Package 'tidyCoverage'

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Title Extract and aggregate genomic coverage over features of interest

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Description `tidyCoverage` framework enables tidy manipulation of collections of genomic tracks and features using `tidySummarizedExperiment` methods. It facilitates the extraction, aggregation and visualization of genomic coverage over individual or thousands of genomic loci, relying on `CoverageExperiment` and `AggregatedCoverage` classes. This accelerates the integration of genomic track data in genomic analysis workflows.

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URL https://github.com/js2264/tidyCoverage

BugReports https://github.com/js2264/tidyCoverage/issues

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AggregatedCoverage aggregate

Description

Bin coverage contained in a CoverageExperiment into an AggregatedCoverage object.

Usage

S4 method for signature 'CoverageExperiment'
aggregate(x, bin = 1, ...)

Arguments

х	a CoverageExperiment object
bin	an integer to bin each assay by. The width of the AggregatedCoverage object should be a multiple of bin.
	ignored

Value

an AggregatedCoverage object

Examples

```
data(ce)
aggregate(ce, bin = 10)
```

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as_tibble-methods as_tibble

Description

Coerce an CoverageExperiment or AggregatedCoverage object into a tibble

Usage

```
## S3 method for class 'AggregatedCoverage'
as_tibble(x, ...)
```

Arguments

Х	an CoverageExperiment object
	ignored

Value

tibble

Examples

data(ac)
as_tibble(ac)

CoverageExperiment CoverageExperiment

Description

#' @description

Usage

```
CoverageExperiment(tracks, features, ...)
```

```
coarsen(x, window, ...)
```

```
## S4 method for signature 'BigWigFileList,GRangesList'
CoverageExperiment(
   tracks,
   features,
   width,
   center = FALSE,
   scale = FALSE,
```

```
ignore.strand = TRUE,
 window = 1,
 BPPARAM = BiocParallel::bpparam()
)
## S4 method for signature 'BigWigFileList,GRanges'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'BigWigFileList,list'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'BigWigFile,GRangesList'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'BigWigFile,GRanges'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'BigWigFile,list'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'list,GRangesList'
CoverageExperiment(
  tracks,
  features,
 width,
  center = FALSE,
  scale = FALSE,
 ignore.strand = TRUE,
 window = 1,
 BPPARAM = BiocParallel::bpparam()
)
## S4 method for signature 'list,GRanges'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'list,list'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'RleList,GRangesList'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'RleList,GRanges'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'RleList,list'
CoverageExperiment(tracks, features, ...)
## S4 method for signature 'CoverageExperiment'
```

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CoverageExperiment

```
coarsen(x, window = 1, BPPARAM = BiocParallel::bpparam())
```

Arguments

tracks	A genomic track imported as a RleList or a <i>named</i> list of genomic tracks.
features	A set of features imported as GRanges or a <i>named</i> GRangesList.
	Passed to the relevant method
х	a CoverageExperiment object
window	an integer to coarsen coverage by.
width	Width to resize each set of genomic features
scale, center	Logical, whether to scale and/or center tracks prior to summarization
ignore.strand	Logical, whether to not take the features strand information
BPPARAM	Passed to BiocParallel.

Details

CoverageExperiment objects store coverages for individual tracks over different sets of features. The coverage assay contains a separate matrix for each combination of track x features. CoverageExperiment objects are instantiated using the CoverageExperiment() #' function, and can be coarsened using the coarsen() function.

Value

A CoverageExperiment object

Examples

```
library(rtracklayer)
library(purrr)
library(plyranges)
TSSs_bed <- system.file("extdata", "TSSs.bed", package = "tidyCoverage")
features <- import(TSSs_bed) |> filter(strand == '+')
## 1. Creating a `CoverageExperiment` object from a single BigWigFile
*****
RNA_fwd <- system.file("extdata", "RNA.fwd.bw", package = "tidyCoverage")</pre>
tracks <- BigWigFile(RNA_fwd)</pre>
CoverageExperiment(tracks, features, width = 5000)
## 2. Creating a `CoverageExperiment` object from a BigWigFileList
RNA_rev <- system.file("extdata", "RNA.rev.bw", package = "tidyCoverage")</pre>
tracks <- BigWigFileList(list(RNA_fwd = RNA_fwd, RNA_rev = RNA_rev))</pre>
CoverageExperiment(tracks, features, width = 5000)
```

```
## 3. Creating a `CoverageExperiment` object from imported bigwig files
tracks <- list(</pre>
  RNA_fwd = system.file("extdata", "RNA.fwd.bw", package = "tidyCoverage"),
  RNA_rev = system.file("extdata", "RNA.rev.bw", package = "tidyCoverage")
) |> map(import, as = 'Rle')
CoverageExperiment(tracks, features, width = 5000)
## 4. Correct for strandness when recovering coverage
TSSs_bed <- system.file("extdata", "TSSs.bed", package = "tidyCoverage")</pre>
features <- list(</pre>
  TSS_fwd = import(TSSs_bed) |> filter(strand == '+'),
  TSS_rev = import(TSSs_bed) |> filter(strand == '-')
)
tracks <- list(</pre>
  RNA_fwd = system.file("extdata", "RNA.fwd.bw", package = "tidyCoverage"),
  RNA_rev = system.file("extdata", "RNA.rev.bw", package = "tidyCoverage")
) |> map(import, as = 'Rle')
CoverageExperiment(tracks, features, width = 5000, ignore.strand = FALSE)
## Aggregating a `CoverageExperiment` object
data(ce)
coarsen(ce, window = 10)
```

data

Example CoverageExperiment and AggregatedCoverage objects

Description

Two example objects are provided in the tidyCoverage package:

- ce: a CoverageExperiment dataset containing stranded RNA-seq coverage (forward and reverse) over Scc1 peaks (± 1kb).
- ac: an AggregatedCoverage object obtained with aggregate(ce).

Usage

data(ce)

data(ac)

Format

CoverageExperiment object containing 1 features set and 2 tracks. AggregatedCoverage object containing 1 features set and 2 tracks.

Details

Data was generated in yeast (S288c) and aligned to reference R64-1-1.

expand, CoverageExperiment

Expand a CoverageExperiment object

Description

A CoverageExperiment object can be coerced into a tibble using the tidySummarizedExperiment package, but this will not turn each coverage matrix into a "long" format. The expand function provided here allows one to coerce a CoverageExperiment object into a long data frame, and adds the ranges and seqnames to the resulting tibble.

Usage

S3 method for class 'CoverageExperiment'
expand(data, ..., .name_repair = NULL)

Arguments

data a CoverageExperiment object

..., .name_repair ignored

Value

a tibble object

Examples

data(ce) ce

expand(ce)

```
reexports
```

Description

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

dplyr as_tibble
S4Vectors aggregate
tidyr expand

Value

Depending on the re-exported function

Examples

1 + 1

show

show

Description

show method for CoverageExperiment and AggregatedCoverage objects

Usage

```
## S4 method for signature 'CoverageExperiment'
show(object)
## S4 method for signature 'AggregatedCoverage'
show(object)
## S3 method for class 'CoverageExperiment'
print(x, ..., n = NULL)
## S3 method for class 'AggregatedCoverage'
print(x, ..., n = NULL)
## S3 method for class 'tidyCoverageExperiment'
tbl_format_header(x, setup, ...)
## S3 method for class 'tidyAggregatedCoverage'
tbl_format_header(x, setup, ...)
```

show

Arguments

object	a CoverageExperiment or AggregatedCoverage object
х	Object to format or print.
	Passed on to tbl_format_setup().
n	Number of rows to show. If NULL, the default, will print all rows if less than the print_max option. Otherwise, will print as many rows as specified by the print_min option.
setup	a setup object returned from pillar::tbl_format_setup().

Value

Prints a message to the console describing the contents of the CoverageExperimentorAggregatedCoverage'objects.

Examples

data(ce)
print(ce)
data(ac)
print(ac)

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