

Package ‘gUtils’

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Title R Package Providing Additional Capabilities and Speed for GenomicRanges Operations

Version 0.2.0

Description R package providing additional capabilities and speed for GenomicRanges operations.

Depends R (>= 3.1.0),
GenomicRanges (>= 1.18),
data.table (>= 1.9)

Imports IRanges (>= 2.0),
S4Vectors (>= 0.4),
GenomeInfoDb (>= 1.2),
parallel,
BiocGenerics(>= 0.12),
methods

Suggests BSgenome.Hsapiens.UCSC.hg19,
testthat,
covr,
rtracklayer

License GPL-2

BugReports <http://github.com/mskilab/gUtils/issues>

LazyData true

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R topics documented:

dt2gr	2
gr.chr	3
gr.dice	3
gr.dist	4
gr.DNAase	4
gr.end	5
gr.findoverlaps	5
gr.fix	6
gr.flatten	7
gr.flipstrand	7
gr.genes	8

gr.match	8
gr.mid	9
gr.nochr	9
gr.rand	10
gr.sample	10
gr.simplify	11
gr.start	12
gr.string	12
gr.tile	13
gr.tile.map	13
gr.trim	14
gr2dt	15
grbind	15
grl.hiC	16
grl.in	16
grl.pivot	17
grl.string	17
grl.unlist	18
grl1	18
grl2	18
grlbind	19
ra.overlaps	19
rrbind	20
si	20
si2gr	20
streduce	21
%*%	22

Index 23

dt2gr	<i>Convert data.table to GRanges</i>
-------	--------------------------------------

Description

Takes as input a data.table which must have the following fields: start, end, strand, seqnames. Will throw an error if any one of these is not present. All of the remaining fields are added as metadata to the GRanges.

Usage

```
dt2gr(dt)
```

Arguments

dt	data.table to convert to GRanges
----	----------------------------------

Value

GRanges object of length = nrow(dt)

Examples

```
gr <- dt2gr(data.table(start=c(1,2), seqnames=c("X", "1"), end=c(10,20), strand = c('+', '-')))
```

gr.chr	<i>Prepend "chr" to GRanges seqlevels</i>
--------	---

Description

Prepend "chr" to GRanges seqlevels

Usage

```
gr.chr(gr)
```

Arguments

gr GRanges object to append 'chr' to

Value

Identical GRanges, but with 'chr' prepended to each seqlevel

Examples

```
gr <- gr.chr(GRanges(c(1,"chrX"), IRanges(c(1,2), 1)))
seqnames(gr)
```

gr.dice	<i>Dice up GRanges into width = 1 GRanges spanning the input (warning can produce a very large object)</i>
---------	--

Description

Dice up GRanges into width = 1 GRanges spanning the input (warning can produce a very large object)

Usage

```
gr.dice(gr)
```

Arguments

gr GRanges object to dice

Value

GRangesList where kth element is a diced pile of GRanges from kth input GRanges

Examples

```
gr.dice(GRanges(c(1,4), IRanges(c(10,10),20)))
```

gr.dist	<i>Pairwise distance between two GRanges</i>
---------	--

Description

Computes matrix of pairwise distance between elements of two GRanges objects of length n and m.

Usage

```
gr.dist(gr1, gr2 = NULL, ignore.strand = FALSE, ...)
```

Arguments

gr1	First GRanges
gr2	Second GRanges
ignore.strand	Don't required elements be on same strand to avoid NA [FALSE]
...	Additional arguments to be supplied to GenomicRanges::distance

Details

Distances are computed as follows:

- NA for ranges on different seqnames
- 0 for overlapping ranges
- min(abs(end1-end2), abs(end1-start2), abs(start1-end2), abs(start1-end1),) for all others

If only gr1 is provided, then will return n x n matrix of gr1 vs itself

If max.dist = TRUE, then will replace min with max above

Value

N by M matrix with the pairwise distances, with gr1 on rows and gr2 on cols

gr.DNAase	<i>DNAaseI hypersensitivity sites for hg19</i>
-----------	--

Description

DNAaseI hypersensitivity sites from UCSC Table Browser hg19, subsampled to 10,000 sites

Format

GRanges

gr.end	<i>Get the right ends of a GRanges</i>
--------	--

Description

Alternative to `GenomicRanges::flank` that will provide end positions **within** intervals

Usage

```
gr.end(x, width = 1, force = FALSE, ignore.strand = TRUE, clip = TRUE)
```

Arguments

x	GRanges object to operate on
width	Specify subranges of greater width including the start of the range. [1]
force	Allows returned GRanges to have ranges outside of its Seqinfo bounds. [FALSE]
ignore.strand	If set to FALSE, will extend '-' strands from the other direction. [TRUE]
clip	Trims returned GRanges so that it does not extend beyond bounds of the input GRanges. [TRUE]

Value

GRanges object of width = width ranges representing end of each genomic range in the input.

Examples

```
gr.end(gr.DNAase, width=200, clip=TRUE)
```

gr.findoverlaps	<i>Wrapper to GenomicRanges::findOverlaps with added functionality</i>
-----------------	--

Description

Returns GRanges of matches with two additional fields:

`$query.id` - index of matching query `$subject.id` - index of matching subject

Optional "by" field is a character scalar that specifies a metadata column present in both query and subject that will be used to additionally restrict matches, i.e. to pairs of ranges that overlap and also have the same values of their "by" fields

Usage

```
gr.findoverlaps(query, subject, ignore.strand = TRUE, first = FALSE,
  qcol = NULL, scol = NULL, type = "any", by = NULL,
  return.type = "same", ...)
```

Arguments

query	Query GRanges pile
subject	Subject GRanges pile
ignore.strand	Don't consider strand information during overlaps. [TRUE]
first	Restrict to only the first match of the subject (default is to return all matches). [FALSE]
qcol	character vector of query meta-data columns to add to results
scol	character vector of subject meta-data columns to add to results
type	type argument as defined by <code>IRanges::findOverlaps("any", "start", "end", "within", "equal")</code> . ["any"]
by	Meta-data column to consider when performing overlaps [NULL]
return.type	Select data format to return (supplied as character): "same", "data.table", "GRanges". ["same"]
...	Additional arguments sent to <code>IRanges::findOverlaps</code> .

Value

GRanges pile of the intersection regions, with `query.id` and `subject.id` marking sources

gr.fix	"Fixes" seqlengths / seqlevels
--------	--------------------------------

Description

If "genome" not specified will replace NA seqlengths in GRanges to reflect largest coordinate per seqlevel and removes all NA seqlevels after this fix.

Usage

```
gr.fix(gr, genome = NULL, gname = NULL, drop = FALSE)
```

Arguments

gr	GRanges object to fix
genome	Genome to fix to: Seqinfo, BSgenome, GRanges (w/seqlengths), GRangesList (w/seqlengths)
gname	Name of the genome (optional, just appends to Seqinfo of the output) [NULL]
drop	Remove ranges that are not present in the supplied genome [FALSE]

Details

if "genome" defined (i.e. as Seqinfo object, or a BSgenome, GRanges, GRangesList object with populated seqlengths), then will replace seqlengths in gr with those for that genome

Value

GRanges pile with the fixed Seqinfo

gr.flatten	<i>Lay ranges end-to-end onto a derivate "chromosome"</i>
------------	---

Description

Takes pile of GRanges and returns into a data.frame with `nrow = length(gr)` with each representing the corresponding input range superimposed onto a single "flattened" chromosome, with ranges laid end-to-end

Usage

```
gr.flatten(gr, gap = 0)
```

Arguments

gr	GRanges to flatten
gap	Number of bases between ranges on the new chromosome [0]

Value

data.frame with start and end coordinates, and all of the original metadata

gr.flipstrand	<i>Flip strand on GRanges</i>
---------------	-------------------------------

Description

Flip strand on GRanges

Usage

```
gr.flipstrand(gr)
```

Arguments

gr	GRanges pile with strands to be flipped
----	---

Value

GRanges with flipped strands (+ to -, * to *, - to *)

Examples

```
gr.flipstrand(GRanges(1, IRanges(c(10,10,10),20), strand=c("+","*","-")))
```

gr.genes	<i>RefSeq genes from UCSC Table Browser hg19, subsampled to 10,000 genes</i>
----------	--

Description

RefSeq genes from UCSC Table Browser hg19, subsampled to 10,000 genes

Format

GRanges

gr.match	<i>Alternative GenomicRanges::match that accepts "by" argument and data.table inputs</i>
----------	--

Description

Wrapper to GenomicRanges::match (uses [gr.findoverlaps](#))

Usage

```
gr.match(query, subject, ...)
```

Arguments

query	Query GRanges pile
subject	Subject GRanges pile
...	Additional arguments to be passed along to gr.findoverlaps .

Value

Vector of length = length(query) with subject indices of *first* subject in query, or NA if none found. This behavior is different from [gr.findoverlaps](#), which will return *all* indices of subject in query (in the case of one query overlaps with multiple subject) ... = additional args for findOverlaps (IRanges version)

gr.mid	<i>Get the midpoints of GRanges ranges</i>
--------	--

Description

Get the midpoints of GRanges ranges

Usage

```
gr.mid(x)
```

Arguments

x	GRanges object to operate on
---	------------------------------

Value

GRanges of the midpoint, calculated from `floor(width(x)/2)`

Examples

```
gr.mid(GRanges(1, IRanges(1000,2000), seqinfo=Seqinfo("1", 2000)))
```

gr.nochr	<i>Remove chr prefix from GRanges seqlevels</i>
----------	---

Description

Remove chr prefix from GRanges seqlevels

Usage

```
gr.nochr(gr)
```

Arguments

gr	GRanges with chr seqlevel prefixes
----	------------------------------------

Value

GRanges without chr seqlevel prefixes

gr.rand	<i>Generate random GRanges on genome</i>
---------	--

Description

Randomly generates non-overlapping GRanges with supplied widths on supplied genome. Seed can be supplied with `set.seed`

Usage

```
gr.rand(w, genome)
```

Arguments

w	Vector of widths (length of w determines length of output)
genome	Genome which can be a GRanges, GRangesList, or Seqinfo object. Default is "hg19" from the BSGenome package.

Value

GRanges with random intervals on the specified "chromosomes"

Note

This function is currently quite slow, needs optimization

Examples

```
## Generate a single random interval of width 10, on "chr" of length 1000
gr.rand(10, Seqinfo("1", 1000))
## Generate 5 non-overlapping regions of width 10 on hg19
library(BSGenome.Hsapiens.UCSC.hg19)
gr.rand(rep(10,5), Hsapiens)
```

gr.sample	<i>Randomly sample GRanges intervals within territory</i>
-----------	---

Description

Samples k intervals of length "len" from a pile of GRanges.

- If k is a scalar then will (uniformly) select k intervals from the summed territory of GRanges
- If k is a vector of length(gr) then will uniformly select k intervals from each.

Usage

```
gr.sample(gr, k, len = 100, replace = TRUE)
```

Arguments

gr	GRanges object defining the territory to sample from
k	Number of ranges to sample
len	Length of the GRanges element to produce [100]
replace	If TRUE, will bootstrap, otherwise will sample without replacement. [TRUE]

Value

GRanges of max length sum(k) [if k is vector] or k*length(gr) (if k is scalar) with labels indicating the originating range.

Note

This is different from `GenomicRanges::sample` function, which just samples from a pile of GRanges

Examples

```
## sample 5 \code{GRanges} of length 10 each from territory of RefSeq genes
gr.sample(reduce(gr.genes), k=5, len=10)
```

gr.simplify	<i>Simplify granges by collapsing all non-overlapping adjacent ranges that share a given "field" value (adjacent == adjacent in the input GRanges object)</i>
-------------	---

Description

Simplify granges by collapsing all non-overlapping adjacent ranges that share a given "field" value (adjacent == adjacent in the input GRanges object)

Usage

```
gr.simplify(gr, field = NULL, include.val = TRUE, split = FALSE,
            pad = 1)
```

Arguments

gr	takes in gr or grl
field	character scalar, corresponding to value field of gr. [NULL]
include.val	scalar logical, will include in out gr values field of first matching record in input gr. [TRUE]
split	Split the output into GRangesList split by "field". [FALSE]
pad	Pad ranges by this amount before doing merge. [1], which merges contiguous but non-overlapping ranges.
val	[NULL]

Value

Simplified GRanges with "field" populated with uniquely contiguous values

gr.start	<i>Get GRanges corresponding to beginning of range</i>
----------	--

Description

Get GRanges corresponding to beginning of range

Usage

```
gr.start(x, width = 1, force = FALSE, ignore.strand = TRUE,
        clip = FALSE)
```

Arguments

x	GRanges object to operate on
width	[default = 1] Specify subranges of greater width including the start of the range.
force	[default = F] Allows returned GRanges to have ranges outside of its Seqinfo bounds.
ignore.strand	If set to FALSE, will extend '-' strands from the other direction [TRUE].
clip	[default = F] Trims returned GRanges so that it does not extend beyond bounds of the input GRanges

Value

GRanges object of width 1 ranges representing start of each genomic range in the input.

Examples

```
gr.start(gr.DNAase, width=200)
gr.start(gr.DNAase, width=200, clip=TRUE)
```

gr.string	<i>Return UCSC style interval string corresponding to GRanges pile (ie chr:start-end)</i>
-----------	---

Description

Return UCSC style interval string corresponding to GRanges pile (ie chr:start-end)

Usage

```
gr.string(gr, add.chr = FALSE, mb = FALSE, round = 3, other.cols = c())
```

Arguments

gr	GRanges pile to get intervals from
add.chr	Prepend seqnames with "chr" [FALSE]
mb	Round to the nearest megabase [FALSE]
round	If mb supplied, how many digits to round to. [3]
other.cols	Names of additional mcols fields to add to the string (seperated by ";")

Examples

```
gr.string(gr.genes, other.cols = c("name", "name2"))
```

gr.tile	<i>Tile ranges across GRanges</i>
---------	-----------------------------------

Description

Tiles interval (or whole genome) with segments of \leq specified width.

Usage

```
gr.tile(gr, w = 1000)
```

Arguments

gr	GRanges, seqlengths or Seqinfo range to tile. If has GRanges has overlaps, will reduce first.
w	Width of each tile

Examples

```
## 10 tiles of width 10
gr1 <- gr.tile(GRanges(1, IRanges(1,100)), w=10)
## make them overlap each other by 5
gr1 + 5
```

gr.tile.map	<i>gr.tile.map</i>
-------------	--------------------

Description

Given two tilings of the genome (e.g. at different resolution) query and subject outputs a length(query) list whose items are integer vectors of indices in subject overlapping that overlap that query (strand non-specific)

Usage

```
gr.tile.map(query, subject, verbose = FALSE)
```

Arguments

query	Query GRanges pile, perhaps created from some tile (e.g. gr.tile), and assumed to have no gaps
subject	Subject GRanges pile, perhaps created from some tile (e.g. gr.tile), and assumed to have no gaps
verbose	Increase the verbosity of the output

Value

list of length = length(query), where each element i is a vector of indices in subject that overlaps element i of query

Note

Assumes that input query and subject have no gaps (including at end) or overlaps, i.e. ignores end() coordinates and only uses "starts"

gr.trim	<i>Trims pile of GRanges relative to the specified <local> coordinates of each range</i>
---------	--

Description

Example: GRanges with genomic coordinates 1:1,000,000-1,001,000 can get the first 20 and last 50 bases trimmed off with start = 20, end = 950. if end is larger than the width of the corresponding gr, then the corresponding output will only have end(gr) as its coordinate.

Usage

```
gr.trim(gr, starts = 1, ends = 1)
```

Arguments

gr	GRanges to trim
starts	Number of bases to trim off of the front[1]
ends	Number of bases to trim off of the back[1]

Details

This is a role not currently provided by the standard GenomicRanges functions (e.g. shift, reduce, restrict, shift, resize, flank)

Examples

```
## trim the first 20 and last 50 bases
gr.trim(GRanges(1, IRanges(1e6, width=1000)), starts=20, ends=950)
## return value: GRanges on 1:1,000,019-1,000,949
```

gr2dt	<i>Converts GRanges to data.table</i>
-------	---------------------------------------

Description

Converts GRanges to data.table

Usage

```
gr2dt(gr)
```

Arguments

gr GRanges pile to convert to data.table

Value

data.table with seqnames, start, end, width, strand and all of the meta data. Width is end-inclusive (e.g. [6,7] width = 2)

Examples

```
gr2dt(gr.genes)
```

grbind	<i>Concatenate GRanges, robust to different mcols</i>
--------	---

Description

Concatenates GRanges objects, taking the union of their features if they have non-overlapping features

Usage

```
grbind(x, ...)
```

Arguments

x First GRanges
... additional GRanges

Value

Concatenated GRanges grbind(gr.genes, gr.DNAase)

Note

Does not fill in the Seqinfo for the output GRanges

grl.hiC	<i>HiC data for chr14 from Lieberman-Aiden 2009 (in hg19), subsampled to 10,000 interactions</i>
---------	--

Description

HiC data for chr14 from Lieberman-Aiden 2009 (in hg19), subsampled to 10,000 interactions

Format

GRangesList

grl.in	<i>Check intersection of GRangesList with windows on genome</i>
--------	---

Description

Like only if the ranges in grl[i] intersect «all», «some», «only» windows in the subject

Usage

```
grl.in(grl, windows, some = FALSE, only = FALSE, ...)
```

Arguments

grl	GRangesList object to query for membership in windows
windows	GRanges pile of windows
some	Will return TRUE for GRangesList elements that intersect at least on window range [FALSE]
only	Will return TRUE for GRangesList elements only if there are no elements of query that fail to intersect with windows [FALSE]
...	Additional parameters to be passed on to GenomicRanges::findOverlaps

Details

eg can use to identify read pairs whose ends are contained inside two genes)

grl.pivot	<i>Pivot a GRangesList, inverting "x" and "y"</i>
-----------	---

Description

"Pivots" grl object "x" by returning a new grl "y" whose kth item is gr object of ranges x[[i]][k] for all i in 1:length(x) e.g. If length(grl) is 50 and length of each GRanges element inside is 2, then grl.pivot will produce a length 3 GRangesList with 50 elements per GRanges

Usage

```
grl.pivot(x)
```

Arguments

x	GRangesList object to pivot
---	-----------------------------

Details

Assumes all grs in "x" are of equal length

Examples

```
grl.pivot(grl.hiC)
```

grl.string	<i>Create string representation of GRangesList</i>
------------	--

Description

Return ucsc style interval string corresponding to each GRanges in the GRangesList. One line per per GRangesList item. GRanges elements themselves are separated by sep

Usage

```
grl.string(grl, mb = FALSE, sep = ",", ...)
```

Arguments

grl	GRangesList to convert to string vector
mb	Will return as MB and round to "round" [FALSE]
sep	Character to separate single GRanges ranges [,]
...	Additional arguments to be passed to gr.string

Value

Character vector where each element is a GRanges pile corresponding to a single GRangesList element

Examples

```
grl.string(grl.hiC, mb=TRUE)
```

grl.unlist	<i>Robust unlisting of GRangesList that keeps track of origin</i>
------------	---

Description

Does a "nice" unlist of a GRangesList object adding a field `grl.ix` denoting which element of the GRangesList each GRanges corresponds to and a field `grl.iix` which saves the (local) index that that gr was in its corresponding GRangesList item

Usage

```
grl.unlist(grl)
```

Arguments

grl	GRangesList object to unlist
-----	------------------------------

Details

In this way, `grl.unlist` is reversible, while `BiocGenerics::unlist` is not.

Value

GRanges with added metadata fields `grl.ix` and `grl.iix`.

Examples

```
grl.unlist(grl.hiC)
```

grl1	<i>Fake rearrangement data (set 1)</i>
------	--

Description

Fake rearrangement data (set 1)

Format

GRangesList

grl2	<i>Fake rearrangement data (set 2)</i>
------	--

Description

Fake rearrangement data (set 2)

Format

GRangesList

grlbind	<i>Concatenate GRangesList objects</i>
---------	--

Description

Concatenates GRangesList objects taking the union of their mcols features if they have non-overlapping features

Usage

```
grlbind(...)
```

Arguments

... Any number of GRangesList to concatenate together

Value

Concatenated GRangesList with NA filled in for mcols fields that are non-overlapping

Examples

```
## Concatenate
grl.hiC2 <- grl.hiC[1:20]
mcols(grl.hiC2)$test = 1
grlbind(grl.hiC2, grl.hiC[1:30])
```

ra.overlaps	<i>Find overlaps between rearrangements represented by GRangesList objects</i>
-------------	--

Description

Determines overlaps between two piles of rearrangement junctions, each GRangesLists of signed locus pairs, against each other. returning a sparseMatrix that is T at entry ij if junction i overlaps junction j.

Usage

```
ra.overlaps(ra1, ra2, pad = 0, ...)
```

Arguments

ra1	GRangesList of pairs of signed ranges representing a rearrangement set
ra2	GRangesList of pairs of signed ranges representing a rearrangement set
pad	Pad each breakpoint when considering overlaps. [0]
...	Additional arguments to be sent to findOverlaps (e.g. ignore.strand)

Value

matrix with the indices of ra1 that overlap with ra2 and vice-versa

rrbind	<i>Improved rbind for intersecting/union columns of data.frames or data.tables</i>
--------	--

Description

Like rbind, but takes the intersecting columns of the data.

Usage

```
rrbind(..., union = TRUE, as.data.table = FALSE)
```

Arguments

...	Any number of data.frame or data.table objects
union	Take union of columns (and put NA's for columns of df1 not in df2 and vice versa). [TRUE]
as.data.table	Return the binded data as a data.table. [FALSE]

Value

data.frame or data.table of the rbind operation

si	<i>Seqinfo object for hg19</i>
----	--------------------------------

Description

Seqinfo object for hg19

Format

Seqinfo

si2gr	<i>Create GRanges from Seqinfo or BSgenome</i>
-------	--

Description

Creates a genomic ranges from seqinfo object ie a pile of ranges spanning the genome

Usage

```
si2gr(si, strip.empty = FALSE)
```

Arguments

si	Seqinfo object or a BSgenome genome
strip.empty	Don't know. [FALSE]

Value

GRanges representing the range of the input genome

Examples

```
## Not run: library(BSgenome.Hsapiens.UCSC.hg19); si2gr(Hsapiens)
```

streduce	<i>Reduce GRanges and GRangesList to minimal footprint</i>
----------	--

Description

Shortcut for `reduce(sort(gr.stripstrand(unlist(x))))`

Usage

```
streduce(gr, pad = 0, sort = TRUE)
```

Arguments

gr	GRanges or GRangesList
pad	Expand the input data before reducing. [0]
sort	Flag to sort the output. [TRUE]

Value

GRanges object with no strand information, representing a minimal footprint

Examples

```
streduce(gr1.hiC, pad=10)  
streduce(gr.genes, pad=1000)
```

%%	<i>Metadata join with coordinates as keys (wrapper to gr.findoverlaps)</i>
----	--

Description

Shortcut for `gr.findoverlaps` with `qcol` and `scol` filled in with all the query and subject metadata names. This function is useful for piping `GRanges` operations together. Another way to think of join of the metadata, with genomic coordinates as the keys.

Example usage:

x

Usage

```
## S4 method for signature 'GRanges'
x %%% y
```

Arguments

x	GRanges
y	GRanges

Value

GRanges containing every pairwise intersection of ranges in x and y with a join of the corresponding metadata

Author(s)

Marcin Imielinski

Examples

```
gr.genes %%% gr.DNAase
```

Index

*Topic **data**

gr.DNAase, [4](#)

gr.genes, [8](#)

grl.hiC, [16](#)

grl1, [18](#)

grl2, [18](#)

si, [20](#)

%%%, GRanges-method (%%%), [22](#)

%%%, [22](#)

dt2gr, [2](#)

findOverlaps, [19](#)

gr.chr, [3](#)

gr.dice, [3](#)

gr.dist, [4](#)

gr.DNAase, [4](#)

gr.end, [5](#)

gr.findoverlaps, [5](#), [8](#), [22](#)

gr.fix, [6](#)

gr.flatten, [7](#)

gr.flipstrand, [7](#)

gr.genes, [8](#)

gr.match, [8](#)

gr.mid, [9](#)

gr.nochr, [9](#)

gr.rand, [10](#)

gr.sample, [10](#)

gr.simplify, [11](#)

gr.start, [12](#)

gr.string, [12](#)

gr.tile, [13](#)

gr.tile.map, [13](#)

gr.trim, [14](#)

gr2dt, [15](#)

grbind, [15](#)

grl.hiC, [16](#)

grl.in, [16](#)

grl.pivot, [17](#)

grl.string, [17](#)

grl.unlist, [18](#)

grl1, [18](#)

grl2, [18](#)

grlbind, [19](#)

ra.overlaps, [19](#)

rrbind, [20](#)

si, [20](#)

si2gr, [20](#)

streduce, [21](#)