
    genome = NULL,
    seqnames = NULL,
    seqlengths = NULL,
    is_circular = NULL
)
get_genome_info(.data)
```

## **Arguments**

genome A character vector of length one indicating the genome build.

seqnames A character vector containing the name of sequences.

seqlengths An optional integer vector containg the lengths of sequences.

is\_circular An optional logical vector indicating whether a sequence is ciruclar.

. data A Ranges object to annotate or retrieve an annotation for.

#### Value

a GRanges object containing annotations. To retrieve the annotations as a Ranges object use get\_genome\_info.

#### See Also

```
GenomeInfoDb::Seqinfo()
```

```
x <- genome_info(genome = "toy",</pre>
                  segnames = letters[1:4],
                  seqlengths = c(100, 300, 15, 600),
                  is_circular = c(NA, FALSE, FALSE, TRUE))
Х
rng <- as_granges(data.frame(seqnames = "a", start = 30:50, width = 10))</pre>
rng <- set_genome_info(rng,</pre>
                        genome = "toy",
                        seqnames = letters[1:4],
                        seqlengths = c(100, 300, 15, 600),
                        is_circular = c(NA, FALSE, FALSE, TRUE))
get_genome_info(rng)
## Not run:
if (interactive()) {
 # requires internet connection
 genome_info(genome = "hg38")
}
```

read\_bam 41

## End(Not run)

read\_bam

Read a BAM file

#### **Description**

Read a BAM file

## Usage

```
read_bam(file, index = file, paired = FALSE)
```

## **Arguments**

file A connection or path to a BAM file

index The path to the BAM index file

paired Whether to treat alignments as paired end (TRUE) or single end (FALSE). De-

fault is FALSE.

## Details

Reading a BAM file is deferred until an action such as using summarise() or mutate() occurs. If paired is set to TRUE, when alignments are loaded, the GRanges has two additional columns called read\_pair\_id and read\_pair\_group corresponding to paired reads and is grouped by the read\_pair\_group.

Certain verbs have different behaviour, after using read\_bam().

For select() valid columns are the fields available in the BAM file. Valid entries are qname (QNAME), flag (FLAG), rname (RNAME), strand, pos (POS), qwidth (width of query), mapq (MAPQ), cigar (CIGAR), mrnm (RNEXT), mpos (PNEXT), isize (TLEN), seq (SEQ), and qual (QUAL). Any two character tags in the BAM file are also valid.

For filter() the following fields are valid, to select the FALSE option place! in front of the field:

- is\_paired Select either unpaired (FALSE) or paired (TRUE) reads.
- is\_proper\_pair Select either improperly paired (FALSE) or properly paired (TRUE) reads. This is dependent on the alignment software used.
- 'is\_unmapped\_query" Select unmapped (TRUE) or mapped (FALSE) reads.
- has\_unmapped\_mate Select reads with mapped (FALSE) or unmapped (TRUE) mates.
- is\_minus\_strand Select reads aligned to plus (FALSE) or minus (TRUE) strand.
- is\_mate\_minus\_strand Select reads where mate is aligned to plus (FALSE) or minus (TRUE) strand.
- is\_first\_mate\_read Select reads if they are the first mate (TRUE) or not (FALSE).

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- is\_second\_mate\_read Select reads if they are the second mate (TRUE) or not (FALSE).
- is\_secondary\_alignment Select reads if their alignment status is secondary (TRUE) or not (FALSE). This might be relevant if there are multimapping reads.
- is\_not\_passing\_quality\_controls Select reads that either pass quality controls (FALSE) or that do not (TRUE).
- is\_duplicate Select reads that are unduplicated (FALSE) or duplicated (TRUE). This may represent reads that are PCR or optical duplicates.

#### Value

A DeferredGenomicRanges object

#### See Also

```
Rsamtools::BamFile(),GenomicAlignments::readGAlignments()
```

# Examples

read\_bed

Read a BED or BEDGraph file

# **Description**

This is a lightweight wrapper to the import family of functions defined in rtracklayer.

Read common interval based formats as GRanges.

# Usage

```
read_bed(file, col_names = NULL, genome_info = NULL, overlap_ranges = NULL)
read_bed_graph(
   file,
   col_names = NULL,
   genome_info = NULL,
```

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```
overlap_ranges = NULL
)

read_narrowpeaks(
  file,
  col_names = NULL,
  genome_info = NULL,
  overlap_ranges = NULL
)
```

## **Arguments**

file A path to a file or a connection.

col\_names An optional character vector for including additional columns in file that are

not part of the BED/narrowPeaks specification.

genome\_info An optional character string or a Ranges object that contains information about

the genome build. For example the USSC identifier "hg19" will add build infor-

mation to the returned GRanges.

overlap\_ranges An optional Ranges object. Only the intervals in the file that overlap the Ranges

will be returned.

#### **Details**

This is a lightweight wrapper to the import family of functions defined in **rtracklayer**. The read\_narrowpeaks function parses the ENCODE narrowPeak BED format (see https://genome.ucsc.edu/FAQ/FAQformat.html#format12 for details.). As such the parser expects four additional columns called (corresponding to the narrowPeaks spec):

- signalValue
- pValue
- qValue
- · peak

#### Value

A GRanges object

# See Also

```
rtracklayer::BEDFile()
```

```
test_path <- system.file("tests", package = "rtracklayer")
bed_file <- file.path(test_path, "test.bed")
gr <- read_bed(bed_file)
gr
gr <- read_bed(bed_file, genome_info = "hg19")
gr</pre>
```

44 read\_bigwig

read\_bigwig

Read a BigWig file

## **Description**

Read a BigWig file

# Usage

```
read_bigwig(file, genome_info = NULL, overlap_ranges = NULL)
```

## Arguments

file A path to a file or URL.

genome\_info An optional character string or a Ranges object that contains information about

the genome build. For example the identifier "hg19" will add build information

to the returned GRanges.

overlap\_ranges An optional Ranges object. Only the intervals in the file that overlap the Ranges

will be loaded.

## Value

a GRanges object

## See Also

```
rtracklayer::BigWigFile()
```

```
if (.Platform$0S.type != "windows") {
  test_path <- system.file("tests", package = "rtracklayer")
  bw_file <- file.path(test_path, "test.bw")
  gr <- read_bigwig(bw_file)
  gr
}</pre>
```

read\_gff 45

read\_gff

Read a GFF/GTF/GVT file

## **Description**

This is a lightweight wrapper to the import family of functions defined in **rtracklayer**.

## Usage

```
read_gff(file, col_names = NULL, genome_info = NULL, overlap_ranges = NULL)
read_gff1(file, col_names = NULL, genome_info = NULL, overlap_ranges = NULL)
read_gff2(file, col_names = NULL, genome_info = NULL, overlap_ranges = NULL)
read_gff3(file, col_names = NULL, genome_info = NULL, overlap_ranges = NULL)
```

## **Arguments**

file A path to a file or a connection.

col\_names An optional character vector for parsing specific columns in file that are part

of the GFF specification. These should name either fixed fields, like source or

type, or, for GFF2 and GFF3, any attribute.

genome\_info An optional character string or a Ranges object that contains information about

the genome build. For example the UCSC identifier "hg19" will add build infor-

mation to the returned GRanges.

overlap\_ranges An optional Ranges object. Only the intervals in the file that overlap the Ranges

will be returned.

## Value

A GRanges object a GRanges object

#### See Also

```
rtracklayer::GFFFile()
```

```
test_path <- system.file("tests", package = "rtracklayer")
# gff3
test_gff3 <- file.path(test_path, "genes.gff3")
gr <- read_gff3(test_gff3)
gr
# alternatively with read_gff
gr <- read_gff(test_gff3, genome_info = "hg19")
gr</pre>
```

46 read\_wig

read\_wig

Read a WIG file

# **Description**

This is a lightweight wrapper to the import family of functions defined in **rtracklayer**.

# Usage

```
read_wig(file, genome_info = NULL, overlap_ranges = NULL)
```

# Arguments

file A path to a file or a connection.

genome\_info An optional character string or a Ranges object that contains information about

the genome build. For example the USSC identifier "hg19" will add build infor-

mation to the returned GRanges.

overlap\_ranges An optional Ranges object. Only the intervals in the file that overlap the Ranges

will be returned.

## Value

A GRanges object

A GRanges object

## See Also

```
rtracklayer::WIGFile()
```

```
test_path <- system.file("tests", package = "rtracklayer")
test_wig <- file.path(test_path, "step.wig")
gr <- read_wig(test_wig)
gr
gr <- read_wig(test_wig, genome_info = "hg19")</pre>
```

reduce\_ranges 47

reduce\_ranges

Reduce then aggregate a Ranges object

# **Description**

Reduce then aggregate a Ranges object

# Usage

```
reduce_ranges(.data, min.gapwidth = 1L, ...)
reduce_ranges_directed(.data, min.gapwidth = 1L, ...)
```

# **Arguments**

.data a Ranges object to reduce
 min.gapwidth Ranges separated by a gap of at least min.gapwidth positions are not merged.
 ... Name-value pairs of summary functions.

## Value

a Ranges object with the

```
set.seed(10)
df <- data.frame(start = sample(1:10),</pre>
                 width = 5,
                 seqnames = "seq1",
                 strand = sample(c("+", "-", "*"), 10, replace = TRUE),
                 gc = runif(10)
rng <- as_granges(df)</pre>
rng %>% reduce_ranges()
rng %>% reduce_ranges(gc = mean(gc))
rng %>% reduce_ranges_directed(gc = mean(gc))
rng %>% reduce_ranges_directed(gc = mean(gc), min.gapwidth = 10)
x <- data.frame(start = c(11:13, 2, 7:6),
               width=3,
               id=sample(letters[1:3], 6, replace = TRUE),
               score= sample(1:6))
x <- as_iranges(x)</pre>
x %>% reduce_ranges()
x %>% reduce_ranges(score = sum(score))
x %>% group_by(id) %>% reduce_ranges(score = sum(score))
```

remove\_names

reexports

Objects exported from other packages.

# Description

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

remove\_names

Tools for working with named Ranges

## **Description**

Tools for working with named Ranges

# Usage

```
remove_names(.data)
names_to_column(.data, var = "name")
id_to_column(.data, var = "id")
```

# **Arguments**

. data a Ranges object

var Name of column to use for names

# **Details**

The function names\_to\_column() and id\_to\_column() always places var as the first column in mcols(.data), shifting all other columns to the left. The id\_to\_column() creates a column with sequential row identifiers starting at 1, it will also remove any existing names.

#### Value

Returns a Ranges object with empty names

select.Ranges 49

## **Examples**

```
ir <- IRanges::IRanges(start = 1:3, width = 4, names = c("a", "b", "c"))
remove_names(ir)
ir_noname <- names_to_column(ir)
ir_noname
ir_with_id <- id_to_column(ir)
ir_with_id</pre>
```

select.Ranges

Select metadata columns of the Ranges object by name or position

## **Description**

Select metadata columns of the Ranges object by name or position

#### Usage

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

# **Arguments**

.data a Ranges object

... One or more metadata column names.

that form the core part of the Ranges object.

#### **Details**

Note that by default select only acts on the metadata columns (and will therefore return a Ranges object) if a core component of a Ranges is dropped or selected without the other required components (this includes the seqnames, strand, start, end, width names), then select will throw an error unless .drop\_ranges is set to TRUE.

#### Value

a Ranges object or a tibble

#### See Also

```
dplyr::select()
```

50 set\_width

## **Examples**

```
df <- data.frame(start = 1:10, width = 5, seqnames = "seq1",</pre>
trand = transple(c("+", "-", "*"), 10, replace = true), gc = runif(10), counts = rpois(10, 2))
rng <- as_granges(df)</pre>
select(rng, -gc)
select(rng, gc)
select(rng, counts, gc)
select(rng, 2:1)
select(rng, seqnames, strand, .drop_ranges = TRUE)
```

 $\operatorname{set\_width}$ 

Functional setters for Ranges objects

# **Description**

Functional setters for Ranges objects

# Usage

```
set_width(x, width)
set_start(x, start = 0L)
set_end(x, end = 0L)
set_seqnames(x, seqnames)
set_strand(x, strand)
```

# Arguments

X	a Ranges object
width	integer amount to modify width by
start	integer amount to modify start by
end	integer amount to modify end by
seqnames	update seqnames column
strand	update strand column

## **Details**

These methods are used internally in mutate() to modify core columns in Ranges objects.

# Value

a Ranges object

shift\_left 51

shift_left Shift all coordinates in a genomic interval left or rig downstream	right, upstream or
--	--------------------

# **Description**

Shift all coordinates in a genomic interval left or right, upstream or downstream

# Usage

```
shift_left(x, shift = 0L)
shift_right(x, shift = 0L)
shift_upstream(x, shift = 0L)
shift_downstream(x, shift = 0L)
```

# **Arguments**

x a Ranges 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.

## **Details**

Shifting left or right will ignore any strand information in the Ranges object, while shifting upstream/downstream will shift coordinates on the positive strand left/right and the negative strand right/left. By default, unstranded features are treated as positive. When using shift\_upstream() or shift\_downstream() when the shift argument is indexed by the strandedness of the input ranges.

# Value

a Ranges object with start and end coordinates shifted.

# See Also

```
IRanges::shift(), GenomicRanges::shift()
```

52 slice.Ranges

```
strand = c("+", "+", "-", "-", "+", "*"))) shift_upstream(gr, 5L) shift_downstream(gr, 5L)
```

slice.Ranges

Choose rows by their position

## **Description**

Choose rows by their position

## Usage

```
## S3 method for class 'Ranges'
slice(.data, ..., .preserve = FALSE)

## S3 method for class 'GroupedGenomicRanges'
slice(.data, ..., .preserve = FALSE)

## S3 method for class 'GroupedIntegerRanges'
slice(.data, ..., .preserve = FALSE)
```

## **Arguments**

... Integer row values indicating rows to keep. If .data has been grouped via group\_by(), then the positions are selected within each group.

.preserve when FALSE (the default) the grouping structure is recomputed, otherwise it is kept as is. Currently ignored.

## Value

a GRanges object

stretch 53

```
# slice with group by finds positions within each group
dplyr::slice(by_strand, n())
dplyr::slice(by_strand, which.max(gc))
# if the index is beyond the number of groups slice are ignored
dplyr::slice(by_strand, 1:3)
```

stretch

Stretch a genomic interval

## **Description**

By default, stretch(x) will anchor by the center of a Ranges object. This means that half of the value of extend will be added to the end of the range and the remaining half subtracted from the start of the Range. The other anchors will leave the start/end fixed and stretch the end/start respectively.

## Usage

```
stretch(x, extend)
```

## **Arguments**

Х

a Ranges object, to fix by either the start, end or center of an interval use anchor\_start(x), anchor\_end(x), anchor\_center(x). To fix by strand use

anchor $_3p(x)$  or anchor $_5p(x)$ .

extend

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 x.

## Value

a Ranges object with modified start or end (or both) coordinates

# See Also

```
anchor(), mutate()
```

54 summarise.Ranges

```
stretch(anchor_3p(grng), 10)
stretch(anchor_5p(grng), 10)
```

summarise.Ranges

Reduce multiple values in a Ranges down to a single value

## Description

Reduce multiple values in a Ranges down to a single value

## Usage

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

## **Arguments**

. data a Ranges object

Name-value pairs of summary functions. The name will be the name of the variable in the result. The value should be an expression that will return a value that has length one or length equal to the number of groups.

## **Details**

Creates one or more variables as a S4Vectors::DataFrame() from the input Ranges object. If the ranges object is grouped, there will be a row for each group. Because grouping may remove whether a Ranges object is valid, a DataFrame is always returned.

## Value

```
A S4Vectors::DataFrame()
```

```
df <- data.frame(start = 1:10, width = 5, seqnames = "seq1",
strand = sample(c("+", "-", "*"), 10, replace = TRUE), gc = runif(10))
rng <- as_granges(df)
rng %>% summarise(gc = mean(gc))
rng %>% group_by(strand) %>% summarise(gc = mean(gc))
```

tile\_ranges 55

tile\_ranges

Slide or tile over a Ranges object

## **Description**

Slide or tile over a Ranges object

# Usage

```
tile_ranges(x, width)
slide_ranges(x, width, step)
```

## **Arguments**

x a Ranges object
 width the maximum width of each window/tile (integer vector of length 1)
 step the distance between start position of each sliding window (integer vector of

length 1)

#### **Details**

The tile\_ranges() function paritions a Ranges object x by the given the width over all ranges in x, truncated by the sequence end. The slide\_ranges() function makes sliding windows within each range of x of size width and sliding by step. Both slide\_ranges() and tile\_ranges() return a new Ranges object with a metadata column called "partition" which contains the index of the input range x that a parition belongs to.

## Value

a Ranges object

## See Also

```
GenomicRanges::tile()
```

56 write\_bed

```
# make sliding windows of width 3, moving window with step size of 2
slide_ranges(gr, width = 3, step = 2)
```

write\_bed

Write a BED or BEDGraph file

## **Description**

This is a lightweight wrapper to the export family of functions defined in **rtracklayer**.

## Usage

```
write_bed(x, file, index = FALSE)
write_bed_graph(x, file, index = FALSE)
write_narrowpeaks(x, file)
```

#### **Arguments**

x A GRanges object

file File name, URL or connection specifying a file to write x to. Compressed files

with extensions such as '.gz' are handled automatically. If you want to index the

file with tabix use the index argument.

index Compress and index the output file with bgzf and tabix (default = FALSE). Note

that tabix indexing will sort the data by chromosome and start.

#### Value

The write functions return a BED(Graph)File invisibly

## See Also

```
rtracklayer::BEDFile()
```

```
## Not run:
    test_path <- system.file("tests", package = "rtracklayer")
    bed_file <- file.path(test_path, "test.bed")
    gr <- read_bed(bed_file)
    bed_file_out <- file.path(tempdir(), "new.bed")
    write_bed(gr, bed_file_out)
    read_bed(bed_file_out)
    #' bedgraph
    bg_file <- file.path(test_path, "test.bedGraph")</pre>
```

write\_bigwig 57

```
gr <- read_bed_graph(bg_file)
bg_file_out <- file.path(tempdir(), "new.bg")
write_bed(gr, bg_file_out)
read_bed(bg_file_out)
# narrowpeaks
np_file <- system.file("extdata", "demo.narrowPeak.gz",package="rtracklayer")
gr <- read_narrowpeaks(np_file, genome_info = "hg19")
np_file_out <- file.path(tempdir(), "new.bg")
write_narrowpeaks(gr, np_file_out)
read_narrowpeaks(np_file_out)
## End(Not run)</pre>
```

write\_bigwig

Write a BigWig file

## **Description**

This is a lightweight wrapper to the export family of functions defined in **rtracklayer**.

## Usage

```
write_bigwig(x, file)
```

# **Arguments**

x A GRanges object

file File name, URL or connection specifying a file to write x to. Compressed files with extensions such as '.gz' are handled automatically.

#### Value

The write functions return a BigWigFile invisibly

## See Also

```
rtracklayer::BigWigFile()
```

```
## Not run:
    if (.Platform$0S.type != "windows") {
        test_path <- system.file("tests", package = "rtracklayer")
        bw_file <- file.path(test_path, "test.bw")
        gr <- read_bigwig(bw_file)
        gr
        bw_out <- file.path(tempdir(), "test_out.bw")
        write_bigwig(gr ,bw_out)
        read_bigwig(bw_out)</pre>
```

58 write\_gff

```
}
## End(Not run)
```

write\_gff

Write a GFF(123) file

# Description

This is a lightweight wrapper to the export family of functions defined in **rtracklayer**.

# Usage

```
write_gff(x, file, index = FALSE)
write_gff1(x, file, index = FALSE)
write_gff2(x, file, index = FALSE)
write_gff3(x, file, index = FALSE)
```

# **Arguments**

x A GRanges object

file Path or connection to write to

index If TRUE the output file will be compressed and indexed using bgzf and tabix.

## Value

The write function returns a GFFFile object invisibly

#### See Also

```
rtracklayer::GFFFile()
```

```
## Not run:
  test_path <- system.file("tests", package = "rtracklayer")
  test_gff3 <- file.path(test_path, "genes.gff3")
  gr <- read_gff3(test_gff3)
  out_gff3 <- file.path(tempdir(), "test.gff3")
  write_gff3(gr, out_gff3)
  read_gff3(out_gff3)
## End(Not run)</pre>
```

write\_wig 59

write\_wig

Write a WIG file

# Description

Write a WIG file

## Usage

```
write_wig(x, file)
```

# **Arguments**

x A GRanges object

file File name, URL or connection specifying a file to write x to. Compressed files

with extensions such as '.gz' are handled automatically.

## Value

The write function returns a WIGFile invisibly.

## See Also

```
rtracklayer::WIGFile()
```

%union%

Row-wise set operations on Ranges objects

# Description

Row-wise set operations on Ranges objects

# Usage

```
x %union% y
x %intersect% y
x %setdiff% y
between(x, y)
span(x, y)
```

60 %union%

## **Arguments**

x, y Ranges objects

#### **Details**

Each of these functions acts on the rows between pairs of Ranges object. The function %union%(). will return the entire range between two ranges objects assuming there are no gaps, if you would like to force gaps use span() instead. The function %intersect%() will create a new ranges object with a hit column indicating whether or not the two ranges intersect. The function %setdiff%()will return the ranges for each row in x that are not in the corresponding row of y. The function between() will return the gaps between two ranges.

#### Value

A Ranges object

#### See Also

[IRanges::punion()][IRanges::pintersect()][IRanges::pgap()][IRanges::psetdiff()]

```
x <- as_iranges(data.frame(start = 1:10, width = 5))</pre>
# stretch x by 3 on the right
y <- stretch(anchor_start(x), 3)</pre>
# take the rowwise union
x %union% y
# take the rowwise intersection
x %intersect% y
# asymetric difference
y %setdiff% x
x %setdiff% y
\# if there are gaps between the rows of each range use span
y <- as_iranges(data.frame(start = c(20:15, 2:5),
width = c(10:15,1:4))
# fill in the gaps and take the rowwise union
span(x,y)
# find the gaps
between(x,y)
```

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