

Package ‘RiboCrypt’

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Type Package

Title Interactive visualization in genomics

Version 1.12.0

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Description R Package for interactive visualization and browsing NGS data.

It contains a browser for both transcript and genomic coordinate view.

In addition a QC and general metaplots are included, among others differential translation plots and gene expression plots. The package is still under development.

biocViews Software, Sequencing, RiboSeq, RNASeq,

Encoding UTF-8

LazyData true

BugReports <https://github.com/m-swirski/RiboCrypt/issues>

URL <https://github.com/m-swirski/RiboCrypt>

Depends R (>= 3.6.0), ORFik (>= 1.13.12)

Imports bslib, BiocGenerics, BiocParallel, Biostrings, data.table,
dplyr, GenomeInfoDb, GenomicFeatures, GenomicRanges, ggplot2,
htmlwidgets, httr, IRanges, jsonlite, knitr, markdown,
NGLVieweR, plotly, rlang, RCurl, shiny, shinycssloaders,
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Suggests testthat, rmarkdown, BiocStyle, BSgenome,
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Author Michal Swirski [aut, cre, cph],
Haakon Tjeldnes [aut, ctb],
Kornel Labun [ctb]

Maintainer Michal Swirski <michal.swirski@uw.edu.pl>

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antisense

Get antisense

Description

Get antisense

Usage

```
antisense(grl)
```

Value

a GRangesList

createSeqPanelPattern *Create sequence panel for RiboCrypt*

Description

Create sequence panel for RiboCrypt

Usage

```
createSeqPanelPattern(
  sequence,
  start_codons = "ATG",
  stop_codons = c("TAA", "TAG", "TGA"),
  frame = 1,
  custom_motif = NULL
)
```

Arguments

start_codons character vector, default "ATG"
 stop_codons character vector, default c("TAA", "TAG", "TGA")
 custom_motif character vector, default NULL.

Value

a ggplot object

 DEG_plot

Differential expression plots (1D or 2D)

Description

Gives you interactive 1D or 2D DE plots

Usage

```
DEG_plot(
  dt,
  draw_non_regulated = FALSE,
  xlim = ifelse(two_dimensions, "bidir.max", "auto"),
  ylim = "bidir.max",
  xlab = ifelse(two_dimensions, "RNA fold change (log2)", "Mean counts (log2)"),
  ylab = ifelse(two_dimensions, "RFP fold change (log2)", "Fold change (log2)"),
  two_dimensions = ifelse("LFC" %in% colnames(dt), FALSE, TRUE),
  color.values = c(`No change` = "black", Significant = "red", Buffering = "purple",
    `mRNA abundance` = "darkgreen", Expression = "blue", Forwarded = "yellow", Inverse =
    "aquamarine", Translation = "orange4")
)
```

Arguments

dt a data.table with results from a differential expression run. Normally from: ORFik::DTEG.analysis(df1, df2)
 draw_non_regulated logical, default FALSE. Should non-regulated rows be included in the plot? Will make the plot faster to render if skipped (FALSE)

| | |
|----------------|---|
| xlim | numeric vector or character preset, default: <code>ifelse(two_dimensions, "bidir.max", "auto")</code> (Equal in both + / - direction, using max value + 0.5 of <code>meanCounts(in 1d) / rna(in 2d)</code> column of <code>dt</code>). If you want <code>ggplot</code> to decide limit, set to "auto". For numeric vector, specify min and max x limit: like <code>c(-5, 5)</code> |
| ylim | numeric vector or character preset, default: "bidir.max" (Equal in both + / - direction, using max value + 0.5 of <code>LFC(in 1d) / rfp(in 2d)</code> column of <code>dt</code>). If you want <code>ggplot</code> to decide limit, set to "auto". For numeric vector, specify min and max x limit: like <code>c(-5, 5)</code> |
| xlab | character, default: <code>ifelse(two_dimensions, "RNA fold change (log2)", "Mean counts (log2)")</code> |
| ylab | character, default: <code>ifelse(two_dimensions, "RFP fold change (log2)", "Fold change (log2)")</code> |
| two_dimensions | logical, default: <code>ifelse("LFC" %in% colnames(dt), FALSE, TRUE)</code> Is this two dimensional, like: Ribo-seq vs RNA-seq. Alternative, FALSE: Then Log fold change vs mean counts |
| color.values | named character vector, default: <code>c("No change" = "black", "Significant" = "red", "Buffering" = "purple", "mRNA abundance" = "darkgreen", "Expression" = "blue", "Forwarded" = "yellow", "Inverse" = "aquamarine", "Translation" = "orange4")</code> |

Value

plotly object

Examples

```
# Load experiment
df <- ORFik.template.experiment()
# 1 Dimensional analysis
dt <- DEG.analysis(df[df$libtype == "RNA",])
dt$Regulation[1] <- "Significant" # Fake sig level
DEG_plot(dt, draw_non_regulated = TRUE)
# 2 Dimensional analysis
dt_2d <- DTEG.analysis(df[df$libtype == "RFP",], df[df$libtype == "RNA",],
                      output.dir = NULL)
dt_2d$Regulation[4] <- "Translation" # Fake sig level
dt_2d$Regulation[5] <- "Buffering" # Fake sig level
DEG_plot(dt_2d, draw_non_regulated = TRUE)
```

distanceToFollowing *Distance to following range*

Description

Distance to following range

Usage

```
distanceToFollowing(gr1, grl2 = gr1, ignore.strand = FALSE)
```

Arguments

gr1 a GRangesList
gr12 a GRangesList, default 'gr1'
ignore.strand logical, default FALSE

Value

numeric vector of distance

| | |
|--------------|----------------------------------|
| fetch_JS_seq | <i>Fetch Javascript sequence</i> |
|--------------|----------------------------------|

Description

Fetch Javascript sequence

Usage

```
fetch_JS_seq(  
  target_seq,  
  nplots,  
  distance = 50,  
  display_dist,  
  aa_letter_code = "one_letter"  
)
```

Arguments

target_seq the target sequence
nplots number of plots
distance numeric, default 50.
display_dist display distance
aa_letter_code "one_letter"

Value

a list of 2 lists, the nt list (per frame, total 3) and AA list (per frame, total 3)

fetch_summary *Fetch summary of uniprot id*

Description

Fetch summary of uniprot id

Usage

```
fetch_summary(qualifier, provider = "alphafold")
```

Arguments

qualifier uniprot ids
provider "pdbe", alternatives: "alphafold", "all"

Value

a character of json

geneTrackLayer *How many rows does the gene track need*

Description

How many rows does the gene track need

Usage

```
geneTrackLayer(gr1)
```

Arguments

gr1 a GRangesList

Value

numeric, the track row index

| | |
|--------------------|-----------------------------|
| getCoverageProfile | <i>Get coverage profile</i> |
|--------------------|-----------------------------|

Description

Get coverage profile

Usage

```
getCoverageProfile(grl, reads, kmers = 1, kmers_type = "mean")
```

Arguments

| | |
|------------|---------------|
| grl | a GRangesList |
| reads | GRanges |
| kmers | 1 |
| kmers_type | "mean" |

Value

data.table of coverage

| | |
|----------|------------------|
| getIndex | <i>Get index</i> |
|----------|------------------|

Description

Get index

Usage

```
getIndex(ref_granges)
```

Arguments

| | |
|-------------|------------------|
| ref_granges | a GRanges object |
|-------------|------------------|

Value

integer vector, indices

ggplotlyHover *Call ggplotly with hoveron defined*

Description

Call ggplotly with hoveron defined

Usage

```
ggplotlyHover(x, ...)
```

Arguments

x a a ggplot argument
... additional arguments for ggplotly

Value

a ggplotly object

matchMultiplePatterns *Match multiple patterns*

Description

Match multiple patterns

Usage

```
matchMultiplePatterns(patterns, Seq)
```

Arguments

patterns character
Seq a DNAStrngSet

Value

integer vector, indices (named with pattern hit)

| | |
|----------------|-------------------------|
| matchToGRanges | <i>Match to GRanges</i> |
|----------------|-------------------------|

Description

Match to GRanges

Usage

```
matchToGRanges(matches, ref_granges)
```

Arguments

| | |
|-------------|-------------------------|
| matches | integer vector, indices |
| ref_granges | GRanges |

Value

GRanges object

| | |
|------------------------|---|
| multiOmicsPlot_animate | <i>Multi-omics animation using list input</i> |
|------------------------|---|

Description

The animation will move with a play button, there is 1 transition per library given.

Usage

```
multiOmicsPlot_animate(  
  display_range,  
  annotation = display_range,  
  reference_sequence,  
  reads,  
  viewMode = c("tx", "genomic")[1],  
  custom_regions = NULL,  
  leader_extension = 0,  
  trailer_extension = 0,  
  withFrames = NULL,  
  frames_type = "lines",  
  colors = NULL,  
  kmers = NULL,  
  kmers_type = c("mean", "sum")[1],  
  ylabels = NULL,  
  lib_to_annotation_proportions = c(0.8, 0.2),  
  lib_proportions = NULL,  
  annotation_proportions = NULL,  
  width = NULL,  
)
```

```

height = NULL,
plot_name = "default",
plot_title = NULL,
display_sequence = c("both", "nt", "aa", "none")[1],
seq_render_dist = 100,
aa_letter_code = c("one_letter", "three_letters")[1],
annotation_names = NULL,
start_codons = "ATG",
stop_codons = c("TAA", "TAG", "TGA"),
custom_motif = NULL,
BPPARAM = BiocParallel::SerialParam()
)

```

Arguments

| | |
|-------------------------------|--|
| display_range | the whole region to visualize, a GRangesList or GRanges object |
| annotation | the whole annotation which your target region is a subset, a GRangesList or GRanges object |
| reference_sequence | the genome reference, a FaFile or FaFile convertible object |
| reads | the NGS libraries, as a list of GRanges with or without score column for replicates. |
| viewMode | character, default "tx" (transcript coordinates, first position is 1, exons are merged into a single sequence) Alternative: "genomic" (genomic coordinates, first position is first position in display_range argument. Introns are displayed). |
| custom_regions | a GRangesList or NULL, default: NULL. The alternative annotation, like self defined uORFs etc. The vertical annotation bars will have a different color. |
| leader_extension | integer, default 0. (How much to extend view upstream) |
| trailer_extension | integer, default 0. (How much to extend view downstream) |
| withFrames | a logical vector, default NULL. Alternative: a length 1 or same length as list length of "reads" argument. |
| frames_type | character, default "lines". Alternative: - columns - stacks - area |
| colors | character, default NULL (automatic colouring). If "withFrames" argument is TRUE, colors are set to c("red", "green", "blue") for the 3 frames. Alternative: Character vector of length 1 or length of "reads" list argument. |
| kmers | numeric (integer), bin positions into kmers. |
| kmers_type | character, function used for kmers sliding window. default: "mean", alternative: "sum" |
| ylabels | character, default NULL. Name of libraries in "reads" list argument. |
| lib_to_annotation_proportions | numeric vector of length 2. relative sizes of profiles and annotation. |
| lib_proportions | numeric vector of length equal to displayed libs. Relative sizes of profiles displayed |

| | |
|------------------------|--|
| annotation_proportions | numeric vector of length 3 (seq displayed), or 2 (seq not displayed). Relative sizes of annotation tracks. |
| width | numeric, default NULL. Width of plot. |
| height | numeric, default NULL. Height of plot. |
| plot_name | = character, default "default" (will create name from display_range name). Alternative: custom name for region. |
| plot_title | character, default NULL. A title for plot. |
| display_sequence | character/logical, default c("both", "nt", "aa", "none")[1]. If TRUE or "both", display nucleotide and aa sequence in plot. |
| seq_render_dist | integer, default 100. The sequences will appear after zooming below this threshold. |
| aa_letter_code | character, when set to "three_letters", three letter amino acid code is used. One letter by default. |
| annotation_names | character, default NULL. Alternative naming for annotation. |
| start_codons | character vector, default "ATG" |
| stop_codons | character vector, default c("TAA", "TAG", "TGA") |
| custom_motif | character vector, default NULL. |
| BPPARAM | how many cores/threads to use? default: BiocParallel::SerialParam(). To see number of threads used for multicores, do BiocParallel::bpparam()\$workers. You can also add a time remaining bar, for a more detailed pipeline. |

Value

the plot object

Examples

```
library(RiboCrypt)
df <- ORFik.template.experiment()[9:10,]
cds <- loadRegion(df, "cds")
mrna <- loadRegion(df, "mrna")
# multiOmicsPlot_animate(mrna[1], annotation = cds[1], reference_sequence = findFa(df),
#                         frames_type = "columns", leader_extension = 30, trailer_extension = 30,
#                         reads = outputLibs(df, type = "pshifted", output.mode = "envirlist",
#                         naming = "full", BPPARAM = BiocParallel::SerialParam()))
```

multiOmicsPlot_list *Multi-omics plot using list input*

Description

Customizable html plots for visualizing genomic data.

Usage

```

multiOmicsPlot_list(
  display_range,
  annotation = display_range,
  reference_sequence,
  reads,
  viewMode = c("tx", "genomic")[1],
  custom_regions = NULL,
  leader_extension = 0,
  trailer_extension = 0,
  withFrames = NULL,
  frames_type = "lines",
  colors = NULL,
  kmers = NULL,
  kmers_type = c("mean", "sum")[1],
  ylabels = NULL,
  lib_to_annotation_proportions = c(0.8, 0.2),
  lib_proportions = NULL,
  annotation_proportions = NULL,
  width = NULL,
  height = NULL,
  plot_name = "default",
  plot_title = NULL,
  display_sequence = c("both", "nt", "aa", "none")[1],
  seq_render_dist = 100,
  aa_letter_code = c("one_letter", "three_letters")[1],
  annotation_names = NULL,
  start_codons = "ATG",
  stop_codons = c("TAA", "TAG", "TGA"),
  custom_motif = NULL,
  AA_code = Biostrings::GENETIC_CODE,
  BPPARAM = BiocParallel::SerialParam(),
  summary_track = FALSE,
  summary_track_type = frames_type,
  export.format = "svg"
)

```

Arguments

| | |
|---------------------------------|---|
| <code>display_range</code> | the whole region to visualize, a GRangesList or GRanges object |
| <code>annotation</code> | the whole annotation which your target region is a subset, a GRangesList or GRanges object |
| <code>reference_sequence</code> | the genome reference, a FaFile or FaFile convertible object |
| <code>reads</code> | the NGS libraries, as a list of GRanges with or without score column for replicates. |
| <code>viewMode</code> | character, default "tx" (transcript coordinates, first position is 1, exons are merged into a single sequence) Alternative: "genomic" (genomic coordinates, first position is first position in <code>display_range</code> argument. Introns are displayed). |

| | |
|-------------------------------|--|
| custom_regions | a GRangesList or NULL, default: NULL. The alternative annotation, like self defined uORFs etc. The vertical annotation bars will have a different color. |
| leader_extension | integer, default 0. (How much to extend view upstream) |
| trailer_extension | integer, default 0. (How much to extend view downstream) |
| withFrames | a logical vector, default NULL. Alternative: a length 1 or same length as list length of "reads" argument. |
| frames_type | character, default "lines". Alternative: - columns - stacks - area |
| colors | character, default NULL (automatic colouring). If "withFrames" argument is TRUE, colors are set to c("red", "green", "blue") for the 3 frames. Alternative: Character vector of length 1 or length of "reads" list argument. |
| kmers | numeric (integer), bin positions into kmers. |
| kmers_type | character, function used for kmers sliding window. default: "mean", alternative: "sum" |
| ylabels | character, default NULL. Name of libraries in "reads" list argument. |
| lib_to_annotation_proportions | numeric vector of length 2. relative sizes of profiles and annotation. |
| lib_proportions | numeric vector of length equal to displayed libs. Relative sizes of profiles displayed |
| annotation_proportions | numeric vector of length 3 (seq displayed), or 2 (seq not displayed). Relative sizes of annotation tracks. |
| width | numeric, default NULL. Width of plot. |
| height | numeric, default NULL. Height of plot. |
| plot_name | = character, default "default" (will create name from display_range name). Alternative: custom name for region. |
| plot_title | character, default NULL. A title for plot. |
| display_sequence | character/logical, default c("both", "nt", "aa", "none")[1]. If TRUE or "both", display nucleotide and aa sequence in plot. |
| seq_render_dist | integer, default 100. The sequences will appear after zooming below this threshold. |
| aa_letter_code | character, when set to "three_letters", three letter amino acid code is used. One letter by default. |
| annotation_names | character, default NULL. Alternative naming for annotation. |
| start_codons | character vector, default "ATG" |
| stop_codons | character vector, default c("TAA", "TAG", "TGA") |
| custom_motif | character vector, default NULL. |

| | |
|--------------------|--|
| AA_code | Genetic code for amino acid display. Default is SGC0 (standard: Vertebrate). See Biostrings::GENETIC_CODE_TABLE for options. To change to bacterial, do: Biostrings::getGeneticCode("11") |
| BPPARAM | how many cores/threads to use? default: BiocParallel::SerialParam(). To see number of threads used for multicores, do BiocParallel::bpparam()\$workers. You can also add a time remaining bar, for a more detailed pipeline. |
| summary_track | logical, default FALSE. Display a top track, that is the sum of all tracks. |
| summary_track_type | character, default is same as 'frames_type' argument |
| export.format | character, default: "svg". alternative: "png". when you click the top right image button export, what should it export as? |

Value

the plot object

Examples

```
library(RiboCrypt)
df <- ORFik.template.experiment()[9:10,]
cds <- loadRegion(df, "cds")
mrna <- loadRegion(df, "mrna")
multiOmicPlot_list(mrna[1], annotation = cds[1], reference_sequence = findFa(df),
                  frames_type = "columns", leader_extension = 30, trailer_extension = 30,
                  reads = outputLibs(df, type = "pshifted", output.mode = "envirlist",
                  naming = "full", BPPARAM = BiocParallel::SerialParam()))
```

multiOmicPlot_ORFikExp

Multi-omics plot using ORFik experiment input

Description

Customizable html plots for visualizing genomic data.

Usage

```
multiOmicPlot_ORFikExp(
  display_range,
  df,
  annotation = "cds",
  reference_sequence = findFa(df),
  reads = outputLibs(df, type = "pshifted", output.mode = "envirlist", naming = "full",
  BPPARAM = BiocParallel::SerialParam()),
  viewMode = c("tx", "genomic")[1],
  custom_regions = NULL,
  leader_extension = 0,
  trailer_extension = 0,
  withFrames = libraryTypes(df, uniqueTypes = FALSE) %in% c("RFP", "RPF", "LSU"),
  frames_type = "lines",
  colors = NULL,
```

```

kmers = NULL,
kmers_type = c("mean", "sum")[1],
ylabels = bamVarName(df),
lib_to_annotation_proportions = c(0.8, 0.2),
lib_proportions = NULL,
annotation_proportions = NULL,
width = NULL,
height = NULL,
plot_name = "default",
plot_title = NULL,
display_sequence = c("both", "nt", "aa", "none")[1],
seq_render_dist = 100,
aa_letter_code = c("one_letter", "three_letters")[1],
annotation_names = NULL,
start_codons = "ATG",
stop_codons = c("TAA", "TAG", "TGA"),
custom_motif = NULL,
BPPARAM = BiocParallel::SerialParam(),
input_id = "",
summary_track = FALSE,
summary_track_type = frames_type,
export.format = "svg"
)

```

Arguments

display_range the whole region to visualize, a [GRangesList](#) or [GRanges](#) object

df an ORFik [experiment](#) or a list containing ORFik experiments. Usually a list when you have split Ribo-seq and RNA-seq etc.

annotation the whole annotation which your target region is a subset, a [GRangesList](#) or [GRanges](#) object

reference_sequence the genome reference, default `ORFik::findFa(df)`

reads the NGS libraries, as a list of [GRanges](#) with or without 'score' column for replicates. Can also be a `covRle` object of precomputed coverage. Default: `outputLibs(df, type = "pshifted", output.mode = "envirlist", naming = "full", BPPARAM = BiocParallel::SerialParam())`

viewMode character, default "tx" (transcript coordinates, first position is 1, exons are merged into a single sequence)
Alternative: "genomic" (genomic coordinates, first position is first position in `display_range` argument. Introns are displayed).

custom_regions a [GRangesList](#) or `NULL`, default: `NULL`. The alternative annotation, like self defined uORFs etc. The vertical annotation bars will have a different color.

leader_extension integer, default 0. (How much to extend view upstream)

trailer_extension integer, default 0. (How much to extend view downstream)

withFrames a logical vector, default `libraryTypes(df, uniqueTypes = FALSE) %in% c("RFP", "RPF", "LSU")` Alternative: a length 1 or same length as list length of "reads" argument.

| | |
|-------------------------------|--|
| frames_type | character, default "lines". Alternative: - columns - stacks - area |
| colors | character, default NULL (automatic colouring). If "withFrames" argument is TRUE, colors are set to c("red", "green", "blue") for the 3 frames. Alternative: Character vector of length 1 or length of "reads" list argument. |
| kmers | numeric (integer), bin positions into kmers. |
| kmers_type | character, function used for kmers sliding window. default: "mean", alternative: "sum" |
| ylabels | character, default bamVarName(df). Name of libraries in "reads" list argument. |
| lib_to_annotation_proportions | numeric vector of length 2. relative sizes of profiles and annotation. |
| lib_proportions | numeric vector of length equal to displayed libs. Relative sizes of profiles displayed |
| annotation_proportions | numeric vector of length 3 (seq displayed), or 2 (seq not displayed). Relative sizes of annotation tracks. |
| width | numeric, default NULL. Width of plot. |
| height | numeric, default NULL. Height of plot. |
| plot_name | character, default "default" (will create name from display_range name). |
| plot_title | character, default NULL. A title for plot. |
| display_sequence | character/logical, default c("both", "nt", "aa", "none")[1]. If TRUE or "both", display nucleotide and aa sequence in plot. |
| seq_render_dist | integer, default 100. The sequences will appear after zooming below this threshold. |
| aa_letter_code | character, when set to "three_letters", three letter amino acid code is used. One letter by default. |
| annotation_names | character, default NULL. Alternative naming for annotation. |
| start_codons | character vector, default "ATG" |
| stop_codons | character vector, default c("TAA", "TAG", "TGA") |
| custom_motif | character vector, default NULL. |
| BPPARAM | how many cores/threads to use? default: BiocParallel::SerialParam(). To see number of threads used for multicores, do BiocParallel::bpparam()\$workers. You can also add a time remaining bar, for a more detailed pipeline. |
| input_id | character path, default: "", id for shiny to display structures, should be "" for local users. |
| summary_track | logical, default FALSE. Display a top track, that is the sum of all tracks. |
| summary_track_type | character, default is same as 'frames_type' argument |
| export.format | character, default: "svg". alternative: "png". when you click the top right image button export, what should it export as? |

Value

the plot object

Examples

```
library(RiboCrypt)
df <- ORFik.template.experiment()[9,] #Use third library in experiment only
cds <- loadRegion(df, "cds")
multiOmicsPlot_ORFikExp(extendLeaders(extendTrailers(cds[1], 30), 30), df = df,
                        frames_type = "columns")
```

organism_input_select *Select box for organism*

Description

Select box for organism

Usage

```
organism_input_select(genomes, ns)
```

Arguments

| | |
|---------|---|
| genomes | name of genomes, returned from list.experiments() |
| ns | the ID, for shiny session |

Value

selectizeInput object

RiboCrypt_app *Create RiboCrypt app*

Description

Create RiboCrypt app

Usage

```
RiboCrypt_app(
  validate.experiments = TRUE,
  options = list(launch.browser = ifelse(interactive(), TRUE, FALSE)),
  all_exp = list.experiments(validate = validate.experiments),
  browser_options = c(),
  init_tab_focus = "browser"
)
```

Arguments

`validate.experiments` logical, default TRUE, set to FALSE to allow starting the app with malformed experiments, be careful will crash if you try to load that experiment!

`options` list of arguments, default `list("launch.browser" = ifelse(interactive(), TRUE, FALSE))`

`all_exp` a data.table, default: `list.experiments(validate = validate.experiments)`. Which experiments do you want to allow your app to see, default is all in your system config path.

`browser_options` named character vector of browser specific arguments:

- `default_experiment` : Which experiment to select, default: first one
- `default_gene` : Which genes to select, default: first one
- `default_libs` : Which libraries to select: first one, else a single string, where libs are separated by "|", like "RFP_WT_r1|RFP_WT_r2".
- `default_kmer` : K-mer windowing size, default: 1
- `default_frame_type` : Ribo-seq line type, default: "lines"
- `plot_on_start` : Plot when starting, default: "FALSE"

`init_tab_focus` character, default "browser". Which tab to open on init.

Value

RiboCrypt shiny app

Examples

```
## Default run
# RiboCrypt_app()
## Plot on start
# RiboCrypt_app(browser_options = c(plot_on_start = "TRUE"))
## Init with an experiment and gene (you must of course have the experiment)

#RiboCrypt_app(validate.experiments = FALSE,
#               browser_options = c(plot_on_start = "TRUE",
#                                   default_experiment = "human_all_merged_150",
#                                   default_gene = "ATF4-ENSG00000128272"))
```

trimOverlaps

Trim overlaps

Description

Trim overlaps

Usage

```
trimOverlaps(overlaps, display_range)
```

trimOverlaps

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Arguments

overlaps GRanges

display_range GRanges

Value

GRanges

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