Package ‘RaggedExperiment’

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Title Representation of Sparse Experiments and Assays Across Samples

Version 1.0.0

Description This package provides a flexible representation of copy number, mutation, and other data that fit into the ragged array schema for genomic location data. The basic representation of such data provides a rectangular flat table interface to the user with range information in the rows and samples/specimen in the columns.

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biocViews Infrastructure, DataRepresentation

BugReports https://github.com/Bioconductor/RaggedExperiment/issues

VignetteBuilder knitr

Depends R (>= 3.4.0), GenomicRanges

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RaggedExperiment-package

RaggedExperiment: Range-based data representation package

Description

RaggedExperiment allows the user to represent, copy number, mutation, and other types of range-based data formats where optional information about samples can be provided. At the backbone of this package is the GRangesList class. The RaggedExperiment class uses this representation and presents the data in a couple of different ways:

- rowRanges
- colData

The rowRanges method will return the internal GRangesList representation of the dataset. A distinction between the GRangesList and the RaggedExperiment classes is that the RaggedExperiment class allows for ragged ranges, meaning that there may be a different number of ranges for each sample.

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See Also

Useful links:

- Report bugs at https://github.com/Bioconductor/RaggedExperiment/issues

assay-functions

Create simplified representation of ragged assay data.

Description

These methods transform assay() from the default (i.e., sparseAssay()) representation to various forms of more dense representation. compactAssay() collapses identical ranges across samples into a single row. disjoinAssay() creates disjoint (non-overlapping) regions, simplifies values within each sample in a user-specified manner, and returns a matrix of disjoint regions x samples.

This method transforms assay() from the default (i.e., sparseAssay()) representation to a reduced representation summarizing each original range overlapping ranges in query. Reduction in each cell can be tailored to individual needs using the simplify argument.
assay-functions

Usage

sparseAssay(x, i = 1, withDimnames = TRUE, background = NA)

compactAssay(x, i = 1, withDimnames = TRUE, background = NA)

disjoinAssay(x, simplify, i = 1, withDimnames = TRUE, background = NA)

qreduceAssay(x, query, simplify, i = 1, withDimnames = TRUE, background = NA)

Arguments

x   A RaggedExperiment object

i   integer(1) or character(1) name of assay to be transformed.

withDimnames logical(1) include dimnames on the returned matrix. When there are no explicit rownames, these are manufactured with as.character(rowRanges(x)); rownames are always manufactured for compactAssay() and disjoinAssay().

background A value (default NA) for the returned matrix after *Assay operations

simplify disjoinAssay: A function operating on a *List, where the elements of the list are all within-sample assay values from ranges overlapping each disjoint range. For instance, to use the simplify=mean of overlapping ranges, where ranges are characterized by integer-valued scores, the entries are calculated as

\[
\begin{align*}
\text{original:} & \quad |-----------| \\
\text{b} & \quad |----------| \\
\text{a a, b b} & \\
\text{disjoint:} & \quad |----|------|---| \\
\end{align*}
\]

\[
\text{values <- IntegerList(a, c(a, b), b) simplify(values)}
\]

qreduceAssay: A function accepting arguments score, range, and qrange:

• score A *List, where each list element corresponds to a cell in the matrix to be returned by qreduceAssay. Vector elements correspond to ranges overlapping query. The *List objects support many vectorized mathematical operations, so simplify can be implemented efficiently.

• range A GRangesList instance, 'parallel' to score. Each element of the list corresponds to a cell in the matrix to be returned by qreduceAssay. Each range in the element corresponds to the range for which the score element applies.

• qrange A GRanges instance with the same length as score, providing the query range to which the corresponding scores apply.

query GRanges providing regions over which reduction is to occur.

Value

sparseAssay(): A matrix() with dimensions dim(x). Elements contain the assay value for the ith range and jth sample.
compactAssay(): Samples with identical range are placed in the same row. Non-disjoint ranges are NOT collapsed.

disjoinAssay(): A matrix with number of rows equal to number of disjoint ranges across all samples. Elements of the matrix are summarized by applying simplify() to assay values of overlapping ranges.

qreduceAssay(): A matrix() with dimensions length(query) x ncol(x). Elements contain assay values for the ith query range and jth sample, summarized according to the function simplify.

Examples

```r
x <- RaggedExperiment(GRangesList(
  GRanges(c("A:1-3", "A:4-5", "A:10-15"), score=1:3),
  GRanges(c("A:4-5", "B:1-3"), score=4:5)
))
query <- GRanges(c("A:1-2", "A:4-5", "B:1-5"))

weightedmean <- function(scores, ranges, qranges)
## weighted average score per query range
  sum(scores * width(ranges)) / sum(width(ranges))
qreduceAssay(x, query, weightedmean)

## Not run:
## Extended example: non-silent mutations, summarized by genic region
suppressPackageStartupMessages({
  library(TxDb.Hsapiens.UCSC.hg19.knownGene)
  library(org.Hs.eg.db)
  library(GenomeInfoDb)
  library(MultiAssayExperiment)
})

## TCGA Multi-assay experiment to RaggedExperiment
url <- "http://s3.amazonaws.com/multiassayexperiments/accMAEO.rds"
## download.file(url, fl <- tempfile())
## fl <- "accMAEO.rds"
mae <- readRDS(fl)[, , c("RNASeq2GeneNorm", "CNASNP", "Mutations")]

## genomic coordinates
gn <- genes(TxDb.Hsapiens.UCSC.hg19.knownGene)
gn <- keepStandardChromosomes(granges(gn), pruning.mode="coarse")
seqlevelsStyle(gn) <- "NCBI"

## reduce mutations, marking any genomic range with non-silent mutation as FALSE
nonsilent <- function(scores, ranges, qranges)
  any(scores != "Silent")
re <- as(mae["Mutations"], "RaggedExperiment")
mutations <- qreduceAssay(re, gn, nonsilent, "Variant_Classification")

## reduce copy number
re <- as(mae["CNASNP"], "RaggedExperiment")
```
RaggedExperiment-class

The `RaggedExperiment` class is a container for storing range-based data, including but not limited to copy number data, and mutation data. It can store a collection of `GRanges` objects, as it is derived from the `GenomicRangesList`.

Usage

```r
RaggedExperiment(..., colData = DataFrame())
## S4 method for signature 'RaggedExperiment'
rowRanges(x, ...)
## S4 method for signature 'RaggedExperiment'
dim(x)
## S4 method for signature 'RaggedExperiment'
dimnames(x)
## S4 replacement method for signature 'RaggedExperiment,list'
dimnames(x) <- value
## S4 method for signature 'RaggedExperiment'
colData(x, ...)
## S4 method for signature 'RaggedExperiment,missing'
assay(x, i, ...)
## S4 method for signature 'RaggedExperiment,ANY'
assay(x, i, ..., withDimnames = TRUE)
## S4 method for signature 'RaggedExperiment'
assays(x, ..., withDimnames = TRUE)
## S4 method for signature 'RaggedExperiment'
assayNames(x, ...)
## S4 method for signature 'RaggedExperiment'
show(object)
## S4 method for signature 'RaggedExperiment,ANY,ANY,ANY'
x[i, j, ..., drop = TRUE]
```
## S4 method for signature 'RaggedExperiment,Vector'
overlapsAny(query, subject, maxgap = 0L,
minoverlap = 1L, type = c("any", "start", "end", "within", "equal"), ...)

## S4 method for signature 'RaggedExperiment,Vector'
subsetByOverlaps(query, subject,
maxgap = 0L, minoverlap = 1L, type = c("any", "start", "end", "within", "equal"), invert = FALSE, ...)

### Arguments

... Constructor: GRanges, list of GRanges, or GRangesList OR assay: Additional arguments for assay. See details for more information.

colData A DataFrame describing samples. Length of rowRanges must equal the number of rows in colData

x A RaggedExperiment object.

value A list of dimension names

i logical(1), integer(1), or character(1) indicating the assay to be reported. For [], i can be any supported Vector object, e.g., GRanges.

withDimnames logical (default TRUE) whether to use dimension names in the resulting object

object A RaggedExperiment object.

j integer(), character(), or logical() index selecting columns from RaggedExperiment

drop logical (default TRUE) whether to drop empty samples

query A RaggedExperiment instance.

subject Each of them can be a Ranges, Views, RangesList, ViewsList, or RangedData object. In addition, if subject is a Ranges object, query can be an integer vector to be converted to length-one ranges.

If query is a RangesList or RangedData, subject must be a RangesList or RangedData. If both lists have names, each element from the subject is paired with the element from the query with the matching name, if any. Otherwise, elements are paired by position. The overlap is then computed between the pairs as described below.

If subject is omitted, query is queried against itself. In this case, and only this case, the drop.self and drop.redundant arguments are allowed. By default, the result will contain hits for each range against itself, and if there is a hit from A to B, there is also a hit for B to A. If drop.self is TRUE, all self matches are dropped. If drop.redundant is TRUE, only one of A->B and B->A is returned.

maxgap Intervals with a separation of maxgap or less and a minimum of minoverlap overlapping positions, allowing for maxgap, are considered to be overlapping. maxgap should be a scalar, non-negative, integer. minoverlap should be a scalar, positive integer.

minoverlap Intervals with a separation of maxgap or less and a minimum of minoverlap overlapping positions, allowing for maxgap, are considered to be overlapping. maxgap should be a scalar, non-negative, integer. minoverlap should be a scalar, positive integer.

type By default, any overlap is accepted. By specifying the type parameter, one can select for specific types of overlap. The types correspond to operations in
Allen’s Interval Algebra (see references). If type is start or end, the intervals are required to have matching starts or ends, respectively. While this operation seems trivial, the naive implementation using outer would be much less efficient. Specifying equal as the type returns the intersection of the start and end matches. If type is within, the query interval must be wholly contained within the subject interval. Note that all matches must additionally satisfy the minoverlap constraint described above.

The maxgap parameter has special meaning with the special overlap types. For start, end, and equal, it specifies the maximum difference in the starts, ends or both, respectively. For within, it is the maximum amount by which the subject may be wider than the query.

invert If TRUE, keep only the query ranges that do not overlap the subject.

Value
constructor returns a RaggedExperiment object
‘rowRanges’ returns a GRanges object summarizing ranges corresponding to assay() rows.
‘assays’ returns a SimpleList
overlapsAny returns a logical vector of length equal to the number of rows in the query; TRUE when the copy number region overlaps the subject.
subsetByOverlaps returns a RaggedExperiment containing only copy number regions overlapping subject.

Methods (by generic)
• rowRanges: rowRanges accessor
• dim: get dimensions (number of sample-specific row ranges by number of samples)
• dimnames: get row (sample-specific) range names and sample names
• dimnames<-. set row (sample-specific) range names and sample names
• colData: get column data
• assay: assay missing method uses first metadata column
• assay: assay numeric method.
• assays: assays
• assayNames: names in each assay
• show: show method
• [: Subset a RaggedExperiment object
• overlapsAny: Determine whether copy number ranges defined by query overlap ranges of subject.
• subsetByOverlaps: Subset the RaggedExperiment to contain only copy number ranges overlapping ranges of subject.

Constructors
RaggedExperiment(..., colData=DataFrame()): Creates a RaggedExperiment object using multiple GRanges objects or a list of GRanges objects. Additional column data may be provided as a DataFrame object.
Subsetting

In the following, 'x' represents a RaggedExperiment object:

\[
x[i, j]: \text{Get ranges or elements (i and j, respectively) with optional metadata columns where i or j can be missing, an NA-free logical, numeric, or character vector.}
\]

Coercion

Coercion possible from `RangedRaggedAssay` to RaggedExperiment. Here object represents a `RangedRaggedAssay`: `as(object, "RaggedExperiment")`

Examples

```r
## Create an empty RaggedExperiment instance
re0 <- RaggedExperiment()
re0

## Create a couple of GRanges objects with row ranges names
sample1 <- GRanges(
  c(a = "chr1:1-10:-", b = "chr1:11-18:+"),
  score = 1:2)
sample2 <- GRanges(
  c(c = "chr2:1-10:-", d = "chr2:11-18:+"),
  score = 3:4)

## Include column data
colDat <- DataFrame(id = 1:2)

## Create a RaggedExperiment object from a couple of GRanges
re1 <- RaggedExperiment(sample1 = sample1, sample2 = sample2, colData = colDat)
re1

## With list of GRanges
lgr <- list(sample1 = sample1, sample2 = sample2)

## Create a RaggedExperiment from a list of GRanges
re2 <- RaggedExperiment(lgr, colData = colDat)
grl <- GRangesList(sample1 = sample1, sample2 = sample2)

## Create a RaggedExperiment from a GRangesList
re3 <- RaggedExperiment(grl, colData = colDat)

## Subset a RaggedExperiment
assay(re3[c(1, 3),])
subsetByOverlaps(re3, GRanges("chr1:1-5"))  # by ranges
```

sparseSummarizedExperiment

Create `SummarizedExperiment` representations by transforming ragged assays to rectangular form.
sparseSummarizedExperiment

Description

These methods transform RaggedExperiment objects to similar SummarizedExperiment objects. They do so by transforming assay data to more rectangular representations, following the rules outlined for similarly names transformations sparseAssay(), compactAssay(), disjoinAssay(), and qreduceAssay(). Because of the complexity of the transformation, it usually only makes sense transform RaggedExperiment objects with a single assay; this is currently enforced at time of coercion.

Usage

sparseSummarizedExperiment(x, i = 1, withDimnames = TRUE)
compactSummarizedExperiment(x, i = 1L, withDimnames = TRUE)
disjoinSummarizedExperiment(x, simplify, i = 1L, withDimnames = TRUE)
qreduceSummarizedExperiment(x, query, simplify, i = 1L, withDimnames = TRUE)

Arguments

x RaggedExperiment
i integer(1), character(1), or logical() selecting the assay to be transformed.
withDimnames logical(1) default TRUE, propagate dimnames to SummarizedExperiment.
simplify function of 1 (for disjoinSummarizedExperiment) or 3 (for qreduceSummarizedExperiment) arguments, used to transform assays. See assay-functions.
query GRanges providing regions over which reduction is to occur.

Value

All functions return RangedSummarizedExperiment.
sparseSummarizedExperiment has rowRanges() identical to the row ranges of x, and assay() data as sparseAssay(). This is very space-inefficient representation of ragged data.
compactSummarizedExperiment has rowRanges() identical to the row ranges of x, and assay() data as compactAssay(). This is space-inefficient representation of ragged data when samples are primarily composed of different ranges.
disjoinSummarizedExperiment has rowRanges() identical to the disjoint row ranges of x, disjoint(rowRanges(x)), and assay() data as disjoinAssay().
qreduceSummarizedExperiment has rowRanges() identical to query, and assay() data as qreduceAssay().

Examples

x <- RaggedExperiment(GRangesList(
    GRanges(c("A:1-5", "A:4-6", "A:10-15"), score=1:3),
    GRanges(c("A:1-5", "B:1-3"), score=4:5)
))
## sparseSummarizedExperiment
sse <- sparseSummarizedExperiment(x)
assay(sse)
rowRanges(sse)

## compactSummarizedExperiment
cse <- compactSummarizedExperiment(x)
assay(cse)
rowRanges(cse)

## disjoinSummarizedExperiment
djoinAssay(x, lengths)
dse <- disjoinSummarizedExperiment(x, lengths)
assay(dse)
rowRanges(dse)

## qreduceSummarizedExperiment
x <- RaggedExperiment(GRangesList(
  GRanges(c("A:1-3", "A:4-5", "A:10-15"), score=1:3),
  GRanges(c("A:4-5", "B:1-3"), score=4:5)
))
query <- GRanges(c("A:1-2", "A:4-5", "B:1-5"))

weightedmean <- function(scores, ranges, qranges)
  ## weighted average score per query range
  sum(scores * width(ranges)) / sum(width(ranges))
qreduceAssay(x, query, weightedmean)
qse <- qreduceSummarizedExperiment(x, query, weightedmean)
assay(qse)
rowRanges(qse)
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