# rtracklayer

April 19, 2010

activeView-methods Accessing the active view

# Description

Get the active view.

# Methods

The following methods are defined by **rtracklayer**.

activeView(object) <- value: Sets the active BrowserView in a browser session.

BasicTrackLine-class

Class "BasicTrackLine"

# Description

The type of UCSC track line used to annotate most types of tracks (every type except Wiggle).

# **Objects from the Class**

Objects can be created by calls of the form new ("BasicTrackLine", ...) or parsed from a character vector track line with as (text, "BasicTrackLine") or converted from a GraphTrackLine using as (wig, "BasicTrackLine").

#### Slots

- itemRgb: Object of class "logical" indicating whether each feature in a track uploaded as BED should be drawn in its specified color.
- useScore: Object of class "logical" indicating whether the data value should be mapped to color.

group: Object of class "character" naming a group to which this track should belong.

- db: Object of class "character" indicating the associated genome assembly.
- offset: Object of class "numeric", a number added to all positions in the track.
- url: Object of class "character" referring to additional information about this track.
- htmlUrl: Object of class "character" referring to an HTML page to be displayed with this track.
- name: Object of class "character" specifying the name of the track.
- description: Object of class "character" describing the track.
- visibility: Object of class "character" indicating the default visible mode of the track, see UCSCTrackModes.

color: Object of class "integer" representing the track color (as from col2rgb).

priority: Object of class "numeric" specifying the rank of the track.

# Extends

Class "TrackLine", directly.

## Methods

as(object, "character") Export line to its string representation.

**as(object,** "GraphTrackLine") Convert this line to a graph track line, using defaults for slots not held in common.

#### Author(s)

Michael Lawrence

#### References

http://genome.ucsc.edu/goldenPath/help/customTrack.html#TRACK for the
official documentation.

# See Also

GraphTrackLine for Wiggle/bedGraph tracks.

Bed15TrackLine-class

Class "Bed15TrackLine"

## Description

A UCSC track line for graphical tracks.

## **Objects from the Class**

Objects can be created by calls of the form new("Bed15TrackLine", ...) or parsed from a character vector track line with as (text, "Bed15TrackLine").

# Slots

expStep: A "numeric" scalar indicating the step size for the heatmap color gradient.

expScale: A positive "numeric" scalar indicating the range of the data to be [-expScale, expScale] for determining the heatmap color gradient.

expNames: A "character" vector naming the the experimental samples.

name: Object of class "character" specifying the name of the track.

description: Object of class "character" describing the track.

visibility: Object of class "character" indicating the default visible mode of the track, see UCSCTrackModes.

color: Object of class "integer" representing the track color (as from col2rgb).

priority: Object of class "numeric" specifying the rank of this track.

# Extends

Class "TrackLine", directly.

#### Methods

as(object, "character") Export line to its string representation.

## Author(s)

Michael Lawrence

# References

Official documentation: http://genomewiki.ucsc.edu/index.php/Microarray\_track.

# See Also

export.bed15 for exporting bed15 tracks.

blocks-methods Get blocks/exons

## Description

Obtains the block ranges (subranges, usually exons) from an object, such as a RangedData imported from a BED file.

# Usage

blocks(x, ...)

## Arguments

Х	The instance from which to obtain the block/exon information. Currently must
	be a RangedData, presumably imported with import.bed.
	Additional arguments for methods

## Details

For the RangedData method, there must be two columns in x: blockStarts and blockSizes, each field of which should be a comma-separated list of block starts and widths, respectively. This comes from the BED specification.

## Author(s)

Michael Lawrence

## See Also

import.bed for importing a track from BED, which can store block information.

browseGenome Browse a genome

# Description

A generic function for launching a genome browser.

# Usage

```
browseGenome(object, ...)
## S4 method for signature 'RangedDataORRangedDataList':
browseGenome(object,
    browser = "UCSC", range = base::range(object),
    view = TRUE, trackParams = list(), viewParams = list(), ...)
```

#### Arguments

object	A list of RangedData instances, e.g. a RangedDataList instance.
browser	The name of the genome browser.
range	The RangesList to display in the initial view.
view	Whether to open a view.
trackParams	Named list of parameters to pass to track<
viewParams	Named list of parameters to pass to browserView.
	Arguments corresponding to slots in RangesList that override those in range.

## Value

Returns a BrowserSession.

# Author(s)

Michael Lawrence

# See Also

BrowserSession and BrowserView, the two main classes for interfacing with genome browsers.

## Examples

```
## Not run:
## open UCSC genome browser:
browseGenome()
## to view a specific range:
range <- GenomicRanges("hg18", "chr22", 20000, 50000)
browseGenome(range = range)
## a slightly larger range:
browseGenome(range = range, end = 75000)
## with a track:
track <- import(system.file("tests", "v1.gff", package = "rtracklayer"))
browseGenome(RangedDataList(track))
```

## End(Not run)

BrowserSession-class

Class "BrowserSession"

# Description

An object representing a genome browser session. Each session corresponds to a set of loaded tracks and a set of BrowserView instances. Note that this is a virtual class; a concrete implementation is provided by each backend driver.

# **Objects from the Class**

A virtual Class: No objects may be created from it. See browserSession for obtaining an instance of an implementation for a particular genome browser.

## Methods

This specifies the API implemented by each browser backend. Note that a backend is not guaranteed to support all operations, and that each backend often has additional parameters for each of the methods. See the backend-specific documentation for more details. The only built-in backend is UCSCSession.

```
browserView(object, range = range(object), track = trackNames(object), ...)
Constructs a BrowserView of range for this session.
```

browserViews (object, ...) Gets the BrowserView instances belonging to this session.

- activeView(object, ...) Returns the BrowserView that is currently active in the session.
- range (x, ...) Gets the RangesList representing the range of the genome currently displayed by the browser (i.e. the range shown by the active view) or a default value (possibly NULL) if no views exist.
- getSeq(object, range = range(object), ...) gets a genomic sequence of range
   from this session.

sequence(object, ...) <- value Loads a sequence into the session.</pre>

- track(object, name = deparse(substitute(track)), view = TRUE, ...) <- value Loads one or more tracks into the session and optionally open a view of the track.
- x[[i]] <- value Loads the track value into session x, under the name i. Shortcut to above.
- x\$name <- value Loads the track value into session x, under the name name. Shortcut to above.

track (object, ...) Gets a track from a session as a RangedData.

x[[i]] Gets the track named i from session x. A shortcut to track.

x\$name Gets the track named name from session x. A shortcut to track.

trackNames (object, ...) Gets the names of the tracks stored in this session.

genome (x), genome (x) <- value Gets or sets the genome identifier (e.g. "hg18") for the session.

close (con, ...) Close this session.

show(object, ...) Output a textual description of this session.

# Author(s)

Michael Lawrence

# See Also

browserSession for obtaining implementations of this class for a particular genome browser.

browserSession-methods

Get a genome browser session

## Description

Methods for getting browser sessions.

#### Methods

The following methods are defined by rtracklayer.

**object = "character"** browserSession(object, ...): Creates a BrowserSession from a genome browser identifier. The identifier corresponds to the prefix of the session class name (e.g. "ucsc" in "UCSCSession"). The arguments in ... are passed to the initialization function of the class.

**object = "browserView"** Gets the BrowserSession for the view.

```
object = "missing" Calls browserSession("ucsc", ...).
```

BrowserView-class Class "BrowserView"

#### Description

An object representing a genome browser view of a particular segment of a genome.

## **Objects from the Class**

A virtual Class: No objects may be created from it directly. See browserView for obtaining an instance of an implementation for a particular genome browser.

#### Slots

session: Object of class "BrowserSession" the browser session to which this view belongs.

## Methods

This specifies the API implemented by each browser backend. Note that a backend is not guaranteed to support all operations. See the backend-specific documentation for more details. The only built-in backend is UCSCView.

browserSession (object) Obtains the BrowserSession to which this view belongs.

```
close (object) Close this view.
```

range(object) Obtains the RangesList displayed by this view.

trackNames (object) Gets the names of the visible tracks in the view.

trackNames(object) <- value Sets the visible tracks by their names.</pre>

show (object) Outputs a textual description of this view.

visible(object) Get a named logical vector indicating whether each track is visible.

```
visible(object) <- value Set a logical vector indicating the visibility of each track, with
the same names and in the same order as that returned by visible(object).
```

# Author(s)

Michael Lawrence

# See Also

browserView for obtaining instances of this class.

browserView-methods

Getting browser views

# Description

Methods for creating and getting browser views.

# Usage

browserView(object, range, track, ...)

# Arguments

object	The object from which to get the views.
range	The RangesList to display.
track	List of track names to make visible in the view.
•••	Arguments to pass to methods

## Methods

The following methods are defined by rtracklayer.

```
object = "UCSCSession" browserView(object, range = range(object), track
= trackNames(object), imagewidth = 800, ...): Creates a BrowserView
of range with visible tracks specified by track. The imagewidth parameter specifies the
width of the track image in pixels. track may be an instance of UCSCTrackModes. Argu-
ments in ... are passed to ucscTrackModes to create the UCSCTrackModes instance
that will override modes indicated by the track parameter.
```

# Examples

```
## Not run:
    session <- browserSession()
    browserView(session, GenomicRanges(20000, 50000, "chr2"))
    ## only view "knownGene" track
    browserView(session, track = "knownGene")
```

## End(Not run)

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browserViews-methods

Getting the browser views

# Description

Methods for getting browser views.

# Methods

The following methods are defined by rtracklayer.

Gets the instances of BrowserView in the session.

## See Also

object = "UCSCSession" browserView for creating a browser view.

# Examples

```
## Not run:
session <- browseGenome()
browserViews(session)
```

## End(Not run)

cpneTrack CPNE1 SNP track

# Description

A RangedData object (created by the GGtools package) with features from a subset of the SNPs on chromosome 20 from 60 HapMap founders in the CEU cohort. Each SNP has an associated data value indicating its association with the expression of the CPNE1 gene according to a Cochran-Armitage 1df test. The top 5000 scoring SNPs were selected for the track.

# Usage

```
data(cpneTrack)
```

## Format

Each feature (row) is a SNP. The association test scores are accessible via score.

## Source

Vince Carey and the GGtools package.

# Examples

```
data(cpneTrack)
plot(start(cpneTrack), score(cpneTrack))
```

export-tracks Export tracks

# Description

These functions output RangedData instances in various formats.

# Usage

# Arguments

object	The object to export, such as a RangedData, or anything coercible to a RangedData If a UCSCData, the track line information is output. In the case of export.bed15, export.bedGraph, export.wig, and export.ucsc, a RangedDataList
	object with possibly multiple tracks is supported.
con	The connection to which the object is exported.
version	The GFF version, either "1", "2" or "3" (default is "1").
source	The source of the GFF information, for GFF.
variant	Which variant of BED lines to output, not for the user.
color	Recycled vector of colors, as interpreted by collrgb for BED features. If NULL, the color column in the featureData is used, if any.
dataFormat	The format of the data lines for WIG tracks, see references. The "auto" format uses the most efficient format possible.
subformat	The format of the tracks within the UCSC container. If "auto", the type is de- termined from the trackline. If object is not a UCSCData, this essentially means "wig" or "bedGraph" (depending on the density) if there is a numeric score, else "bed".
expNames	Names of the columns in object that hold the experimental data. Defaults to all column names, unless object is a UCSCData, in which case the expNames field is taken from the track line, if it exists.
	For export.gff1, export.gff2 and export.gff3: arguments to pass to export.gff. For export.bed: arguments to pass to methods. For export.bed15, export.bedGraph and export.wig: arguments to pass to export.ucsc. For export.ucsc: arguments to pass to export.subformat or to set on the slots of the TrackLine subclass corresponding to subformat.

#### export-tracks

#### Details

The following is some advice for choosing a file format.

- GFF The General Feature Format is meant to represent any set of genomic features, with applicationspecific columns represented as "attributes". There are three principal versions (1, 2, and 3). This is a good format for interoperating with other genomic tools. UCSC supports GFF1, but it needs to be encapsulated in the UCSC metaformat, i.e. export.ucsc(subformat = "gff1").
- BED The Browser Extended Display format is for displaying tracks in a genome browser, in particular UCSC. There are many options to control the appearance of the track, see GraphTrackLine. To output a track line when object is not a UCSCData, call export.ucsc(subformat = "bed").
- **BED15** An extension of BED with 15 columns, Bed15 is meant to represent data from microarray experiments. Multiple samples/columns are supported, and the data is displayed as a compact heatmap. With 15 columns per feature, this format is probably too verbose for e.g. ChIP-seq coverage (use multiple WIG tracks instead).
- **BEDGRAPH** A variant of BED that represents experimental data more compactly than BED and especially BED15, although only one sample is supported. The data is displayed as a bar or line graph. For dense data, WIG is preferred.
- WIG The Wiggle format is meant for storing dense numerical data, such as the coverage from a ChIP-seq experiment. The data is displayed as a bar or line graph.

In summary, BED is usually best for displaying qualitative features or sparse quantiative features (like ChIP-seq peaks), while WIG is usually best for displaying dense data like coverage.

# Value

If con is missing, a character vector containing the string output, otherwise nothing.

### Author(s)

Michael Lawrence

#### References

GFF1 and GFF2 http://www.sanger.ac.uk/Software/formats/GFF
GFF3 http://www.sequenceontology.org/gff3.shtml
BED http://genome.ucsc.edu/goldenPath/help/customTrack.html#BED
WIG http://genome.ucsc.edu/goldenPath/help/wiggle.html
UCSC http://genome.ucsc.edu/goldenPath/help/customTrack.html

#### See Also

See export for the high-level interface to these functions.

#### Examples

```
dummy <- file() # dummy file connection for demo
track <- import(system.file("tests", "bed.wig", package = "rtracklayer"))
## output a track as GFF2
export.gff(track, dummy, version = "2")
## equivalently</pre>
```

```
export.gff2(track, dummy)
## output as WIG string in variableStep format
wig <- export.wig(track, dummy, dataFormat = "variableStep")
## output multiple tracks in UCSC meta-format
track2 <- import(system.file("tests", "v1.gff", package = "rtracklayer"))
## output to WIG
export.ucsc(RangedDataList(track, track2), dummy, subformat = "wig")</pre>
```

```
export
```

# Export objects to connections

# Description

Exports (serializes) an object in a given format to a given connection.

# Usage

```
export(object, con, format, ...)
```

## Arguments

object	The object to export.
con	The connection to which the object is exported. If this is a character vector, it is assumed to be a filename and a corresponding file connection is created and then closed after exporting the object. If missing, the function will return the output as a character vector, rather than writing to a connection.
format	The format of the output. If missing and $con$ is a filename, the format is derived from the file extension.
	Parameters to pass to the format-specific export routine.

# Details

This function delegates to another function, depending on the specified format. The name of the delegate is of the form export.format where format is specified by the format argument.

# Value

If con is missing, a character vector containing the string output. Otherwise, nothing is returned.

# Author(s)

Michael Lawrence

#### See Also

import for the reverse

#### genomeBrowsers

# Examples

```
track <- import(system.file("tests", "v1.gff", package = "rtracklayer"))
## Not run: export(track, "my.gff", version = "3")
## equivalently,
## Not run: export(track, "my.gff3")
## or
## Not run:
con <- file("my.gff3")
export(track, con, "gff3")
close(con)
## End(Not run)
## or as a string
export(track, format = "gff3")</pre>
```

genomeBrowsers Get available genome browsers

# Description

Gets the identifiers of the loaded genome browser drivers.

# Usage

```
genomeBrowsers(where = topenv(parent.frame()))
```

## Arguments

where The environment in which to search for drivers.

# Details

This searches the specified environment for classes that extend BrowserSession. The prefix of the class name, e.g. "ucsc" in "UCSCSession", is returned for each driver.

#### Value

A character vector of driver identifiers.

## Author(s)

Michael Lawrence

# See Also

browseGenome and browserSession that create browserSession implementations given an identifier returned from this function.

getSeq-methods *Retrieving a genome sequence* 

# Description

Methods for retrieving the sequence of a RangesList from an object.

# Methods

```
The following methods are defined by rtracklayer for getSeq(object, range = range(object), ...).
```

```
object = "UCSCSession" getSeq(object, range = range(object), track = "Assembly"):
    Gets the sequence in range and track from the session.
```

# See Also

sequence<- for storing sequences.

import.gff	Importing tracks

# Description

These are the functions for importing RangedData instances from connections or text.

# Usage

# Arguments

con	The connection from which to receive the input.
version	The version of GFF ("1", "2" or "3").
genome	The genome to set on the imported track.
variant	Variant of BED lines, not for the user.
trackLine	Whether the BED data has a track line (it normally does though track lines are not mandatory).

#### import

subformat	The expected subformat of the UCSC data. If "auto", automatic detection of the subformat is attempted.
drop	If TRUE and there is only one track in the UCSC data, return the track instead of a list.
	For import.gff1, import.gff2 and import.gff3: arguments to pass to import.gff. For import.ucsc: arguments to pass on to import.subformat. For the others, arguments to pass to methods.

## Value

An instance of RangedData or one of its subclasses, except for import.ucsc, which returns a RangedDataList instance, unless there is one track and the drop parameter is TRUE.

# Author(s)

Michael Lawrence

# References

GFF1 and GFF2 http://www.sanger.ac.uk/Software/formats/GFF
GFF3 http://www.sequenceontology.org/gff3.shtml
BED http://genome.ucsc.edu/goldenPath/help/customTrack.html#BED
WIG http://genome.ucsc.edu/goldenPath/help/wiggle.html
UCSC http://genome.ucsc.edu/goldenPath/help/customTrack.html

#### See Also

import for the high-level interface to these routines.

## Examples

```
import
```

Importing objects

# Description

Imports an object from a connection according to a specified format.

# Usage

import(con, format, text, ...)

# Arguments

con	The connection through which the data is received. If this is a character vector, it is assumed to be a filename.
format	The format in which to expect the input. If omitted and con is a filename, the format is taken from the file extension.
text	If $\operatorname{con}$ is missing, this can be a character vector directly providing the string data to import.
•••	Arguments to pass to the format-specific import routines.

# Details

This function delegates to a format-specific function named according to the scheme import.format where format is specified by the format parameter.

# Value

The object parsed from the connection or text.

## Author(s)

Michael Lawrence

## See Also

export to do the reverse.

# Examples

```
track <- import(system.file("tests", "bed.wig", package = "rtracklayer"))
track <- import(system.file("tests", "v1.gff", package = "rtracklayer"), version = "1")
# or
track <- import(system.file("tests", "v1.gff", package = "rtracklayer"), "gff1")</pre>
```

sequence <- methods Load a sequence

# Description

Methods for loading sequences.

#### Methods

No methods are defined by **rtracklayer** for the sequence (object, ...) <- value generic.

track<-methods Laying tracks

#### Description

Methods for loading RangedData instances (tracks) into genome browsers.

# Usage

```
## S4 method for signature 'BrowserSession,RangedData':
track(object, name = deparse(substitute(track)), view = FALSE, ...) <- value</pre>
```

## Arguments

object	A BrowserSession into which the track is loaded.
value	The track(s) to load.
name	The name(s) of the track(s) being loaded.
view	Whether to create a view of the track after loading it.
	Arguments to pass on to methods.

## Methods

The following methods are defined by **rtracklayer**. A browser session implementation must implement a method for either RangedData or RangedDataList. The base browserSession class will delegate appropriately.

object = "BrowserSession", value = "RangedData" Load this track into the session.

**object = "BrowserSession", value = "RangedDataList"** Load all tracks into the session.

```
object = "UCSCSession", value = "RangedDataList" track(object, name = deparse(substitute(t)
view = FALSE, format = "gff") <- value: Load the tracks into the session us-
ing the specified format.
```

## See Also

track for getting a track from a session.

## Examples

```
## Not run:
    session <- browserSession()
    track <- import(system.file("tests", "v1.gff", package = "rtracklayer"))
    track(session, "My Track") <- track
## End(Not run)
```

RangedData-methods Data on a Genome

#### Description

The rtracklayer package adds convenience methods on top of RangedData manipulate data on genomic ranges. The spaces are now called chromosomes (but could still refer to some other type of sequence). The universe refers to the genome.

## Accessors

In the code snippets below, x is a RangedData object.

- chrom(x), chrom(x) <- value: Gets or sets the chromosome names for x. The length
   of value should equal the length of x. This is an alias for names(x).</pre>
- genome(x), genome(x) <- value: Gets or sets the genome (a single string or NULL) for the ranges in x; simple wrappers around universe and universe<-, respectively.</pre>

#### Constructor

GenomicData(ranges, ..., strand = NULL, chrom = NULL, genome = NULL): Constructs a RangedData instance with the given ranges and variables in ... (see the RangedData constructor). If non-NULL, strand specifies the strand of each range. It should be a character vector or factor of length equal to that of ranges. All values should be either -, +, \* or NA. To get these levels, call levels(strand()). chrom is analogous to splitter in RangedData; if non-NULL it should be coercible to a factor indicating how the ranges, variables and strand should be split up across the chromosomes. The genome argument should be a scalar string and is treated as the RangedData universe. See the examples.

If ranges is not a Ranges object, this function calls as (ranges, "RangedData") and returns the result if successful. As a special case, the "chrom" column in a data.framelike object is renamed to "space", for convenience. Thus, one could pass a data.frame with columns "start", "end" and, optionally, "chrom".

# Author(s)

Michael Lawrence

# Examples

```
range1 <- IRanges(start=c(1,2,3), end=c(5,2,8))
## just ranges</pre>
```

```
gr <- GenomicData(rangel)
## with a genome (universe)
gr <- GenomicData(rangel, genome = "hg18")
genome(gr) ## "hg18"
## with some data
filter <- c(11 01 11)</pre>
```

```
filter <- c(1L, 0L, 1L)
score <- c(10L, 2L, NA)
```

#### RangesList-methods

```
strand <- factor(c("+", NA, "-"), levels = levels(strand()))</pre>
gr <- GenomicData(range1, score, genome = "hg18")</pre>
gr[["score"]]
strand(gr) ## all NA
gr <- GenomicData(rangel, score, filt = filter, strand = strand)</pre>
gr[["filt"]]
strand(gr) ## equal to 'strand'
range2 <- IRanges(start=c(15,45,20,1), end=c(15,100,80,5))</pre>
ranges <- c(range1, range2)</pre>
score <- c(score, c(0L, 3L, NA, 22L))</pre>
chrom <- paste("chr", rep(c(1,2), c(length(range1), length(range2))), sep="")</pre>
gr <- GenomicData(ranges, score, chrom = chrom, genome = "hg18")</pre>
chrom(gr) # equal to 'chrom'
gr[["score"]] # unlists over the chromosomes
score(gr)
gr[1][["score"]] # equal to score[1:3]
df <- as.data.frame(gr)</pre>
GenomicData(df)
```

RangesList-methods Ranges on a Genome

#### Description

Genomic coordinates are often specified in terms of a genome identifier, chromosome name, start position and end position. This information can be represented by a RangesList instance, and the rtracklayer package adds convenience methods to RangesList for the manipulation of genomic ranges. The spaces (or names) of RangesList are the chromosome names. The universe slot indicates the genome, usually as given by UCSC (e.g. "hg18").

## Accessors

In the code snippets below, x is a RangesList object.

- chrom(x), chrom(x) <- value: Gets or sets the chromosome names for x. This is an alias for names(x).
- genome(x), genome(x) <- value: Gets or sets the genome (a single string or NULL) for the ranges in x; simple wrappers around universe and universe<-, respectively.</pre>

## Constructor

GenomicRanges (start, end, chrom = NULL, genome = NULL): Constructs a RangesList containing ranges specified by start and end, optionally split into elements based on chrom, a vector of chromosome identifiers (or NULL for no splitting). The genome argument should be a scalar string and is treated as the RangesList universe. See the examples.

## Author(s)

Michael Lawrence

targets

# Examples

```
GenomicRanges(c(1,2,3), c(5,2,8))
GenomicRanges(c(1,2,3), c(5,2,8), c("chr1", "chr1", "chr2"))
GenomicRanges(c(1,2,3), c(5,2,8), genome = "hg18")
```

targets

#### microRNA target sites

# Description

A data frame of human microRNA target sites retrieved from MiRBase. This is a subset of the hsTargets data frame in the microRNA package. See the rtracklayer vignette for more details.

## Usage

data(targets)

# Format

A data frame with 2981 observations on the following 6 variables.

name The miRBase ID of the microRNA.

target The Ensembl ID of the targeted transcript.

chrom The name of the chromosome for target site.

start Target start position.

end Target stop position.

strand The strand of the target site, "+", or "-".

# Source

The microRNA package, dataset hsTargets. Originally MiRBase (http://microrna.sanger.ac.uk/).

# Examples

```
data(targets)
targetTrack <- with(targets,
    GenomicData(IRanges(start, end),
        strand = strand, chrom = chrom))</pre>
```

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tracks-methods Accessing track names

## Description

Methods for getting and setting track names.

#### Methods

The following methods are defined by **rtracklayer** for **getting** track names via the generic trackNames(object, ...).

Get the tracks loaded in the session.

- **object = "UCSCSessiohject = "UCSCTrackModes"** Get the visible tracks according to the modes (all tracks not set to "hide").
- **object = "UCSCView"** Get the visible tracks in the view.

The following methods are defined by **rtracklayer** for **setting** track names via the generic trackNames(object) <- value.

- **object = "UCSCTrackModes"** Sets the tracks that should be visible in the modes. All specified tracks with mode "hide" in object are set to mode "full". Any tracks in object that are not specified in the value are set to "hide". No other modes are changed.
- **object = "UCSCView"** Sets the visible tracks in the view. This opens a new web browser with only the specified tracks visible.

UCSCData-class Class "UCSCData"

# Description

Each track in UCSC has an associated TrackLine that contains metadata on the track.

## Slots

trackLine: Object of class "TrackLine" holding track metadata.

# Methods

- export.bed(object, con, variant = c("base", "bedGraph", "bed15"), color, trackLi
  Exports the track and its track line (if trackLine is TRUE) to con in the Browser Extended
  Display (BED) format. The arguments in ... are passed to export.ucsc.
- export.bed15(object, con, expNames = NULL, ...) Exports the track and its track line(if trackLine is TRUE) to con in the Bed15 format. The data is taken from the columns named in expNames, which defaults to the expNames in the track line, if any, otherwise all column names. The arguments in ... are passed to export.ucsc.
- export.gff(object) Exports the track and its track line (as a comment) to con in the General Feature Format (GFF).

as (object, "UCSCData") Constructs a UCSCData from a RangedData instance, by adding a default track line and ensuring that the sequence/chromosome names are compliant with UCSC conventions. If there is a numeric score, the track line type is either "bedGraph" or "wig", depending on the feature density. Otherwise, "bed" is chosen.

# Author(s)

Michael Lawrence

# See Also

import and export for reading and writing tracks to and from connections (files), respectively.

ucscGenomes Get available genomes on UCSC

## Description

Get a list of genome identifiers for the UCSC genome browser.

# Usage

```
ucscGenomes()
```

## Value

A character vector of genome identifiers, with names indicating the species.

#### Author(s)

Michael Lawrence

# See Also

UCSCSession for details on specifying the genome.

UCSCSession-class Class "UCSCSession"

# Description

An implementation of BrowserSession for the UCSC genome browser.

## **Objects from the Class**

Objects can be created by calls of the form browserSession ("ucsc", url = "http://genome.ucsc.edu bin", ...). The arguments in ... correspond to libcurl options, see getCurlHandle. Setting these options may be useful e.g. for getting past a proxy.

#### UCSCSession-class

# Slots

url: Object of class "character" holding the base URL of the UCSC browser.

- hguid: Object of class "numeric" holding the user identification code.
- views: Object of class "environment" containing a list stored under the name "instances". The list holds the instances of BrowserView for this session.

## Extends

Class "BrowserSession", directly.

#### Methods

browserView(object, range = range(object), track = trackNames(object), ...)
Creates a BrowserView of range with visible tracks specified by track. track may be
an instance of UCSCTrackModes. Arguments in ... should override slots in range or
else match parameters to a ucscTrackModes method for creating a UCSCTrackModes
instance that will override modes indicated by the track parameter.

browserViews(object) Gets the BrowserView instances for this session.

range(x) Gets the RangesList last displayed in this session.

- genome (x) Gets the genome identifier of the session, i.e. genome (range (x)).
- range(x) <- value Sets value, a RangesList, as the range of session x. Note that this setting only lasts until a view is created or manipulated. This mechanism is useful, for example, when treating the UCSC browser as a database, rather than a genome viewer.
- genome (x) <- value Sets the genome identifier on the range of session x.
- getSeq(object, range, track = "Assembly") Gets the sequence in range and track.
- track(object, name = names(track), view = TRUE, format = "gff", ...) <- value Loads a track, stored under name and formatted as format. The arguments in ... are passed on to export.ucsc, so they could be slots in a TrackLine subclass or parameters to pass on to the export function for format.
- track (object, name, range = range (object), table = NULL) Retrieves a RangedData
  with features in range from track named name. Some built-in tracks have multiple series,
  each stored in a separate database table. A specific table may be retrieved by passing its name
  in the table parameter. See tableNames for a way to list the available tables.
- trackNames(object) Gets the names of the tracks stored in the session.

ucscTrackModes(object) Gets the default view modes for the tracks in the session.

#### Author(s)

Michael Lawrence

## See Also

browserSession for creating instances of this class.

UCSCTableQuery-class

Querying UCSC Tables

# Description

The UCSC genome browser is backed by a large database, which is exposed by the Table Browser web interface. Tracks are stored as tables, so this is also the mechanism for retrieving tracks. The UCSCTableQuery class represents a query against the Table Browser. Storing the query fields in a formal class facilitates incremental construction and adjustment of a query.

## Details

There are five supported fields for a table query:

- session The UCSCSession instance from the tables are retrieved. Although all sessions are based on the same database, the set of user-uploaded tracks, which are represented as tables, is not the same, in general.
- **trackName** The name of a track from which to retrieve a table. Each track can have multiple tables. Many times there is a primary table that is used to display the track, while the other tables are supplemental. Sometimes, tracks are displayed by aggregating multiple tables.
- **tableName** The name of the specific table to retrieve. May be NULL, in which case the behavior depends on how the query is executed, see below.
- range A RangesList indicating the portion of the table to retrieve, in genome coordinates. The genome indicated by the RangesList also determines which tracks are available and must always be non-NULL. If the RangesList is empty, the table is downloaded for the entire genome.
- names Names/accessions of the desired features

A common workflow for querying the UCSC database is to create an instance of UCSCTableQuery using the ucscTableQuery constructor, invoke tableNames to list the available tables for a track, and finally to retrieve the desired table either as a data.frame via getTable or as a RangedData track via track. See the examples.

The reason for a formal query class is to facilitate multiple queries when the differences between the queries are small. For example, one might want to query multiple tables within the track and/or same genomic region, or query the same table for multiple regions. The UCSCTableQuery instance can be incrementally adjusted for each new query. Some caching is also performed, which enhances performance.

## Constructor

#### **Executing Queries**

Below, object is a UCSCTableQuery instance.

- track (object): Retrieves the indicated table as a track, i.e. a RangedData instance. Note that not all tables are available as tracks.
- getTable(object): Retrieves the indicated table as a data.frame. Note that not all tables
   are output in parseable form.
- tableNames (object): Gets the names of the tables available for the session, track and range specified by the query.

## Accessor methods

In the code snippets below, x/object is a UCSCTableQuery object.

- browserSession(object), browserSession(object) <- value: Get or set the UCSCSession to query.
- trackNames (x) List the names of the tracks available for retrieval for the assigned genome.
- tableName(x), tableName(x) <- value: Get or set the single string indicating the name of the table to retrieve. May be NULL, in which case the table is automatically determined.
- range(x), range(x) <- value: Get or set the RangesList indicating the portion of the table to retrieve in genomic coordinates. Any missing information, such as the genome identifier, is filled in using range (browserSession(x)).
- names(x), names(x) <- value: Get or set the names of the features to retrieve. If NULL, this filter is disabled.

#### Author(s)

Michael Lawrence

# Examples

```
## Not run:
session <- browserSession()</pre>
genome(session) <- "mm9"</pre>
trackNames(session) ## list the track names
## choose the Conservation track for a portion of mm9 chr1
query <- ucscTableQuery(session, "Conservation",</pre>
                          GenomicRanges(57795963, 57815592, "chr12"))
## list the table names
tableNames(query)
## get the phastCons30way track
tableName(query) <- "phastCons30way"</pre>
## retrieve the track data
track(query)
## get a data.frame summarizing the multiple alignment
tableName(query) <- "multiz30waySummary"</pre>
getTable(query)
genome(session) <- "hg18"</pre>
query <- ucscTableQuery(session, "snp129",</pre>
```

```
names = c("rs10003974", "rs10087355", "rs10075230"))
```

```
getTable(query)
```

## End(Not run)

TrackLine-class Class "TrackLine"

## Description

An object representing a "track line" in the UCSC format. There are two concrete types of track lines: BasicTrackLine (used for most types of tracks) and GraphTrackLine (used for graphical tracks). This class only declares the common elements between the two.

## **Objects from the Class**

Objects can be created by calls of the form new("TrackLine", ...) or parsed from a character vector track line with as (text, "TrackLine"). But note that UCSC only understands one of the subclasses mentioned above.

## Slots

name: Object of class "character" specifying the name of the track.

description: Object of class "character" describing the track.

visibility: Object of class "character" indicating the default visible mode of the track, see UCSCTrackModes.

color: Object of class "integer" representing the track color (as from col2rgb).

priority: Object of class "numeric" specifying the rank of this track.

## Methods

as(object, "character") Export line to its string representation.

# Author(s)

Michael Lawrence

#### References

http://genome.ucsc.edu/goldenPath/help/customTrack.html#TRACK for the
official documentation.

## See Also

BasicTrackLine (used for most types of tracks) and GraphTrackLine (used for Wiggle/bedGraph tracks).

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UCSCTrackModes-class

Class "UCSCTrackModes"

#### Description

A vector of view modes ("hide", "dense", "full", "pack", "squish") for each track in a UCSC view.

#### **Objects from the Class**

Objects may be created by calls of the form ucscTrackModes (object = character(), hide = character(), dense = character(), pack = character(), squish = character(), full = character()), where object should be a character vector of mode names (with its names attribute specifying the corresponding track names). The other parameters should contain track names that override the modes in object.

# Slots

- .Data: Object of class "character" holding the modes ("hide", "dense", "full", "pack", "squish"), with its names attribute holding corresponding track names.

#### Extends

Class "character", from data part. Class "vector", by class "character", distance 2. Class characterORMIAME, by class "character", distance 2.

### Methods

- trackNames(object) <- value Sets the names of the visible tracks. Any tracks named in value are set to "full" if the are currently set to "hide" in this object. Any tracks not in value are set to "hide". All other modes are preserved.
- object[i] Gets the track mode of the tracks indexed by i, which can be any type of index supported by character vector subsetting. If i is a character vector, it indexes first by the internal track IDs (the names on .Data) and then by the user-level track names (the labels slot).

#### Author(s)

Michael Lawrence

#### See Also

UCSCView on which track view modes may be set.

ucscTrackModes-methods

Accessing UCSC track modes

# Description

Generics for getting and setting UCSC track visibility modes ("hide", "dense", "full", "pack", "squish").

## Methods

The following methods are defined by **rtracklayer** for **getting** the track modes through the generic ucscTrackModes(object, ...).

function(object, hide = character(), dense = character(), pack = character(), squish = character(), full = character()) Creates an instance of UCSCTrackModes from object, a character vector of mode names, with the corresponding track names given in the names attribute. Note that object can be a UCSCTrackModes instance. The other parameters are character vectors naming the tracks for each mode and overriding the modes specified by object.

- **object = "charactdrject = "missing"** The same interface as above, except object defaults to an empty character vector.
- object = "UCSCView" Gets modes for tracks in the view.
- **object = "UCSCSession"** Gets default modes for the tracks in the session. These are the modes that will be used as the default for a newly created view.

The following methods are defined by **rtracklayer** for **setting** the track modes through the generic ucscTrackModes(object) <- value.

- object = "UCSCView", value = "UCSCTrackModes" Sets the modes for the tracks in the view.
- **object = "UCSCView", value = "character"** Sets the modes from a character vector of mode names, with the corresponding track names given in the names attribute.

## See Also

trackNames and trackNames<- for just getting or setting which tracks are visible (not of mode "hide").

# Examples

```
# Tracks "foo" and "bar" are fully shown, "baz" is hidden
modes <- ucscTrackModes(full = c("foo", "bar"), hide = "baz")
# Update the modes to hide track "bar"
modes2 <- ucscTrackModes(modes, hide = "bar")</pre>
```

UCSCView-class Class "UCSCView"

# Description

An object representing a view of a genome in the UCSC browser.

## **Objects from the Class**

Objects are created by calling browserView (session, ...) where session is a UCSCSession.

# Slots

hgsid: Object of class "numeric", which identifies this view to UCSC.

session: Object of class "BrowserSession" to which this view belongs.

# Extends

Class "BrowserView", directly.

# Methods

activeView (object) Obtains a logical indicating whether this view is the active view.

range (object) Obtains the RangesList displayed by this view.

range(object) <- value Sets the RangesList displayed by this view.</pre>

trackNames (object) Gets the names of the visible tracks in this view.

trackNames(object) <- value Sets the visible tracks by name.</pre>

visible (object) Get a named logical vector indicating whether each track is visible.

visible(object) <- value Set a logical vector indicating the visibility of each track, in the same order as returned by visible(object).

ucscTrackModes (object) Obtains the UCSCTrackModes for this view.

ucscTrackModes(object) <- value Sets the UCSCTrackModes for this view. The value may be either a UCSCTrackModes instance or a character vector that will be coerced by a call to ucscTrackModes.

## Author(s)

Michael Lawrence

#### See Also

browserView for creating instances of this class.

GraphTrackLine-class

Class "GraphTrackLine"

# Description

A UCSC track line for graphical tracks.

## **Objects from the Class**

Objects can be created by calls of the form new ("GraphTrackLine", ...) or parsed from a character vector track line with as (text, "GraphTrackLine") or converted from a BasicTrackLine using as (basic, "GraphTrackLine").

# Slots

altColor: Object of class "integer" giving an alternate color, as from col2rgb.

autoScale: Object of class "logical" indicating whether to automatically scale to min/max of the data.

gridDefault: Object of class "logical" indicating whether a grid should be drawn.

- maxHeightPixels: Object of class "numeric" of length three (max, default, min), giving the allowable range for the vertical height of the graph.
- graphType: Object of class "character", specifying the graph type, either "bar" or "points".
- viewLimits: Object of class "numeric" and of length two specifying the data range (min, max) shown in the graph.
- yLineMark: Object of class "numeric" giving the position of a horizontal line.
- yLineOnOff: Object of class "logical" indicating whether the yLineMark should be visible.
- windowingFunction: Object of class "character", one of "maximum", "mean", "minimum", for removing points when the graph shrinks.
- smoothingWindow: Object of class "numeric" giving the window size of a smoother to pass
  over the graph.
- type: Scalar "character" indicating the type of the track, either "wig" or "bedGraph".
- name: Object of class "character" specifying the name of the track.
- description: Object of class "character" describing the track.
- visibility: Object of class "character" indicating the default visible mode of the track, see UCSCTrackModes.
- color: Object of class "integer" representing the track color (as from col2rgb).
- priority: Object of class "numeric" specifying the rank of this track.

## Extends

Class "TrackLine", directly.

# GraphTrackLine-class

# Methods

as(object, "character") Export line to its string representation.

**as(object,** "BasicTrackLine") Convert this line to a basic UCSC track line, using defaults for slots not held in common.

## Author(s)

Michael Lawrence

# References

Official documentation: http://genome.ucsc.edu/goldenPath/help/wiggle.html.

# See Also

export.wig, export.bedGraph for exporting graphical tracks.

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