Package ‘tracktables’

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Type Package
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Description Methods to create complex IGV genome browser sessions and
dynamic IGV reports in HTML pages.
biocViews Sequencing, ReportWriting
VignetteBuilder knitr
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R topics documented:

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Parameters for displaying bigwigs, bams and intervals in IGV

Description

Use `igvParam(object)` to create a parameter object to control IGV display invoked from `make-tracktable()` report or from `makeIGVSession()` XML files. Parameters for bigwig, bam and intervals files may be provided. See IGV manual for a full description of options.

Usage

```
igvParam(bigwig.altColor = "darkgrey", bigwig.color = "darkgrey",
        bigwig.autoScale = "true", bigwig.displayMode = "COLLAPSED",
        bigwig.featureVisibilityWindow = -1, bigwig.fontSize = 10,
        bigwig.normalize = "false", bigwig.renderer = "BAR_CHART",
        bigwig.sortable = "true", bigwig.visible = "true",
        bigwig.windowFunction = "mean", bigwig.baseline = 0,
        bigwig.drawBaseline = "true", bigwig.flipAxis = "false",
        bigwig.maximum = 50, bigwig.minimum = 0, bigwig.type = "LINEAR",
        interval.altColor = "darkgrey", interval.color = "darkgrey",
        interval.autoScale = "true", interval.displayMode = "character",
        interval.featureVisibilityWindow = -1, interval.fontSize = 10,
        interval.height = 40, interval.normalise = "false",
        interval.renderer = "BASIC_FEATURE", interval.sortable = "true",
        interval.visible = "true", interval.windowFunction = "count",
        bam.altColor = "darkgrey", bam.color = "darkgrey",
        bam.autoScale = "true", bam.displayMode = "EXPANDED",
        bam.featureVisibilityWindow = -1, bam.fontSize = 10,
        bam.showSpliceJunctions = "false", bam.colorByTag = "",
        bam.colorOption = "UNEXPECTED_PAIR", bam.flagUnmappedPairs = "false",
        bam.groupByTag = "", bam.maxInsertSize = 1000, bam.minInsertSize = 50,
        bam.shadeBasesOption = "QUALITY", bam.shadeCenters = "true",
        bam.showAllBases = "false", bam.sortByTag = ""
)
```
igvParam

bam.showSpliceJunctions = "false", bam.colorByTag = "",
bam.colorOption = "UNEXPECTED_PAIR", bam.flagUnmappedPairs = "false",
bam.groupByTag = "", bam.maxInsertSize = 1000, bam.minInsertSize = 50,
bam.shadeBasesOption = "QUALITY", bam.shadeCenters = "true",
bam.showAllBases = "false", bam.sortByTag = ""

Arguments

bigwig.altColor
A character vector of alternate colour for bigwigs displayed in IGV.

bigwig.color
A character vector of main colour for bigwigs displayed in IGV.

bigwig.autoScale
A character vector ("true"/"false") to indicate whether data is autoscaled for big-
wigs displayed in IGV (Default "true").

bigwig.displayMode
A character vector of display mode for bigwigs displayed in IGV.

bigwig.featureVisibilityWindow
A numeric vector of feature visibility window for bigwigs displayed in IGV
(Default -1).

bigwig.fontSize
A numeric vector of font size for bigwigs displayed in IGV.

bigwig.normalize
A character vector ("true"/"false") to indicate whether data is normalised for
bigwigs displayed in IGV (Default "false").

bigwig.renderer
A character vector of renderer for bigwigs displayed in IGV (Default "BAR_CHART").

bigwig.sortable
A character vector ("true"/"false") to indicate whether data is sortable for big-
wigs displayed in IGV (Default "true").

bigwig.visible
A character vector ("true"/"false") to indicate whether data is visible for bigwigs
displayed in IGV (Default "true").

bigwig.windowFunction
A character vector ("true"/"false") of window function for bigwigs displayed in
IGV (Default "mean").

bigwig.baseline
A numeric vector of baseline bigwigs displayed in IGV.

bigwig.drawBaseline
A character vector ("true"/"false") of whether to draw baseline for bigwigs
displayed in IGV (Default "true").

bigwig.flipAxis
A character vector ("true"/"false") to indicate whether to flip axis for bigwigs
displayed in IGV (Default "false").

bigwig.maximum
A numeric vector of maximum value to display for bigwigs displayed in IGV
(Default 50).

bigwig.minimum
A numeric vector of minimum value to display for bigwigs displayed in IGV
(Default 0).

bigwig.type
A character vector of display type for bigwigs displayed in IGV (Default "LIN-
EAR").
interval.altColor
A character vector of alternate colour for intervals displayed in IGV.

interval.color
A character vector of main colour for intervals displayed in IGV.

interval.autoScale
A character vector ("true"/"false") to indicate whether data is autoscaled for intervals displayed in IGV (Default "false").

interval.displayMode
A character vector of display mode for intervals displayed in IGV (Default "COLLAPSED").

interval.featureVisibilityWindow
A numeric vector of feature visibility window for intervals displayed in IGV (Default -1).

interval.fontSize
A numeric vector of font size for intervals displayed in IGV.

interval.height
A numeric vector of height for intervals displayed in IGV.

interval.normalize
A character vector ("true"/"false") to indicate whether data is normalised for intervals displayed in IGV (Default "false").

interval.renderer
A character vector of renderer for intervals displayed in IGV (Default "BASIC_FEATURE").

interval.sortable
A character vector ("true"/"false") to indicate whether data is sortable for intervals displayed in IGV (Default "true").

interval.visible
A character vector ("true"/"false") to indicate whether data is visible for intervals displayed in IGV (Default "true").

interval.windowFunction
A character vector ("true"/"false") of window function for intervals displayed in IGV (Default "count").

bam.altColor
A character vector of alternate colour for bam files displayed in IGV.

bam.color
A character vector of main colour for bam files displayed in IGV.

bam.autoScale
A character vector ("true"/"false") to indicate whether data is autoscaled for bam files displayed in IGV (Default "false").

bam.displayMode
A character vector of display mode for bam files displayed in IGV (Default "EXPANDED").

bam.featureVisibilityWindow
A numeric vector of feature visibility window for bam files displayed in IGV (Default -1).

bam.fontSize
A numeric vector of font size for intervals displayed in IGV.

bam.showSpliceJunctions
A character vector ("true"/"false") to indicate whether to show splice junctions for bam files displayed in IGV (Default "false").

bam.colorByTag
A character vector to indicate whether to colour reads by Tags for Bam files (Default = "").
**igvParam**

bam.colorOption  
A character vector of option to highlight Tags for Bam files (Default = "UNEXPECTED_PAIR").

bam.flagUnmappedPairs  
A character vector ("true"/"false") to indicate whether to flag unmapped pairs for bam files displayed in IGV (Default "false").

bam.groupByTag  
A character vector ("true"/"false") to indicate how to groups reads by Tag for bam files displayed in IGV (Default ")

bam.maxInsertSize  
A numeric vector of maximum insert size to display for Bam files.

bam.minInsertSize  
A numeric vector of minimum insert size to display for Bam files.

bam.shadeBasesOption  
A character vector of option to shade bases for Bam files (Default "QUALITY").

bam.shadeCenters  
A character vector ("true"/"false") to indicate whether to shade centres for bam files displayed in IGV (Default "false").

bam.showAllBases  
A character vector ("true"/"false") to indicate to show all bases for bam files displayed in IGV (Default "false").

bam.sortByTag  
A character vector ("true"/"false") to indicate how to sort reads by Tag for bam files displayed in IGV (Default "").

**Value**

An igvParam class object to use with maketracktable, MakeIGVSession and makeIGVSessionMXL

**Author(s)**

Thomas Carroll

**Examples**

```r
## Simple initialisation of an IGVParam object  
igvDisplayParams <- igvParam()

## More custom initialisation of an IGVParam object .  
igvDisplayParams <- igvParam(bigwig.color="red",bigwig.autoScale = "false",  
bigwig.minimum = 10,bigwig.maximum = 100)

# See full parameters and IGV online manual for more details on customistions

## Use igvParams with maketracktables function to customise bigwig display colour and data range.
fileLocations <- system.file("extdata",package="tracktables")
bigwigs <- dir(fileLocations,pattern="*.bw",full.names=TRUE)
intervals <- dir(fileLocations,pattern="*.bed",full.names=TRUE)
bigWigMat <- cbind(gsub("_Example.bw","",basename(bigwigs)),
```

```
bigwigs)

intervalsMat <- cbind(gsub("_Peaks.bed","",basename(intervals)),
                   intervals)

fileSheet <- merge(bigWigMat,intervalsMat,all=TRUE)

fileSheet <- as.matrix(cbind(fileSheet,NA))

colnames(fileSheet) <- c("SampleName","bigwig","interval","bam")

SampleSheet <- cbind(as.vector(fileSheet[,"SampleName"]),
                      c("EBF","H3K4me3","H3K9ac","RNAPol2"),
                      c("ProB","ProB","ProB","ProB"))

colnames(SampleSheet) <- c("SampleName","Antibody","Species")

MakeIGVSession(SampleSheet,fileSheet,
               igvdirectory=getwd(),"Example","mm9",
               igvParams=igvDisplayParams)

---

### Intervals

**Example genomic intervals**

<table>
<thead>
<tr>
<th>Intervals</th>
<th>Example genomic intervals</th>
</tr>
</thead>
</table>

**Description**

This dataset contains peaks from an in-house EBF1 ChIP-seq

**Usage**

data(Intervals)

**Details**

- Intervals GRanges object containing EBF1 peaks

**Value**

A GRanges object with two rows

---

### makebedtable

**Make HTML pages for interval files or GRanges.**

**Description**

Creates HTML pages for interval files or GRanges (Tracktables Interval Report).

**Usage**

makebedtable(grangesObject, name, basedirectory)
**MakeIGVSampleMetadata**

Make sample metadata file for use with IGV.

**Description**

Creates sample metadata file for IGV from a Samplesheet of metadata and FileSheet of file locations.

**Usage**

MakeIGVSampleMetadata(SampleSheet, fileSheet, igvDirectory)

**Arguments**

- **SampleSheet**  
  A data.frame object with metadata information for samples. First column must contain unique sample ids.

- **fileSheet**  
  A data.frame of file locations. First column must contain the unique sample ids.

- **igvDirectory**  
  A character of the directory to which sample metadata file is written.

**Value**

A character of file location for the IGV sample information file.

**Author(s)**

Thomas Carroll
Examples

```r
fileLocations <- system.file("extdata", package="tracktables")
bigwigs <- dir(fileLocations, pattern="*.bw", full.names=TRUE)
intervals <- dir(fileLocations, pattern="*.bed", full.names=TRUE)
bigWigMat <- cbind(gsub("_Example.bw","", basename(bigwigs)),
          bigwigs)
intervalsMat <- cbind(gsub("_Peaks.bed","", basename(intervals)),
          intervals)
fileSheet <- merge(bigWigMat, intervalsMat, all=TRUE)
fileSheet <- as.matrix(cbind(fileSheet, NA))
colnames(fileSheet) <- c("SampleName", "bigwig", "interval", "bam")
SampleSheet <- cbind(as.vector(fileSheet[, "SampleName"]),
          c("EBF", "H3K4me3", "H3K9ac", "RNAPol2"),
          c("ProB", "ProB", "ProB", "ProB"))
colnames(SampleSheet) <- c("SampleName", "Antibody", "Species")
MakeIGVSampleMetadata(SampleSheet, fileSheet, igvdirectory=getwd())
```

Description

Make IGV session XML and sample information file from a Samplesheet of metadata and FileSheet of file locations.

Usage

```r
MakeIGVSession(SampleSheet, fileSheet, igvdirectory, XMLname, genomeName,
      locusName = "All", colourBy = NULL, igvParams = igvParam(),
      writedirectory = NULL, full.xml.paths = FALSE, full.file.paths = FALSE,
      use.path.asis = FALSE)
```

Arguments

- **SampleSheet**: A data.frame object with metadata information for samples. First column must contain unique sample ids.
- **fileSheet**: A data.frame of file locations. First column must contain the unique sample ids.
- **igvdirectory**: A character of the directory to which sample metadata file is written.
- **XMLname**: A character of the name for IGV session xml
- **genomeName**: A character of genome for IGV (See IGV user guide for details)
- **locusName**: A character of locus to display in igv on loading (See IGV user guide for details)
- **colourBy**: Character defining which sample metadata to be used for colouring bigwig files
**MakeIGVSession**

**igvParams**  
An object of class igvParam containing display parameters for IGV. When providing a list, this list must be same length as number of samples and each element have two numeric values corresponding to minimum and maximum value to be used in setting data range. Currently only "autoscale" or a list of minimum and maximum values are accepted.

**writedirectory**  
A character of the directory to which files will be written. Default is set to igvdirectory argument.

**full.xml.paths**  
Boolean of whether reference to XML and sample information files should be by relative or absolute paths. Default is FALSE

**full.file.paths**  
Boolean of whether reference to sample bigWig/Bam/interval files should be by relative or absolute paths. Default is FALSE

**use.path.asis**  
Boolean of whether paths to files in samplesheet should be used as is. If TRUE overrides sample files paths specified by full.file.paths argument. Default is FALSE

**Value**  
A character of file location for the IGV session XML

**Author(s)**  
Thomas Carroll

**Examples**

```r
fileLocations <- system.file("extdata",package="tracktables")
bigwigs <- dir(fileLocations,pattern="*.bw",full.names=TRUE)
intervals <- dir(fileLocations,pattern="*.bed",full.names=TRUE)
bigWigMat <- cbind(gsub("_Example.bw","",basename(bigwigs)),
bigwigs)
intervalsMat <- cbind(gsub("_Peaks.bed","",basename(intervals)),
intervals)
fileSheet <- merge(bigWigMat,intervalsMat,all=TRUE)
fileSheet <- as.matrix(cbind(fileSheet,NA))
colnames(fileSheet) <- c("SampleName","bigwig","interval","bam")
SampleSheet <- cbind(as.vector(fileSheet[,"SampleName"]),
c("EBF","H3K4me3","H3K9ac","RNAPol2"),
c("ProB","ProB","ProB","ProB")
colnames(SampleSheet) <- c("SampleName","Antibody","Species")
MakeIGVSession(SampleSheet,fileSheet,igvdirectory=getwd(),"Example","mm9")
```
MakeIGVSessionXML

Description

Creates session XML for IGV from a FileSheet of file locations.

Usage

MakeIGVSessionXML(fileSheet, igvdirectory, XMLname, genomeName,  
locusName = "All", colourBy = NULL, igvParams = igvParam(), 
writedirectory = NULL, full.xml.paths = FALSE, full.file.paths = FALSE,  
use.path.asis = FALSE)

Arguments

fileSheet A data.frame of file locations. First column must contain the unique sample ids.
igvdirectory A character of the directory to which IGV XML session is written.
XMLname A character of the name for IGV session xml
genomeName A character of genome for IGV (See IGV user guide for details)
locusName A character of locus to display in igv on loading (See IGV user guide for details)
colourBy Character vector of RGB colours to use for colouring displayed BigWigs
igvParams An object of class igvParam containing display parameters for IGV. When providing a list, this list must be same length as number of samples and each element have two numeric values corresponding to minimum and maximum value to be used in setting data range. Currently only "autoscale" or a list of minimum and maximum values are accepted.
writedirectory A character of the directory to which files will be written. Default is set to igvdirectory argument.
full.xml.paths Boolean of whether reference to XML and sample information files should be by relative or absolute paths. Default is FALSE
full.file.paths Boolean of whether reference to sample bigWig/Bam/interval files should be by relative or absolute paths. Default is FALSE
use.path.asis Boolean of whether paths to files in samplesheet should be used as is. If TRUE overrides sample files paths specified by full.file.paths argument. Default is FALSE

Value

A character of file location for the IGV session XML

Author(s)

Thomas Carroll
Examples

```r
fileLocations <- system.file("extdata",package="tracktables")
bigwigs <- dir(fileLocations,pattern="*.bw",full.names=TRUE)
intervals <- dir(fileLocations,pattern="*.bed",full.names=TRUE)
bigWigMat <- cbind(gsub("_Example.bw","",basename(bigwigs)),
                  bigwigs)
intervalsMat <- cbind(gsub("_Peaks.bed","",basename(intervals)),
                     intervals)
fileSheet <- merge(bigWigMat,intervalsMat,all=TRUE)
fileSheet <- as.matrix(cbind(fileSheet,NA))
colnames(fileSheet) <- c("SampleName","bigwig","interval","bam")
MakeIGVSessionXML(fileSheet,igvdirectory=getwd(),"Example","mm9")
```

### Description

Creates HTML table of sample metadata and all required files for interacting with IGV.

### Usage

```r
maketracktable(fileSheet, SampleSheet, filename, basedirectory, genome,
                colourBy = NULL, igvParams = igvParam(), writedirectory = NULL,
                full.xml.paths = FALSE, full.file.paths = FALSE, use.path.asis = FALSE)
```

### Arguments

- **fileSheet**: A data frame containing sample file locations (e.g. BigWig locations).
- **SampleSheet**: A data frame containing sample metadata
- **filename**: Character of name for tracktables HTML report. (.html prefix is added automatically)
- **basedirectory**: Character of directory for tracktables HTML report, IGV sessions and any interval files
- **genome**: Character of genome for IGV (See IGV user guide for details)
- **colourBy**: Character defining which sample metadata to be used for colouring bigwig files
- **igvParams**: An object of class igvParam containing display parameters for IGV. When providing a list, this list must be same length as number of samples and each element have two numeric values corresponding to minimum and maximum value to be used in setting data range. Currently only "autoscale" or a list of minimum and maximum values are accepted.
maketracktable

writedirectory A character of the directory to which files will be written. Default is set to igvdirectory argument.

full.xml.paths Boolean of whether reference to XML and sample information files should be by relative or absolute paths. Default is FALSE

full.file.paths Boolean of whether reference to sample bigWig/Bam/interval files should be by relative or absolute paths. Default is FALSE

use.path.asis Boolean of whether paths to files in samplesheet should be used as is. If TRUE overrides sample files paths specified by full.file.paths argument. Default is FALSE

Value

Author(s)
Thomas Carroll

Examples

code:

fileLocations <- system.file("extdata",package="tracktables")
bigwigs <- dir(fileLocations,pattern="*.bw",full.names=TRUE)
intervals <- dir(fileLocations,pattern="*.bed",full.names=TRUE)
bigWigMat <- cbind(gsub("_Example.bw"","",basename(bigwigs)),
                  bigwigs)
intervalsMat <- cbind(gsub("_Peaks.bed"","",basename(intervals)),
                      intervals)
fileSheet <- merge(bigWigMat,intervalsMat,all=TRUE)
fileSheet <- as.matrix(cbind(fileSheet,NA))

colnames(fileSheet) <- c("SampleName","bigwig","interval","bam")
SampleSheet <- cbind(as.vector(fileSheet[,"SampleName"]),
                     c("EBF","H3K4me3","H3K9ac","RNAPol2"),
                     c("ProB","ProB","ProB","ProB"))

colnames(SampleSheet) <- c("SampleName","Antibody","Species")

HTMLreport <- maketracktable(fileSheet,SampleSheet,
                             "IGV_Example.html",
                             basedirectory=getwd(),
                             "mm9")
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