

# Package ‘wiggleplotr’

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**Title** Make read coverage plots from BigWig files

**Version** 1.32.0

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**Description** Tools to visualise read coverage from sequencing experiments together with genomic annotations (genes, transcripts, peaks). Introns of long transcripts can be rescaled to a fixed length for better visualisation of exonic read coverage.

**Depends** R (>= 3.6)

**Imports** dplyr, ggplot2 (>= 2.2.0), GenomicRanges, rtracklayer, cowplot, assertthat, purrr, S4Vectors, IRanges, GenomeInfoDb

**License** Apache License 2.0

**LazyData** true

**RoxygenNote** 6.1.1

**Suggests** knitr, rmarkdown, biomaRt, GenomicFeatures, testthat, ensemblldb, EnsDb.Hsapiens.v86, org.Hs.eg.db, TxDb.Hsapiens.UCSC.hg38.knownGene, AnnotationDbi, AnnotationFilter

**VignetteBuilder** knitr

**biocViews** ImmunoOncology, Coverage, RNASeq, ChIPSeq, Sequencing, Visualization, GeneExpression, Transcription, AlternativeSplicing

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`getGenotypePalette`      *Returns a three-colour palette suitable for visualising read coverage stratified by genotype*

### Description

Returns a three-colour palette suitable for visualising read coverage stratified by genotype

### Usage

```
getGenotypePalette(old = FALSE)
```

### Arguments

`old`      Return old colour palette (now deprecated).

### Value

Vector of three colours.

### Examples

```
getGenotypePalette()
```

---

|                   |  |
|-------------------|--|
| makeManhattanPlot | <i>Make a Manhattan plot of p-values</i> |
|-------------------|--|

---

## Description

The Manhattan plots is compatible with wigggleplotr read coverage and transcript strucutre plots. Can be appended to those using the cowplot::plot\_grid() function.

## Usage

```
makeManhattanPlot(pvalues_df, region_coords, color_R2 = FALSE,  
                  data_track = TRUE)
```

## Arguments

|               |   |
|---------------|---|
| pvalues_df    | Data frame of association p-values (required columns: track_id, p_nominal, pos)   |
| region_coords | Start and end coordinates of the region to plot.  |
| color_R2      | Color the points according to R2 from the lead variant. Require R2 column in the pvalues_df data frame.   |
| data_track    | If TRUE, then remove all information from x-axis. Makes it easy to append to read coverage or transcript strcture plots using cowplot::plot_grid(). |

## Value

ggplot2 object

## Examples

```
data = dplyr::data_frame(track_id = "GWAS", pos = sample(c(1:1000), 200), p_nominal = runif(200, min = 0.0000001, 1  
makeManhattanPlot(data, c(1,1000), data_track = FALSE)
```

---

|            |  |
|------------|--|
| ncoa7_cdss | <i>Coding sequences from 9 protein coding transcripts of NCOA7</i> |
|------------|--|

---

## Description

A dataset containing start and end coordinates of coding sequences (CDS) from nine protein coding transcripts of NCOA7.

## Usage

```
ncoa7_cdss
```

**Format**

A GRangesList object with 9 elements:

**element** CDS start and end coordinates for a single transcript (GRanges object) ...

**Source**

<http://www.ensembl.org/>

---

ncoa7\_exons

*Exons from 9 protein coding transcripts of NCOA7*

---

**Description**

A dataset containing start and end coordinates of exons from nine protein coding transcripts of NCOA7.

**Usage**

`ncoa7_exons`

**Format**

A GRangesList object with 9 elements:

**element** Exon start and end coordinates for a single transcript (GRanges object) ...

**Source**

<http://www.ensembl.org/>

---

ncoa7\_metadata

*Gene metadata for NCOA7*

---

**Description**

A a list of transcripts for NCOA7.

**Usage**

`ncoa7_metadata`

**Format**

A data.frame object with 4 columns:

**transcript\_id** Ensembl transcript id.  
**gene\_id** Ensembl gene id.  
**gene\_name** Human readable gene name.  
**strand** Strand of the transcript (either +1 or -1). ...

**Source**

<http://www.ensembl.org/>

---

pasteFactors

*Paste two factors together and preserved their joint order.*

---

**Description**

Paste two factors together and preserved their joint order.

**Usage**

```
pasteFactors(factor1, factor2)
```

**Arguments**

|         |               |
|---------|---------------|
| factor1 | First factor  |
| factor2 | Second factor |

**Value**

Factors factor1 and factor2 pasted together.

---

plotCoverage

*Plot read coverage across genomic regions*

---

**Description**

Also supports rescaling introns to constant length. Does not work on Windows, because rtracklayer cannot read BigWig files on Windows.

**Usage**

```
plotCoverage(exons, cdss = NULL, transcript_annotations = NULL,
            track_data, rescale_introns = TRUE, new_intron_length = 50,
            flanking_length = c(50, 50), plot_fraction = 0.1, heights = c(0.75,
            0.25), alpha = 1, fill_palette = c("#a1dab4", "#41b6c4", "#225ea8"),
            mean_only = TRUE, connect_exons = TRUE, transcript_label = TRUE,
            return_subplots_list = FALSE, region_coords = NULL,
            coverage_type = "area")
```

**Arguments**

|                                     |   |
|-------------------------------------|---|
| <code>exons</code>                  | list of GRanges objects, each object containing exons for one transcript. The list must have names that correspond to transcript_id column in transcript_annotations data.frame.  |
| <code>cdss</code>                   | list of GRanges objects, each object containing the coding regions (CDS) of a single transcript. The list must have names that correspond to transcript_id column in transcript_annotations data.frame. If cdss is not specified then exons list will be used for both arguments. (default: NULL).  |
| <code>transcript_annotations</code> | Data frame with at least three columns: transcript_id, gene_name, strand. Used to construct transcript labels. (default: NULL)  |
| <code>track_data</code>             | data.frame with the metadata for the bigWig read coverage files. Must contain the following columns: <ul style="list-style-type: none"> <li>• sample_id - unique id for each sample.</li> <li>• track_id - if multiple samples (bigWig files) have the same track_id they will be overlayed on the same plot, track_id is also used as the facet label on the right.</li> <li>• bigWig - path to the bigWig file.</li> <li>• scaling_factor - normalisation factor for each sample, useful if different samples sequenced to different depth and bigWig files not normalised for that.</li> <li>• colour_group - additional column to group samples into, is used as the colour of the coverage track.</li> </ul> |
| <code>rescale_introns</code>        | Specifies if the introns should be scaled to fixed length or not. (default: TRUE)   |
| <code>new_intron_length</code>      | length (bp) of introns after scaling. (default: 50)   |
| <code>flanking_length</code>        | Lengths of the flanking regions upstream and downstream of the gene. (default: c(50,50))  |
| <code>plot_fraction</code>          | Size of the random sub-sample of points used to plot coverage (between 0 and 1). Smaller values make plotting significantly faster. (default: 0.1)  |
| <code>heights</code>                | Specifies the proportion of the height that is dedicated to coverage plots (first value) relative to transcript annotations (second value). (default: c(0.75,0.25))   |

|                                   |   |
|-----------------------------------|---|
| <code>alpha</code>                | Transparency (alpha) value for the read coverage tracks. Useful to set to something < 1 when overlaying multiple tracks (see <code>track_id</code> ). (default: 1)  |
| <code>fill_palette</code>         | Vector of fill colours used for the coverage tracks. Length must be equal to the number of unique values in <code>track_data\$colour_group</code> column.   |
| <code>mean_only</code>            | Plot only mean coverage within each combination of <code>track_id</code> and <code>colour_group</code> values. Useful for example for plotting mean coverage stratified by genotype (which is specified in the <code>colour_group</code> column) (default: TRUE). |
| <code>connect_exons</code>        | Print lines that connect exons together. Set to FALSE when plotting peaks (default: TRUE).  |
| <code>transcript_label</code>     | If TRUE then transcript labels are printed above each transcript. (default: TRUE).  |
| <code>return_subplots_list</code> | Instead of a joint plot return a list of subplots that can be joined together manually.   |
| <code>region_coords</code>        | Start and end coordinates of the region to plot, overrides <code>flanking_length</code> parameter.  |
| <code>coverage_type</code>        | Specifies if the read coverage is represented by either 'line', 'area' or 'both'. The 'both' option tends to give better results for wide regions. (default: area).   |

## Value

Either object from `cow_plot::plot_grid()` function or a list of subplots (if `return_subplots_list == TRUE`)

## Examples

```
require("dplyr")
require("GenomicRanges")
sample_data = dplyr::data_frame(sample_id = c("aipt_A", "aipt_C", "bima_A", "bima_C"),
  condition = factor(c("Naive", "LPS", "Naive", "LPS"), levels = c("Naive", "LPS")),
  scaling_factor = 1) %>%
  dplyr::mutate(bigWig = system.file("extdata", paste0(sample_id, ".str2.bw")), package = "wiggleplotr"))

track_data = dplyr::mutate(sample_data, track_id = condition, colour_group = condition)

selected_transcripts = c("ENST00000438495", "ENST00000392477") #Plot only two transcripts of the gens
## Not run:
plotCoverage(ncoa7_exons[selected_transcripts], ncoa7_cdss[selected_transcripts],
  ncoa7_metadata, track_data,
  heights = c(2,1), fill_palette = getGenotypePalette())

## End(Not run)
```

**plotCoverageFromEnsemblDb**

*Plot read coverage directly from ensemblDb object.*

**Description**

A wrapper around the `plotCoverage` function. See the documentation for ([plotCoverage](#)) for more information.

**Usage**

```
plotCoverageFromEnsemblDb(ensemblDb, gene_names, transcript_ids = NULL,
...)
```

**Arguments**

|                             |   |
|-----------------------------|---|
| <code>ensemblDb</code>      | ensemblDb object.   |
| <code>gene_names</code>     | List of gene names to be plotted.                                 |
| <code>transcript_ids</code> | Optional list of transcript ids to be plotted.                    |
| <code>...</code>            | Additional parameters to be passed to <code>plotCoverage</code> . |

**Value**

`ggplot2` object

**Examples**

```
require("EnsDb.Hsapiens.v86")
require("dplyr")
require("GenomicRanges")
sample_data = dplyr::data_frame(sample_id = c("aipt_A", "aipt_C", "bima_A", "bima_C"),
condition = factor(c("Naive", "LPS", "Naive", "LPS"), levels = c("Naive", "LPS")),
scaling_factor = 1) %>%
dplyr::mutate(bigWig = system.file("extdata", paste0(sample_id, ".str2.bw"), package = "wiggleplotr"))

track_data = dplyr::mutate(sample_data, track_id = condition, colour_group = condition)
## Not run:
plotCoverageFromEnsemblDb(EnsDb.Hsapiens.v86, "NC0A7", transcript_ids = c("ENST00000438495", "ENST00000392477"),
track_data, heights = c(2,1), fill_palette = getGenotypePalette())

## End(Not run)
```

---

plotCoverageFromUCSC *Plot read coverage directly from UCSC OrgDb and TxDb objects.*

---

## Description

A wrapper around the plotCoverage function. See the documentation for ([plotCoverage](#)) for more information.

## Usage

```
plotCoverageFromUCSC(orgdb, txdb, gene_names, transcript_ids = NULL, ...)
```

## Arguments

|                |   |
|----------------|---|
| orgdb          | UCSC OrgDb object.                                  |
| txdb           | UCSC TxDb obejct.                                   |
| gene_names     | List of gene names to be plotted.                   |
| transcript_ids | Optional list of transcript ids to be plotted.      |
| ...            | Additional parameters to be passed to plotCoverage. |

## Value

ggplot2 object

## Examples

```
require("dplyr")
require("GenomicRanges")
require("org.Hs.eg.db")
require("TxDb.Hsapiens.UCSC.hg38.knownGene")

orgdb = org.Hs.eg.db
txdb = TxDb.Hsapiens.UCSC.hg38.knownGene

sample_data = dplyr::data_frame(sample_id = c("aipt_A", "aipt_C", "bima_A", "bima_C"),
  condition = factor(c("Naive", "LPS", "Naive", "LPS"), levels = c("Naive", "LPS")),
  scaling_factor = 1) %>%
  dplyr::mutate(bigWig = system.file("extdata", paste0(sample_id, ".str2.bw")), package = "wiggleplotr"))

track_data = dplyr::mutate(sample_data, track_id = condition, colour_group = condition)
## Not run:
#Note: This example does not work, becasue UCSC and Ensembl use different chromosome names
plotCoverageFromUCSC(orgdb, txdb, "NCOA7", transcript_ids = c("ENST00000438495.6", "ENST00000368357.7"),
  track_data, heights = c(2,1), fill_palette = getGenotypePalette())

## End(Not run)
```

**plotTranscripts**      *Quickly plot transcript structure without read coverage tracks*

## Description

Quickly plot transcript structure without read coverage tracks

## Usage

```
plotTranscripts(exons, cdss = NULL, transcript_annotations = NULL,
  rescale_introns = TRUE, new_intron_length = 50,
  flanking_length = c(50, 50), connect_exons = TRUE,
  transcript_label = TRUE, region_coords = NULL)
```

## Arguments

|                                     |   |
|-------------------------------------|---|
| <code>exons</code>                  | list of GRanges objects, each object containing exons for one transcript. The list must have names that correspond to transcript_id column in transcript_annotations data.frame.  |
| <code>cdss</code>                   | list of GRanges objects, each object containing the coding regions (CDS) of a single transcript. The list must have names that correspond to transcript_id column in transcript_annotations data.frame. If cdss is not specified then exons list will be used for both arguments. (default: NULL) |
| <code>transcript_annotations</code> | Data frame with at least three columns: transcript_id, gene_name, strand. Used to construct transcript labels. (default: NULL)  |
| <code>rescale_introns</code>        | Specifies if the introns should be scaled to fixed length or not. (default: TRUE)   |
| <code>new_intron_length</code>      | length (bp) of introns after scaling. (default: 50)   |
| <code>flanking_length</code>        | Lengths of the flanking regions upstream and downstream of the gene. (default: c(50,50))  |
| <code>connect_exons</code>          | Print lines that connect exons together. Set to FALSE when plotting peaks (default: TRUE).  |
| <code>transcript_label</code>       | If TRUE then transcript labels are printed above each transcript. (default: TRUE).  |
| <code>region_coords</code>          | Start and end coordinates of the region to plot, overrides flanking_length parameter.   |

## Value

ggplot2 object

## Examples

```
plotTranscripts(ncoa7_exons, ncoa7_cdss, ncoa7_metadata, rescale_introns = FALSE)
```

---

### plotTranscriptsFromEnsemblDb

*Plot transcripts directly from ensemblDb object.*

---

## Description

A wrapper around the plotTranscripts function. See the documentation for ([plotTranscripts](#)) for more information.

## Usage

```
plotTranscriptsFromEnsemblDb(ensemblDb, gene_names,  
                             transcript_ids = NULL, ...)
```

## Arguments

|                |   |
|----------------|---|
| ensemblDb      | ensemblDb object.                                     |
| gene_names     | List of gene names to be plotted.                     |
| transcript_ids | Optional list of transcript ids to be plotted.        |
| ...            | Additional parameters to be passed to plotTranscripts |

## Value

ggplot2 object

## Examples

```
require("EnsDb.Hsapiens.v86")  
plotTranscriptsFromEnsemblDb(EnsDb.Hsapiens.v86, "NCOA7", transcript_ids = c("ENST00000438495", "ENST00000392477"))
```

---

**plotTranscriptsFromUCSC**

*Plot transcripts directly from UCSC OrgDb and TxDb objects.*

---

**Description**

A wrapper around the `plotTranscripts` function. See the documentation for ([plotTranscripts](#)) for more information. Note that this function is much slower than ([plotTranscripts](#)) or ([plotTranscriptsFromEnsemblDb](#)) functions, because individually extracting exon coordinates from txdb objects is quite inefficient.

**Usage**

```
plotTranscriptsFromUCSC(orgdb, txdb, gene_names, transcript_ids = NULL,  
...)
```

**Arguments**

|                             |  |
|-----------------------------|--|
| <code>orgdb</code>          | UCSC OrgDb object.   |
| <code>txdb</code>           | UCSC TxDb obejct.  |
| <code>gene_names</code>     | List of gene genaes to be plot.                                    |
| <code>transcript_ids</code> | Optional list of transcript ids to be plot. (default = NULL)       |
| <code>...</code>            | Additional parameters to be passed to <code>plotTranscripts</code> |

**Value**

Transcript plot.

**Examples**

```
#Load OrgDb and TxDb objects with UCSC gene annotations  
require("org.Hs.eg.db")  
require("TxDb.Hsapiens.UCSC.hg38.knownGene")  
orgdb = org.Hs.eg.db  
txdb = TxDb.Hsapiens.UCSC.hg38.knownGene  
  
plotTranscriptsFromUCSC(orgdb, txdb, "NCOA7", transcript_ids = c("ENST00000438495.6", "ENST00000368357.7"))
```

---

wiggleplotr

---

*wiggleplotr*

---

## Description

wiggleplotr package provides tools to visualise transcript annotations ([plotTranscripts](#)) and plot sequencing read coverage over annotated transcripts ([plotCoverage](#)).

## Details

You can also use convenient wrapper functions ([plotTranscriptsFromEnsemblDb](#)), ([plotCoverageFromEnsemblDb](#)), ([plotTranscriptsFromUCSC](#)) and ([plotCoverageFromUCSC](#)).

To learn more about wiggleplotr, start with the vignette: `browseVignettes(package = "wiggleplotr")`

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