Package ‘chimeraviz’

May 29, 2024

Type Package
Title Visualization tools for gene fusions
Version 1.30.0
Description chimeraviz manages data from fusion gene finders and provides useful visualization tools.
License Artistic-2.0
LazyData TRUE
Imports methods, grid, Rsamtools, GenomeInfoDb, GenomicAlignments, RColorBrewer, graphics, AnnotationDbi, RCircos, org.Hs.eg.db, org.Mm.eg.db, rmarkdown, graph, Rgraphviz, DT, plyr, dplyr, BiocStyle, checkmate, gtools, magick
Depends Biostrings, GenomicRanges, IRanges, Gviz, S4Vectors, ensembldb, AnnotationFilter, data.table
Suggests testthat, roxygen2, devtools, knitr, lintr
SystemRequirements bowtie, samtools, and egrep are required for some functionalities
RoxygenNote 7.1.1
VignetteBuilder knitr
biocViews Infrastructure, Alignment
Encoding UTF-8
URL https://github.com/stianlagstad/chimeraviz

BugReports https://github.com/stianlagstad/chimeraviz/issues
git_url https://git.bioconductor.org/packages/chimeraviz
git_branch RELEASE_3_19
git_last_commit 830e53e
git_last_commit_date 2024-04-30
Repository Bioconductor 3.19
Date/Publication 2024-05-29
add_fusion_reads_alignment

Add fusion reads alignment to fusion object

Description

This function lets you add a fusion read alignment file to a fusion object. If you’ve mapped the reads supporting a fusion against the fusion junction sequence, and have the resulting bamfile, use this function to add the information (as a Gviz::GAlignmentPairs object) to the fusion object.

Usage

add_fusion_reads_alignment(fusion, bamfile)
chimeraviz-internals-fusions_to_gene_label_data

Arguments

- fusion: The fusion object to add a genomic alignment to.
- bamfile: The bam file containing the fusion reads plotted to the fusion sequence.

Value

An updated fusion object with fusion@fusion_reads_alignment set.

Examples

```r
# Load data
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
# Find the specific fusion we have aligned reads for
fusion <- get_fusion_by_id(fusions, 5267)
# Get reference to the bamfile with the alignment data
bamfile5267 <- system.file(
  "extdata",
  "5267readsAligned.bam",
  package="chimeraviz")
# Add the bam file of aligned fusion reads to the fusion object
fusion <- add_fusion_reads_alignment(fusion, bamfile5267)
```

chimeraviz

chimeraviz: A package for working with and visualizing fusion genes.

Description

chimeraviz manages data from fusion gene finders and provides useful visualization tools.

chimeraviz-internals-fusions_to_gene_label_data

Create gene label data for RCircos from the given fusions.

Description

This function takes a list of Fusion objects and creates a data frame in the format that RCircos.Gene.Name.Plot() expects for gene label data.

Usage

`.fusions_to_gene_label_data(fusion_list)`
Arguments

fusion_list A list of Fusion objects.

Value

A data frame with fusion gene label data compatible with RCircos::Gene.Name.Plot()

# @examples # Apparently examples shouldn't be set on private functions
defuse833ke <- system.file("extdata", "defuse_833ke_results.filtered.tsv", package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 3)
labelData <- chimeraviz::fusions_to_gene_label_data(fusions) # This labelData can be used with RCircos::Gene.Connector.Plot() and RCircos::Gene.Name.Plot()

Description

This function takes a list of Fusion objects and creates a data frame in the format that RCircos::RCircos.Link.Plot() expects for link data.

Usage

.fusions_to_link_data(fusion_list, min_link_width = 1, max_link_width = 10)

Arguments

fusion_list A list of Fusion objects.
min_link_width The minimum link line width. Default = 1
max_link_width The maximum link line width. Default = 10

Value

A data frame with fusion link data compatible with RCircos::RCircos.Link.Plot()

# @examples # Apparently examples shouldn't be set on private functions
defuse833ke <- system.file("extdata", "defuse_833ke_results.filtered.tsv", package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 3)
linkData <- chimeraviz::fusions_to_link_data(fusions) # This linkData can be used with RCircos::RCircos.Link.Plot()
chimeraviz-internals-scaleListToInterval

Scale a vector of numeric values to an interval.

Description

This function takes a vector of numeric values as well as an interval \([\text{new}_{\text{min}}, \text{new}_{\text{max}}]\) that the numeric values will be scaled (normalized) to.

Usage

```
.scale_list_to_interval(the_list, new_min, new_max)
```

Arguments

- `the_list` A vector of numeric values.
- `new_min` Minimum value for the new interval.
- `new_max` Maximum value for the new interval.

Value

A data frame with fusion link data compatible with `RCircos::RCircos.Link.Plot()`

```
# @examples # Apparently examples shouldn't be set on private functions
list012 <- c(0,1,2)
.scale_list_to_interval(list012, 1, 3) # [1] 1 2 3
```

create_fusion_report

Create a Fusion Report

Description

This function will create a html report with an overplot and a sortable, searchable table with the fusion data.

Usage

```
create_fusion_report(fusions, output_filename, quiet = TRUE)
```

Arguments

- `fusions` A list of Fusion objects.
- `output_filename` Output html-file filename.
- `quiet` Parameter passed to `rmarkdown::render()` to toggle its output.
decide_transcript_category

Value

Creates a html report with an overplot and a sortable, searchable table with the fusion data.

Examples

# Load data
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz"
)fusions <- import_defuse(defuse833ke, "hg19", 3)
# Temporary file to store the report
random_filename <- paste0(
  paste0(sample(LETTERS, 5, replace = TRUE), collapse=''),
  ".png"
)
# Create report
create_fusion_report(fusions, random_filename)
# Delete the file
file.remove(random_filename)

decide_transcript_category

   Retrieves transcripts for partner genes in a Fusion object using EnsemblDb

Description

This function will check where in the transcript (the GRanges object) the fusion breakpoint is located, and return either "exonBoundary", "withinExon", "withinIntron", or "intergenic".

Usage

decide_transcript_category(gr, fusion)

Arguments

  gr          The GRanges object containing the transcript to be checked.
  fusion      The fusion object used to check the transcript.

Value

Either "exonBoundary", "withinExon", "withinIntron", or "intergenic" depending on where in the transcript the breakpoint hits.
Examples

# Load fusion data and choose a fusion object:
defuseData <- system.file(
    "extdata",
    "defuse_833ke_results.filtered.tsv",
    package="chimeraviz"
)fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Create edb object
edbSqliteFile <- system.file(
    "extdata",
    "Homo_sapiens.GRCh37.74.sqlite",
    package="chimeraviz"
)edb <- ensembldb::EnsDb(edbSqliteFile)
# Get all exons for all transcripts in the genes in the fusion transcript
allTranscripts <- ensembldb::exonsBy(edb,
    filter = list(
        AnnotationFilter::GeneIdFilter(
            c(partner_gene_ensembl_id(upstream_partner_gene(fusion)),
              partner_gene_ensembl_id(downstream_partner_gene(fusion))))),
    columns = c(  
        "gene_id",
        "gene_name",
        "tx_id",
        "tx_cds_seq_start",
        "tx_cds_seq_end",
        "exon_id"))
# Extract one of the GRanges objects
gr <- allTranscripts[[1]]
# Check where in the transcript the fusion breakpoint hits
decide_transcript_category(gr, fusion)
# "exonBoundary"
# Check another case
gr <- allTranscripts[[3]]
decide_transcript_category(gr, fusion)
# "withinIntron"

downstream_partner_gene

Get the downstream fusion partner gene

Description

This getter retrieves the downstream PartnerGene object.

This sets the downstream PartnerGene object of a Fusion object.
downstream_partner_gene

Usage

downstream_partner_gene(x)

## S4 method for signature 'Fusion'
downstream_partner_gene(x)

downstream_partner_gene(object) <- value

## S4 replacement method for signature 'Fusion'
downstream_partner_gene(object) <- value

Arguments

x The Fusion object you wish to retrieve the downstream PartnerGene object for.

object The Fusion object you wish to set a new downstream PartnerGene object for.

value The new PartnerGene object.

Value

The downstream PartnerGene object.

Examples

# Load data
defuseData <- system.file("extdata", 
  "defuse_833ke_results.filtered.tsv", 
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Get the downstream fusion partner gene
downstream_partner_gene(fusion)

# Set the downstream PartnerGene object to be the same as the upstream PartnerGene object
downstream_partner_gene(fusion) <- upstream_partner_gene(fusion)
### down_shift

*Remove introns and shift exons leftward*

**Description**
This function takes a GRanges object and moves each IRanges object within next to each other starting at 1. This effectively removes the introns from the GRanges object.

**Usage**
```
down_shift(transcript)
```

**Arguments**
- **transcript**
  The GRanges object to remove introns from.

**Value**
A GRanges object with introns removed.

**Examples**
```
# Create a simple GRanges object:
gr <- IRanges::IRanges(
  start = c(13, 40, 100),
  end = c(20, 53, 110))
# Downshift it and see the introns are removed:
down_shift(gr)
```

---

### fetch_reads_from_fastq

*Fetch reads from fastq files*

**Description**
This function will fetch read sequences from fastq files and put them into new fastq files.

**Usage**
```
fetch_reads_from_fastq(
  reads,
  fastq_file_in1,
  fastq_file_in2,
  fastq_file_out1,
  fastq_file_out2
)
```
Fusion-class

Arguments

- **reads**: List of read IDs that is to be fetched.
- **fastq_file_in1**: First fastq file to search in.
- **fastq_file_in2**: Second fastq file to search in.
- **fastq_file_out1**: First fastq file with results.
- **fastq_file_out2**: Second fastq file with results.

Details

Note: This function runs (read only) bash commands on your system. Therefore the function will only work on a unix system.

Value

The files `fastqFileOut1` and `fastqFileOut2` populated with the specified reads.

Examples

```r
## Not run:
# fastq files that has the supporting reads
fastq1 <- system.file("extdata", "reads.1.fq", package="chimeraviz")
fastq2 <- system.file("extdata", "reads.2.fq", package="chimeraviz")
# Which read ids to extract
reads <- c("13422259", "19375605", "29755061",
           "31632876", "32141428", "33857245")
# Extract the actual reads and put them in the tmp files "fastqFileOut1" and
# "fastqFileOut2"
fastqFileOut1 <- tempfile(pattern = "fq1", tmpdir = tempdir())
fastqFileOut2 <- tempfile(pattern = "fq2", tmpdir = tempdir())
fetch_reads_from_fastq(reads, fastq1, fastq2,
                        fastqFileOut1, fastqFileOut2)
# We now have the reads supporting fusion 5267 in the two files.

## End(Not run)
```

Fusion-class

An S4 class to represent a fusion event.

Description

The Fusion class represents a fusion event, holding data imported from a fusion tool.
Slots

- **id** A unique id representing a fusion event. For deFuse data this will be the cluster id.
- **fusion_tool** Name of the fusion tool.
- **genome_version** Name of the genome used to map reads.
- **spanning_reads_count** The number of spanning reads supporting the fusion.
- **split_reads_count** The number of split reads supporting the fusion.
- **fusion_reads_alignment** A Gviz::AlignmentsTrack object holding the fusion reads aligned to the fusion sequence.
- **gene_upstream** A PartnerGene object holding information of the upstream fusion partner gene.
- **gene_downstream** A PartnerGene object holding information of the downstream fusion partner gene.
- **inframe** A logical value indicating whether or not the downstream fusion partner gene is inframe or not. Not all fusion-finders report this.
- **fusion_tool_specific_data** A list that will hold fields of importance for a specific fusion finder. This field is used because many fusion-finders report important values that are hard to fit into a standardized format. Examples of values that are added to this list is probability from deFuse and EricScore from EricScript.

---

**fusion_spanning_reads_count**

*Get the spanning reads count from a Fusion object*

---

**Description**

This getter retrieves the spanning reads count from a Fusion object.

**Usage**

```r
fusion_spanning_reads_count(x)
```

## S4 method for signature 'Fusion'
fusion_spanning_reads_count(x)

**Arguments**

- **x** The Fusion object you wish to retrieve the spanning reads count for.

**Value**

The Fusion spanning reads count.
fusion_split_reads_count

Examples

# Load data
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Get the spanning reads count
fusion_spanning_reads_count(fusion)

fusion_split_reads_count

*Get the split reads count from a Fusion object*

Description

This getter retrieves the split reads count from a Fusion object

Usage

fusion_split_reads_count(x)

## S4 method for signature 'Fusion'
fusion_split_reads_count(x)

Arguments

x

The Fusion object you wish to retrieve the split reads count for.

Value

The Fusion split reads count.

Examples

# Load data
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Get the split reads count
fusion_split_reads_count(fusion)
fusion_to_data_frame  

**Coerce Fusion object to data.frame**

**Description**

This function is used in create_fusion_report() to convert Fusion objects to a data.frame-format.

**Usage**

fusion_to_data_frame(fusion)

**Arguments**

- fusion
  - The Fusion object to coerce.

**Value**

A data.frame with the fusion object.

**See Also**

create_fusion_report

**Examples**

```r
# Load data
defuse833ke <- system.file(  
  "extdata",  
  "defuse_833ke_results.filtered.tsv",  
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
# Find the fusion object to create a data frame from
fusion <- get_fusion_by_id(fusions, 5267)
# Create the data frame
dffusion <- fusion_to_data_frame(fusion)
```

get_ensembl_ids  

**Get ensembl ids for a fusion object**

**Description**

This function will get the ensembl ids from the org.Hs.eg.db/org.Mm.eg.db package given the gene names of the fusion event.

**Usage**

get_ensembl_ids(fusion)
### get_fusion_by_chromosome

**Find fusions that involves genes in the given chromosome.**

**Description**

Helper function to retrieve the Fusion objects that involves genes in the given chromosome name.

**Usage**

```r
get_fusion_by_chromosome(fusion_list, chr)
```

**Arguments**

- `fusion_list` A list of Fusion objects.
- `chr` The chromosome name we're looking for fusions in.

**Examples**

```r
# Import the filtered defuse results
defuse833keFiltered <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz"
)fusions <- import_defuse(defuse833keFiltered, "hg19", 1)
# Get a specific fusion
fusion <- get_fusion_by_id(fusions, 5267)
# See the ensembl ids:
partner_gene_ensembl_id(upstream_partner_gene(fusion))
# [1] "ENSG00000180198"
partner_gene_ensembl_id(downstream_partner_gene(fusion))
# [1] "ENSG00000162639"
# Reset the fusion objects ensembl ids
partner_gene_ensembl_id(upstream_partner_gene(fusion)) <- ""
partner_gene_ensembl_id(downstream_partner_gene(fusion)) <- ""
# Get the ensembl ids
fusion <- get_ensembl_ids(fusion)
# See that we now have the same ensembl ids again:
partner_gene_ensembl_id(upstream_partner_gene(fusion))
# [1] "ENSG00000180198"
partner_gene_ensembl_id(downstream_partner_gene(fusion))
# [1] "ENSG00000162639"
```
Value

A list of Fusion objects.

Examples

defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
length(get_fusion_by_chromosome(fusions, "chr1"))
# [1] 1

get_fusion_by_gene_name

Find fusions that includes the given gene.

Description

Helper function to retrieve the Fusion objects that has geneName as one of the partner genes.

Usage

get_fusion_by_gene_name(fusion_list, gene_name)

Arguments

fusion_list A list of Fusion objects.
gene_name The gene name we’re looking for.

Details

Note: get_fusion_by_gene_name(fusionList, "MT") will match both MT-ND5 and MT-ND4.

Value

A list of Fusion objects.

Examples

defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
length(get_fusion_by_gene_name(fusions, "RCC1"))
# [1] 1
**get_fusion_by_id**

*Find a specific fusion object in a list by id*

---

**Description**

Helper function to retrieve the Fusion object with the given id.

**Usage**

```r
get_fusion_by_id(fusion_list, id)
```

**Arguments**

- `fusion_list`: A list of Fusion objects.
- `id`: The id (e.g. the cluster_id from a deFuse run) we’re looking for.

**Value**

A Fusion object.

**Examples**

```r
defuse833ke <- system.file(  
  "extdata",  
  "defuse_833ke_results.filtered.tsv",  
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# This should be the Fusion object:
fusion
# [1] "Fusion object"
# [1] "id: 5267"
# [1] "Fusion tool: defuse"
# [1] "Genome version: hg19"
# [1] "Gene names: RCC1-HENMT1"
# [1] "Chromosomes: chr1-chr1"
# [1] "Strands: +,-"
```

---

**get_transcripts_ensembl_db**

*Retrieves transcripts for partner genes in a Fusion object using EnsemblDb*
import_aeron

Description

This function will retrieve transcripts for both genes in a fusion. It will check all transcripts and
decide for each transcript if the fusion breakpoint happens at 1) an exon boundary, 2) within an
exon, or 3) within an intron. This is done because fusions happening at exon boundaries are more
likely to produce biologically interesting gene products. The function returns an updated Fusion
object, where the fusion@gene_upstream@transcriptsX slots are set with transcript information.

Usage

get_transcripts_ensembl_db(fusion, edb)

Arguments

fusion The fusion object to find transcripts for.
edb The edb object used to fetch data from.

Value

An updated fusion object with transcript data stored.

Examples

# Load fusion data and choose a fusion object:
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Create edb object
edbSqliteFile <- system.file(
  "extdata",
  "Homo_sapiens.GRCh37.74.sqlite",
  package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)
# Add transcripts data to fusion object
fusion <- get_transcripts_ensembl_db(fusion, edb)
# The transcripts are now accessible through fusion@gene_upstream@transcripts and
# fusion@gene_downstream@transcripts .

import_aeron

Import results from an Aeron run into a list of Fusion objects.

Description

A function that imports the results from an Aeron run into a list of Fusion objects.
Usage

import_aeron(
    filename_fusion_support,
    filename_fusion_transcript,
    genome_version,
    limit
)

Arguments

filename_fusion_support
    Filename for the Aeron result file fusion_support..txt.

filename_fusion_transcript
    Filename for the Aeron result file fusion_transcripts..txt.

genome_version
    Which genome was used in mapping (hg19, hg38, etc.).

limit
    A limit on how many lines to read.

Details

Note that the strands and breakpoint positions are not included in the result files from Aeron. These have to be retrieved manually, using the ensembl identifiers (which are included in the result files, and will be available in the Fusion objects after importing).

Value

A list of Fusion objects.

Examples

aeronfusionsupportfile <- system.file("extdata", "aeron_fusion_support.txt", package="chimeraviz")
aeronfusiontranscriptfile <- system.file("extdata", "aeron_fusion_transcripts.fa", package="chimeraviz")
fusions <- import_aeron(    aeronfusionsupportfile,    aeronfusiontranscriptfile,    "hg19",    3)
# This should import a list of 3 fusions described in Fusion objects.
import_chimpipe

Import results from a ChimPipe run into a list of Fusion objects.

Description
A function that imports the results from a ChimPipe run, typically from a chimericJunctions_.txt file, into a list of Fusion objects.

Usage
import_chimpipe(filename, genome_version, limit)

Arguments
- filename: Filename for the ChimPipe results.
- genome_version: Which genome was used in mapping (hg19, hg38, etc.).
- limit: A limit on how many lines to read.

Value
A list of Fusion objects.

Examples
chimpipefile <- system.file(
  "extdata",
  "chimericJunctions_MCF-7.txt",
  package="chimeraviz"
)fusions <- import_chimpipe(chimpipefile, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.

import_defuse

Import results from a deFuse run into a list of Fusion objects.

Description
A function that imports the results from a deFuse run, typically from a results.filtered.tsv file, into a list of Fusion objects.

Usage
import_defuse(filename, genome_version, limit)
import_ericscript

Arguments

- `filename` Filename for the deFuse results .tsv file.
- `genome_version` Which genome was used in mapping (hg19, hg38, etc.).
- `limit` A limit on how many lines to read.

Value

A list of Fusion objects.

Examples

defuse833ke <- system.file(  
"extdata",  
"defuse_833ke_results.filtered.tsv",  
package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 3)  
# This should import a list of 3 fusions described in Fusion objects.

import_ericscript

Import results from a EricScript run into a list of Fusion objects.

Description

A function that imports the results from a EricScript run into a list of Fusion objects.

Usage

import_ericscript(filename, genome_version, limit)

Arguments

- `filename` Filename for the EricScript results file.
- `genome_version` Which genome was used in mapping (hg19, hg38, etc.).
- `limit` A limit on how many lines to read.

Value

A list of Fusion objects.

Examples

ericscriptData <- system.file(  
"extdata",  
"ericScript_SRR1657556.results.total.tsv",  
package = "chimeraviz")
fusions <- import_ericscript(ericscriptData, "hg19", 3)  
# This should import a list of 3 fusions described in Fusion objects.
import_function_non_ucsc

*Alternative import function for Gviz::AlignmentsTrack*

**Description**

This alternative import function for use with Gviz::AlignmentsTrack imports a bamfile with non-UCSC chromosome names.

**Usage**

`import_function_non_ucsc(file, selection)`

**Arguments**

- **file**
  - The bamfile.

- **selection**
  - Which regions to get from the bamfile.

**Value**

A GRanges object with coverage data for the selection.

---

import_fusioncatcher

*Import results from a Fusioncatcher run into a list of Fusion objects.*

**Description**

A function that imports the results from a Fusioncatcher run, typically from a final-list-candidate-fusion-genes.txt file, into a list of Fusion objects.

**Usage**

`import_fusioncatcher(filename, genome_version, limit)`

**Arguments**

- **filename**
  - Filename for the Fusioncatcher final-list-candidate-fusion-genes.txt results file.

- **genome_version**
  - Which genome was used in mapping (hg19, hg38, etc.).

- **limit**
  - A limit on how many lines to read.

**Value**

A list of Fusion objects.
import_fusionmap

Examples

```
fusioncatcher833ke <- system.file(
    "extdata",
    "fusioncatcher_833ke_final-list-candidate-fusion-genes.txt",
    package = "chimeraviz")
fusions <- import_fusioncatcher(fusioncatcher833ke, "hg38", 3)
# This should import a list of 3 fusions described in Fusion objects.
```

---

**import_fusionmap**

*Import results from a FusionMap run into a list of Fusion objects.*

**Description**

A function that imports the results from a FusionMap run, typically from a `InputFastq.FusionReport.txt` file, into a list of Fusion objects.

**Usage**

```
import_fusionmap(filename, genome_version, limit)
```

**Arguments**

- `filename` 
  Filname for the FusionMap PairedEndFusionReport.txt results file.
- `genome_version` 
  Which genome was used in mapping (hg19, hg38, etc.).
- `limit` 
  A limit on how many lines to read.

**Value**

A list of Fusion objects.

**Examples**

```
fusionmapData <- system.file( 
    "extdata",
    "FusionMap_01_TestDataset_InputFastq.FusionReport.txt",
    package = "chimeraviz")
fusions <- import_fusionmap(fusionmapData, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.
```
import_infusion

Import results from an InFusion run into a list of Fusion objects.

**Description**

A function that imports the results from an InFusion run into a list of Fusion objects.

**Usage**

    import_infusion(filename, genome_version, limit)

**Arguments**

- **filename**  
  Filename for the jaffa_results.csv file.
- **genome_version**  
  Which genome was used in mapping (hg19, hg38, etc.).
- **limit**  
  A limit on how many lines to read.

**Value**

A list of Fusion objects.

**Examples**

```r
infusionData <- system.file(
  "extdata",
  "infusion_fusions.txt",
  package = "chimeraviz")
fusions <- import_infusion(infusionData, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.
```

import_jaffa

Import results from a JAFFA run into a list of Fusion objects.

**Description**

A function that imports the results from a JAFFA run, typically from a jaffa_results.csv file, into a list of Fusion objects.

**Usage**

    import_jaffa(filename, genome_version, limit)
import_oncofuse

Arguments

filename Filename for the jaffa_results.csv file.

genome_version Which genome was used in mapping (hg19, hg38, etc.).

limit A limit on how many lines to read.

Value

A list of Fusion objects.

Examples

jaffaData <- system.file(
  "extdata",
  "jaffa_results.csv",
  package = "chimeraviz")
fusions <- import_jaffa(jaffaData, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.

import_oncofuse Import results from a oncofuse run into a list of Fusion objects.

Description

A function that imports the results from a oncofuse run, typically from a results.filtered.tsv file, into a list of Fusion objects.

Usage

import_oncofuse(filename, genome_version, limit)

Arguments

filename Filename for the oncofuse results .tsv file.

genome_version Which genome was used in mapping (hg19, hg38, etc.).

limit A limit on how many lines to read.

Details

This import function was contributed by Lavinia G, ref https://github.com/stianlagstad/chimeraviz/issues/47#issuecomment-409773158

Value

A list of Fusion objects.
import_prada

Examples

```r
oncofuse833ke <- system.file(
  "extdata",
  "oncofuse.outfile",
  package="chimeraviz")
fusions <- import_oncofuse(oncofuse833ke, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.
```

---

**import_prada**

Import results from a PRADA run into a list of Fusion objects.

---

**Description**

A function that imports the results from a PRADA run into a list of Fusion objects.

**Usage**

```r
import_prada(filename, genome_version, limit)
```

**Arguments**

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>filename</td>
<td>Filename for the PRADA results file.</td>
</tr>
<tr>
<td>genome_version</td>
<td>Which genome was used in mapping (hg19, hg38, etc.).</td>
</tr>
<tr>
<td>limit</td>
<td>A limit on how many lines to read.</td>
</tr>
</tbody>
</table>

**Value**

A list of Fusion objects.

**Examples**

```r
pradaData <- system.file(
  "extdata",
  "PRADA.acc.fusion.fq.TAF.tsv",
  package = "chimeraviz")
fusions <- import_prada(pradaData, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.
```
import_soapfuse

Import results from a SOAPfuse run into a list of Fusion objects.

Description

A function that imports the results from a SOAPfuse run, typically from a final.Fusion.specific.for.genes file, into a list of Fusion objects.

Usage

import_soapfuse(filename, genome_version, limit)

Arguments

filename Filename for the SOAPfuse final-list-candidate-fusion-genes.txt results file.

genome_version Which genome was used in mapping (hg19, hg38, etc.).

limit A limit on how many lines to read.

Value

A list of Fusion objects.

Examples

```r
soapfuse833ke <- system.file(  
  "extdata",  
  "soapfuse_833ke_final.Fusion.specific.for.genes",  
  package = "chimeraviz")
fusions <- import_soapfuse(soapfuse833ke, "hg19", 3)  
# This should import a list of 3 fusions described in Fusion objects.
```

import_squid

Import results from a SQUID run into a list of Fusion objects.

Description

A function that imports the results from a SQUID run into a list of Fusion objects.

Usage

import_squid(filename, genome_version, limit)
import_starfusion

Arguments

filename Filename for the SQUID results.
genome_version Which genome was used in mapping (hg19, hg38, etc.).
limit A limit on how many lines to read.

Value

A list of Fusion objects.

Examples

squidfile <- system.file(
  "extdata",
  "squid_hcc1954_sv.txt",
  package="chimeraviz")
fusions <- import_squid(squidfile, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.

import_starfusion Import results from a STAR-Fusion run into a list of Fusion objects.

Description

A function that imports the results from a STAR-Fusion run, typically from a star-fusion.fusion_candidates.final.abridged file, into a list of Fusion objects.

Usage

import_starfusion(filename, genome_version, limit)

Arguments

filename Filename for the STAR-Fusion star-fusion.fusion_candidates.final.abridged results file.
genome_version Which genome was used in mapping (hg19, hg38, etc.).
limit A limit on how many lines to read.

Value

A list of Fusion objects.
Examples

```r
starfusionData <- system.file(
  "extdata",
  "star-fusion.fusion_candidates.final.abridged.txt",
  package = "chimeraviz")
fusions <- import_starfusion(starfusionData, "hg19", 3)
# This should import a list of 3 fusions described in Fusion objects.
```

---

**PartnerGene-class**

*An S4 class to represent a gene partner in a fusion*

---

**Description**

The PartnerGene class represents one of the genes in a fusion event.

**Slots**

- `name` Character containing name of the gene.
- `ensembl_id` Character containing ensembl id for the gene.
- `chromosome` Character containing chromosome name.
- `breakpoint` Numeric containing the fusion breakpoint.
- `strand` Character containing gene strand.
- `junction_sequence` Biostrings::DNAString containing the sequence right before/after the fusion breakpoint.
- `transcripts` GenomicRanges::GRangesList containing three GenomicRanges::Granges() objects, one for each "transcript type". The transcript types are: 1) Transcripts where the fusion breakpoint hits an exon boundary, 2) transcripts where the fusion breakpoint is within an exon, 3) transcripts where the fusion breakpoint is within an intron.

---

**partner_gene_ensembl_id**

*Get the Ensembl ID from a PartnerGene object*

---

**Description**

This getter retrieves the Ensembl ID from a PartnerGene object.

This sets the Ensembl ID of a PartnerGene object.
Usage

partner_gene_ensembl_id(x)

## S4 method for signature 'PartnerGene'
partner_gene_ensembl_id(x)

partner_gene_ensembl_id(object) <- value

## S4 replacement method for signature 'PartnerGene'
partner_gene_ensembl_id(object) <- value

Arguments

x The PartnerGene object you wish to retrieve the Ensembl ID for.
object The PartnerGene object you wish to set a new Ensembl ID for.
value The new Ensembl ID.

Value

The upstream fusion partner gene Ensembl ID.

Examples

# Load data
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Get the Ensembl ID from the upstream fusion partner gene
partner_gene_ensembl_id(upstream_partner_gene(fusion))

# Load data
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Set the downstream PartnerGene object to be the same as the upstream PartnerGene object
partner_gene_ensembl_id(upstream_partner_gene(fusion)) <- "test"
partner_gene_junction_sequence

*Get the junction sequence from a PartnerGene object*

**Description**

This getter retrieves the junction sequence from a PartnerGene object.

**Usage**

```r
partner_gene_junction_sequence(x)
```

### S4 method for signature 'PartnerGene'

```r
partner_gene_junction_sequence(x)
```

**Arguments**

- `x` The PartnerGene object you wish to retrieve the junction sequence for.

**Value**

The upstream fusion partner gene junction sequence.

**Examples**

```r
# Load data
defuseData <- system.file("extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Get the junction sequence from the upstream fusion partner gene
partner_gene_junction_sequence(upstream_partner_gene(fusion))
```

---

**plot_circle**

*Create a circle plot of the given fusions.*

**Description**

This function takes a list of Fusion objects and creates a circle plot indicating which chromosomes the fusion genes in the list consists of.

**Usage**

```r
plot_circle(fusion_list)
```
Arguments

- fusion_list: A list of Fusion objects.

Details

Note that only a limited number of gene names can be shown in the circle plot due to the limited resolution of the plot. RCircos will automatically limit the number of gene names shown if there are too many.

Value

Creates a circle plot.

Examples

```r
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz"
)
fusions <- import_defuse(defuse833ke, "hg19", 3)
# Temporary file to store the plot
pngFilename <- tempfile(
  pattern = "circlePlot",
  fileext = ".png",
  tmpdir = tempdir())
# Open device
png(pngFilename, width = 1000, height = 750)
# Plot!
plot_circle(fusions)
# Close device
dev.off()
```

plot_fusion

Plot a fusion event with transcripts, coverage and ideograms.

Description

This function creates a plot with information about transcripts, coverage, location and more.

Usage

```r
plot_fusion(
  fusion,
  edb = NULL,
  bamfile = NULL,
  which_transcripts = "exonBoundary",
  ylim = c(0, 1000),
)```
non_ucsc = TRUE,
reduce_transcripts = FALSE,
bedgraphfile = NULL
)

plot_fusion_separate(
fusion,
edb,
bamfile = NULL,
which_transcripts = "exonBoundary",
ylim = c(0, 1000),
non_ucsc = TRUE,
reduce_transcripts = FALSE,
bedgraphfile = NULL
)

plot_fusion_together(
fusion,
edb,
bamfile = NULL,
which_transcripts = "exonBoundary",
ylim = c(0, 1000),
non_ucsc = TRUE,
reduce_transcripts = FALSE,
bedgraphfile = NULL
)

Arguments

fusion The Fusion object to plot.
edb The ensemblldb object that will be used to fetch data.
bamfile The bamfile with RNA-seq data.
which_transcripts This character vector decides which transcripts are to be plotted. Can be "exonBoundary", "withinExon", "withinIntron", "intergenic", or a character vector with specific transcript ids. Default value is "exonBoundary".
ylim Limits for the coverage y-axis.
non_ucsc Boolean indicating whether or not the bamfile used has UCSC-styled chromosome names (i.e. with the "chr" prefix). Setting this to true lets you use a bamfile with chromosome names like "1" and "X", instead of "chr1" and "chrX".
reduce_transcripts Boolean indicating whether or not to reduce all transcripts into a single transcript for each partner gene.
bedgraphfile A bedGraph file to use instead of the bamfile to plot coverage.
Details

plot_fusion() will dispatch to either plot_fusion_separate() or plot_fusion_together(). plot_fusion_separate() will plot the fusion gene partners in separate graphs shown next to each other, while plot_fusion_together() will plot the fusion gene partners in the same graph with the same x-axis. plot_fusion() will dispatch to plot_fusion_together() if the fusion gene partners are on the same strand, same chromosome and are close together (≤50,000 bp apart).

Value

Creates a fusion plot.

Examples

```r
# Load data and example fusion event
defuse833ke <- system.file("extdata", "defuse_833ke_results.filtered.tsv", package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)

# Load edb
edbSqliteFile <- system.file("extdata", "Homo_sapiens.GRCh37.74.sqlite", package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)

# bamfile with reads in the regions of this fusion event
bamfile5267 <- system.file("extdata", "fusion5267and11759reads.bam", package="chimeraviz")

# Temporary file to store the plot
pngFilename <- tempfile(pattern = "fusionPlot", fileext = ".png", tmpdir = tempdir())

# Open device
png(pngFilename, width = 1000, height = 750)

# Plot!
plot_fusion(
  fusion = fusion,
  bamfile = bamfile5267,
  edb = edb,
  non_ucsc = TRUE)

# Close device
dev.off()

# Example using a .bedGraph file instead of a .bam file:
# Load data and example fusion event
defuse833ke <- system.file("extdata", "defuse_833ke_results.filtered.tsv", package="chimeraviz")
```

plot_fusion_reads

Create a plot of the reads supporting the given fusion.

Description

This function takes a Fusion object and plots the reads supporting the fusion on top of the fusion sequence (fusion@junction_sequence), provided that add_fusion_reads_alignment() has been run earlier in order to add fusion reads alignment data to the fusion object.

Usage

plot_fusion_reads(fusion, show_all_nucleotides = TRUE, nucleotide_amount = 10)

Arguments

fusion The Fusion object to plot.
plot_fusion_reads

show_all_nucleotides

  Boolean indicating whether or not to show all nucleotides. If FALSE, then only nucleotide_amount amount of nucleotides will be shown on each end of the fusion junction. If TRUE, then the whole fusion junction sequence will be shown.

nucleotide_amount

  The number of nucleotides to show on each end of the fusion junction sequence. Defaults to 10. Only applicable if show_all_nucleotides is set to TRUE.

Details

  Note that the package used for plotting, Gviz, is strict on chromosome names. If the plot produced doesn’t show the reads, the problem might be solved by naming the fusion sequence “chrNA”.

Value

  Creates a fusion reads plot.

See Also

  add_fusion_reads_alignment

Examples

  # Load data
defuseData <- system.file(
    "extdata",
    "defuse_833ke_results.filtered.tsv",
    package="chimeraviz"
)fusions <- import_defuse(defuseData, "hg19", 1)
# Find the specific fusion we have aligned reads for
fusion <- get_fusion_by_id(fusions, 5267)
bamfile <- system.file(
    "extdata",
    "5267readsAligned.bam",
    package="chimeraviz"
)# Add the bam file of aligned fusion reads to the fusion object
fusion <- add_fusion_reads_alignment(fusion, bamfile)
# Temporary file to store the plot
pngFilename <- tempfile(
    pattern = "fusionPlot",
    fileext = ".png",
    tmpdir = tempdir())
# Calculate image size based on supporting reads and length of junction
# sequence.
imageWidth <- (nchar(partner_gene_junction_sequence(upstream_partner_gene(fusion))) +
    nchar(partner_gene_junction_sequence(downstream_partner_gene(fusion)))) * 15
imageHeight <- ((fusion_split Reads count(fusion)+fusion_spanning reads count(fusion)) * 20
# Open device
png(pngFilename, width = imageWidth, height = imageHeight)
# Now we can plot
plot_fusion_reads(fusion)
# Close device
Plot possible fusion transcripts based on annotation.

Description
This function takes a fusion object and an ensemblDb object and plots the reduced version of the fusion transcript. This transcript consist of the "mashed together" version of all possible fusion transcripts based on known annotations. If a bamfile is specified, the fusion transcript will be plotted with coverage information.

Usage
plot_fusion_transcript(
  fusion,
  edb = NULL,
  bamfile = NULL,
  which_transcripts = "exonBoundary",
  bedgraphfile = NULL
)

Arguments

  fusion            The Fusion object to plot.
  edb              The edb object that will be used to fetch data.
  bamfile          The bamfile with RNA-seq data.
  which_transcripts This character vector decides which transcripts are to be plotted. Can be "exonBoundary", "withinExon", "withinIntron", "intergenic", or a character vector with specific transcript ids. Default value is "exonBoundary".
  bedgraphfile     A bedGraph file to use instead of the bamfile to plot coverage.

Details
Note that the transcript database used (the edb object) must have the same seqnames as any bamfile used. Otherwise the coverage data will be wrong.

Value
Creates a fusion transcript plot.
Examples

```r
# Load data and example fusion event
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz"
)fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Load edb
edbSqliteFile <- system.file(
  "extdata",
  "Homo_sapiens.GRCh37.74.sqlite",
  package="chimeraviz"
)edb <- ensembldb::EnsDb(edbSqliteFile)
# bamfile with reads in the regions of this fusion event
bamfile5267 <- system.file(
  "extdata",
  "fusion5267and11759reads.bam",
  package="chimeraviz"
)# Temporary file to store the plot
pngFilename <- tempfile(
  pattern = "fusionPlot",
  fileext = ".png",
  tmpdir = tempdir())
# Open device
png(pngFilename, width = 500, height = 500)
# Plot!
plot_fusion_transcript(
  fusion = fusion,
  bamfile = bamfile5267,
  edb = edb)
# Close device
dev.off()

# Example using a .bedGraph file instead of a .bam file:
# Load data and example fusion event
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz"
)fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Load edb
edbSqliteFile <- system.file(
  "extdata",
  "Homo_sapiens.GRCh37.74.sqlite",
  package="chimeraviz"
)edb <- ensembldb::EnsDb(edbSqliteFile)
# bedgraphfile with coverage data from the regions of this fusion event
bedgraphfile <- system.file(
  "extdata",
  "fusion5267and11759reads.bedGraph",
  package="chimeraviz"
)
```
```
package="chimeraviz"
# Temporary file to store the plot
pngFilename <- tempfile(
  pattern = "fusionPlot",
  fileext = ".png",
  tmpdir = tempdir())
# Open device
png(pngFilename, width = 500, height = 500)
# Plot!
plot_fusion_transcripts(  
  fusion = fusion,
  bamfile = bamfile5267,
  edb = edb)
# Close device
dev.off()
```

---

**plot_fusion_transcripts_graph**

*Graph plot of possible fusion transcripts.*

**Description**

This function takes a fusion object and a TranscriptDb object and plots a graph showing the possible fusion transcripts.

**Usage**

```r
plot_fusion_transcripts_graph(
  fusion,
  edb = NULL,
  which_transcripts = "exonBoundary",
  rankdir = "TB"
)
```

**Arguments**

- **fusion**
  - The Fusion object to plot.
- **edb**
  - The edb object that will be used to fetch data.
- **which_transcripts**
  - This character vector decides which transcripts are to be plotted. Can be "exonBoundary", "withinExon", "withinIntron", "intergenic", or a character vector with specific transcript ids. Default value is "exonBoundary".
- **rankdir**
  - Choose whether the graph should be plotted from left to right ("LR"), or from top to bottom ("TB"). This parameter is given to Rgraphviz::plot().

**Value**

Creates a fusion transcripts graph plot.
Examples

```r
# Load data and example fusion event
defuse833ke <- system.file(
    "extdata",
    "defuse_833ke_results.filtered.tsv",
    package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Load edb
edbSqliteFile <- system.file(
    "extdata",
    "Homo_sapiens.GRCh37.74.sqlite",
    package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)
# Temporary file to store the plot
pngFilename <- tempfile(
    pattern = "fusionPlot",
    fileext = ".png",
    tmpdir = tempdir())
# Open device
png(pngFilename, width = 500, height = 500)
# Plot!
plot_fusion_transcripts_graph(
    fusion = fusion,
    edb = edb)
# Close device
dev.off()
```

plot_fusion_transcript_with_protein_domain

\textit{Plot a specific fusion transcript with protein domain annotations}

Description

This function takes a fusion object, an ensembldb object, a bedfile with protein domain data and two specific transcript ids. The function plots the specific fusion transcript along with annotations of protein domains. If a bamfile is specified, the fusion transcript will be plotted with coverage information.

Usage

```r
plot_fusion_transcript_with_protein_domain(
    fusion, 
edb = NULL, 
bamfile = NULL, 
bedfile = NULL, 
gene_upstream_transcript = "", 
gene_downstream_transcript = "",
```
plot_fusion_transcript_with_protein_domain

plot_downstream_protein_domains_if_fusion_is_out_of_frame = FALSE
)

Arguments

fusion        The Fusion object to plot.
edb           The edb object that will be used to fetch data.
bamfile       The bamfile with RNA-seq data.
bedfile       The bedfile with protein domain data.
gene_upstream_transcript
              The transcript id for the upstream gene.
gene_downstream_transcript
              The transcript id for the downstream gene.
plot_downstream_protein_domains_if_fusion_is_out_of_frame
              Setting this to true makes the function plot protein domains in the downstream
gene even though the fusion is out of frame.

Details

Note that the transcript database used (the edb object) must have the same seqnames as any bamfile
used. Otherwise the coverage data will be wrong.

Value

Creates a fusion transcript plot with annotations of protein domains.

Examples

# Load data and example fusion event
defuse833ke <- system.file(
    "extdata",
    "defuse_833ke_results.filtered.tsv",
    package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Select transcripts
gene_upstream_transcript <- "ENST00000434290"
gene_downstream_transcript <- "ENST00000370031"
# Load edb
edbSqliteFile <- system.file(
    "extdata",
    "Homo_sapiens.GRCh37.74.sqlite",
    package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)
# bamfile with reads in the regions of this fusion event
bamfile5267 <- system.file(
    "extdata",
    "Fusion5267and11759reads.bam",
    package="chimeraviz")
# bedfile with protein domains for the transcripts in this example
bedfile <- system.file(
  "extdata",
  "protein_domains_5267.bed",
  package="chimeraviz")
# Temporary file to store the plot
pngFilename <- tempfile(
  pattern = "fusionPlot",
  fileext = ".png",
  tmpdir = tempdir())
# Open device
png(pngFilename, width = 500, height = 500)
# Plot!
plot_fusion_transcript_with_protein_domain(
  fusion = fusion,
  edb = edb,
  bamfile = bamfile5267,
  bedfile = bedfile,
  gene_upstream_transcript = gene_upstream_transcript,
  gene_downstream_transcript = gene_downstream_transcript,
  plot_downstream_protein_domains_if_fusion_is_out_of_frame = TRUE)
# Close device
dev.off()

plot_transcripts

Plot transcripts for each partner gene in a fusion event.

Description

This function takes a fusion object and an ensembldb object and plots transcripts for both genes, showing which parts of each genes are included in the fusion event. If the bamfile parameter is set, then the coverage is plotted beneath the transcripts.

Usage

plot_transcripts(
  fusion,
  edb = NULL,
  bamfile = NULL,
  which_transcripts = "exonBoundary",
  non_ucsc = TRUE,
  ylim = c(0, 1000),
  reduce_transcripts = FALSE,
  bedgraphfile = NULL
)
**Arguments**

- **fusion**
  The Fusion object to plot.
- **edb**
  The edb object that will be used to fetch data.
- **bamfile**
  The bamfile with RNA-seq data.
- **which_transcripts**
  This character vector decides which transcripts are to be plotted. Can be "exonBoundary", "withinExon", "withinIntron", "intergenic", or a character vector with specific transcript ids. Default value is "exonBoundary".
- **non_ucsc**
  Boolean indicating whether or not the bamfile used has UCSC-styled chromosome names (i.e. with the "chr" prefix). Setting this to true lets you use a bamfile with chromosome names like "1" and "X", instead of "chr1" and "chrX".
- **ylim**
  Limits for the coverage y-axis.
- **reduce_transcripts**
  Boolean indicating whether or not to reduce all transcripts into a single transcript for each partner gene.
- **bedgraphfile**
  A bedGraph file to use instead of the bamfile to plot coverage.

**Value**

Creates a fusion transcripts plot.

**Examples**

```r
# Load data and example fusion event
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Load edb
edbSQLiteFile <- system.file(
  "extdata",
  "Homo_sapiens.GRCh37.74.sqlite",
  package="chimeraviz")
edb <- ensembldb::EnsDb(edbSQLiteFile)
# bamfile with reads in the regions of this fusion event
bamfile5267 <- system.file(
  "extdata",
  "fusion5267and11759reads.bam",
  package="chimeraviz")
# Temporary file to store the plot
pngFilename <- tempfile(
  pattern = "fusionPlot",
  fileext = ".png",
  tmpdir = tempdir())
# Open device
png(pngFilename, width = 500, height = 500)
# Plot!
```
plot_transcripts(
    fusion = fusion,
    edb = edb,
    bamfile = bamfile5267,
    non_ucsc = TRUE)
# Close device
dev.off()

# Example using a .bedGraph file instead of a .bam file:
# Load data and example fusion event
defuse833ke <- system.file(
    "extdata",
    "defuse_833ke_results.filtered.tsv",
    package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Load edb
edbSqliteFile <- system.file(
    "extdata",
    "Homo_sapiens.GRCh37.74.sqlite",
    package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)
# bedgraphfile with coverage data from the regions of this fusion event
bedgraphfile <- system.file(
    "extdata",
    "fusion5267and11759reads.bedGraph",
    package="chimeraviz")
# Temporary file to store the plot
pngFilename <- tempfile(
    pattern = "fusionPlot",
    fileext = ".png",
    tmpdir = tempdir())
# Open device
png(pngFilename, width = 500, height = 500)
# Plot!
plot_transcripts(
    fusion = fusion,
    edb = edb,
    bedgraphfile = bedgraphfile,
    non_ucsc = TRUE)
# Close device
dev.off()

---

ChimPipe data

Description

Documentation for the ChimPipe example data.
Documentation for the SQUID example data.
chimericJunctions_MCF-7.txt

This is example data from the ChimPipe tutorial all-in-one package located at https://chimpipe.readthedocs.io/en/latest/tutorial.html#download-all-in-one-package It was downloaded March 10th 2020

squid_hcc1954_sv.txt

This is example data for SQUID provided on GitHub here: https://github.com/Kingsford-Group/squid/issues/20#issuecomment-598888472 It was downloaded March 17th 2020

raw_cytobandhg19 Cytoband information HG19

Description

Cytoband information for the HG19 assembly from UCSC. Downloaded from http://hgdownload.cse.ucsc.edu/goldenpath/hg19/database/UCSC.HG19.Human.CytoBandIdeogram.txt

This data is used with RCircos in plot_circle().

raw_cytobandhg38 Cytoband information HG38

Description

Cytoband information for the HG38 assembly from UCSC. Downloaded from http://hgdownload.cse.ucsc.edu/goldenpath/hg38/database

Details

All _alt or _random entries has been manually removed, as has the chrM entry.

UCSC.HG38.Human.CytoBandIdeogram.txt

This data is used with RCircos in plot_circle().
Description

Documentation for the deFuse example data.

defuse_833ke_results.filtered.tsv

This file has the results from a run of deFuse-0.7.0 on the 833ke cell line. The program was ran with the standard configuration, but with the parameter span_count_threshold=5 instead of the standard 3. The resulting results.filtered.tsv file was then manually filtered to only include 17 fusion events in the interest of saving computing time for tests and examples. The original results contained 171 fusion events.

reads_supporting_defuse_fusion_5267.*.fq

These two files, reads_supporting_defuse_fusion_5267.1.fq and reads_supporting_defuse_fusion_5267.2.fq, contains the reads that support the fusion event with cluster_id 5267.

5267readsAligned.bam

The bamfile 5267readsAligned.bam and the 5267readsAligned.bam.bai index file contains the reads supporting the fusion event with cluster_id 5267 aligned to the fusion sequence. It is used with plot_fusion_reads().

Description

Documentation for the EricScript example data.

ericscript_SRR1657556.results.total.tsv

This is example data thankfully provided by EricScript author Matteo Benelli.
protein_domains_5267.bed

This file is an excerpt from a larger file that we created by: - downloading domain name annotation from Pfam database (PfamA version 31) and domain region annotation from Ensembl database through BioMart API - switching the domain coordinates in the protein level to these in transcript level.

fusion5267and11759reads.bam

This file is the result of running these commands:

```
samtools view -b original_bamfile.bam "1:28831455-28866812" "1:109189912-109205148" "12:8608225-8677832" > fusion5267and11759reads.bam samtools index fusion5267and11759reads.bam fusion5267and11759reads.bam.bai
```

where we extract the reads mapping to the region where we know the fusions with cluster_id=5267 and cluster_id=11759 is.

The original_bamfile.bam is from a study of the 833KE cell line by Andreas M. Hoff et al., documented in the paper [Identification of Novel Fusion Genes in Testicular Germ Cell Tumors](http://cancerres.aacrjournals.org/content/76/1/108.full).
raw_fusion5267readsBedGraph

Fusion5267and11759 bedGraph file

Description

Documentation for the fusion5267and11759reads.bedGraph file containing read count data from the regions of the fusion event with cluster_id=5267.

fusion5267and11759reads.bedGraph

This file is the result of running this command:

```
bedtools genomecov -ibam fusion5267and11759reads.bam -bg > fusion5267and11759reads.bam.bedGraph
```

fusion5267and11759reads.bam has its own documentation entry for how it was created.

raw_fusioncatcher

Fusioncatcher data

Description

Documentation for the Fusioncatcher example data.

fusioncatcher_833ke_final-list-candidate-fusion-genes.txt

This file has the results from a run of Fusioncatcher-0.99.3e on the 833ke cell line. The program was ran with the standard configuration file and with the parameters "-p 8 -z -keep-preliminary".

raw_fusionmap

FusionMap data

Description

Documentation for the FusionMap example data.

FusionMap_01_TestDataset_InputFastq.FusionReport.txt

This is example data provided with the FusionMap version released 2015-03-31.
Homo_sapiens.GRCh37.74_subset.gtf

The Homo_sapiens.GRCh37.74.gtf file is a subset version of the Ensembl Homo_sapiens.GRCh37.74.gtf file, located here: ftp://ftp.ensembl.org/pub/release-74/gtf/homo_sapiens. This gtf file contains transcripts for the partner genes in two of the fusion transcripts from the deFuse example data provided with this package: The fusion transcript with cluster_id=5267, and the fusion transcript with cluster_id=11759.

The file is the result of running this command:

```
# grep "ENST00000373831\|ENST00000373832\|ENST00000373833\|ENST00000398958\|ENST00000411533\|ENST00000478950\" Homo_sapiens.GRCh37.74.gtf > Homo_sapiens.GRCh37.74_subset.gtf
```

The transcript names given in the command above are all transcripts available for the genes CLEC6A, CLEC4D, HENMT1, and RCC1 in Ensembl version 74.

Homo_sapiens.GRCh37.74.sqlite

The Homo_sapiens.GRCh37.74.sqlite file is the sqlite database that the Ensembldb package creates from the corresponding gtf file. It was created using this command:

```
# ensDbFromGtf( # gtf = "Homo_sapiens.GRCh37.74_subset.gtf", # organism = "Homo_sapiens", # genomeVersion = "GRCh37", # version = 74)
```

infusion_fusions.txt

This is example data from the InFusion getting started page located at https://bitbucket.org/kokonech/infusion/wiki/Getting
## Description

Documentation for the JAFFA example data.

### jaffa_results.csv

This is example data from the described JAFFA example run documented at https://github.com/Oshlack/JAFFA/wiki/Example.

## Description

Documentation for the oncofuse example data.

### oncofuse.outfile

The example output from oncofuse was kindly provided by Lavinia G here: https://github.com/stianlagstad/chimeraviz/issues/47#issuecomment-409773158.

## Description

Documentation for the PRADA example data.

### PRADA.acc.fusion.fq.TAE.tsv

This is example data thankfully provided by PRADA authors Siyuan Zheng and Roeland Verhaak.

## Description

Documentation for the SOAPfuse example data.

### soapfuse_833ke_final.Fusion.specific.for.genes

This file has the results from a run of soapfuse-1.26 on the 833ke cell line. The program was ran with the standard configuration file.
Description

Documentation for the STAR-Fusion example data.

star-fusion.fusion_candidates.final.abridged.txt

This example data was retrieved from the STAR-Fusion github page June 2nd 2017.

select_transcript  Select which transcript to use (for plotting) for a GenePartner object

Description

This function takes a GenePartner object and creates a transcript data.frame with transcript information, including only the transcripts given by the parameter which_transcripts

Usage

select_transcript(gene_partner, which_transcripts = "exonBoundary")

Arguments

gene_partner  The GenePartner object to select a transcript for.
which_transcripts  This character vector decides which transcripts are to be plotted. Can be "exonBoundary", "withinExon", "withinIntron", "intergenic", or a character vector with specific transcript ids. Default value is "exonBoundary".

Details

select_transcript() selects which transcript to create by this prioritization:

1. Exon boundary transcripts. 2. Within exon transcripts. 3. Within intron transcripts. 4. Intergenic transcripts.

Value

A data.frame with transcript data.
Examples

```r
# Load data and example fusion event
defuse833ke <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuse833ke, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Load edb
edbSqliteFile <- system.file(
  "extdata",
  "Homo_sapiens.GRCh37.74.sqlite",
  package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)
# Get transcripts
fusion <- get_transcripts_ensembl_db(fusion, edb)
# Select transcript
transcriptsA <- select_transcript(upstream_partner_gene(fusion))
```

show.Fusion-method  
Show method for the Fusion class.

Description

Show method for the Fusion class.

Usage

```r
## S4 method for signature 'Fusion'
show(object)
```

Arguments

- `object`  
A Fusion object

Value

Shows information about a Fusion object.
show,PartnerGene-method

Show method for the PartnerGene class.

Description

Show method for the PartnerGene class.

Usage

```r
## S4 method for signature 'PartnerGene'
show(object)
```

Arguments

- `object`: A PartnerGene object

Value

Shows information about a PartnerGene object.

split_on_utr_and_add_feature

Split GRanges object based on cds

Description

This function will look for ranges (exons) in the GRanges object that has the coding DNA sequence starting or stopping within it. If found, these exons are split, and each exon in the GRanges object will be tagged as either "protein_coding", "5utr", or "3utr". The returned GRanges object will have feature values set in mcols(gr)$feature reflecting this.

Usage

```r
split_on_utr_and_add_feature(gr)
```

Arguments

- `gr`: The GRanges object we want to split and tag with feature info.

Value

An updated GRanges object with feature values set.
Examples

# Load fusion data and choose a fusion object:
defuseData <- system.file(
    "extdata",
    "defuse_833ke_results.filtered.tsv",
    package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- get_fusion_by_id(fusions, 5267)
# Create edb object
edbSqliteFile <- system.file(
    "extdata",
    "Homo_sapiens.GRCh37.74.sqlite",
    package="chimeraviz")
edb <- ensembldb::EnsDb(edbSqliteFile)
# Get all exons for all transcripts in the genes in the fusion transcript
allTranscripts <- ensembldb::exonsBy(edb,
    filter = list(
        AnnotationFilter::GeneIdFilter(
            c(
                partner_gene_ensembl_id(upstream_partner_gene(fusion)),
                partner_gene_ensembl_id(downstream_partner_gene(fusion))))),
    columns = c(
        "gene_id",
        "gene_name",
        "tx_id",
        "tx_cds_seq_start",
        "tx_cds_seq_end",
        "exon_id"))
# Extract one of the GRanges objects
gr <- allTranscripts[[1]]
# Check how many ranges there are here
length(gr)
# Should be 9 ranges
# Split the ranges containing the cds start/stop positions and add feature values:
gr <- split_on_utr_and_add_feature(gr)
# Check the length again
length(gr)
# Should be 11 now, as the range containing the cds_strat position and the range containing the cds_stop position has been split into separate ranges

upstream_partner_gene 

Get the upstream fusion partner gene

Description

This getter retrieves the upstream PartnerGene object.
This sets the upstream PartnerGene object of a Fusion object
upstream_partner_gene

Usage

upstream_partner_gene(x)

## S4 method for signature 'Fusion'
upstream_partner_gene(x)

upstream_partner_gene(object) <- value

## S4 replacement method for signature 'Fusion'
upstream_partner_gene(object) <- value

Arguments

x The Fusion object you wish to retrieve the upstream PartnerGene object for.

object The Fusion object you wish to set a new upstream PartnerGene object for.

value The new PartnerGene object.

Value

The upstream PartnerGene object.

Examples

# Load data
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Get the upstream fusion partner gene
upstream_partner_gene(fusion)

# Load data
defuseData <- system.file(
  "extdata",
  "defuse_833ke_results.filtered.tsv",
  package="chimeraviz")
fusions <- import_defuse(defuseData, "hg19", 1)
fusion <- fusions[[1]]
# Set the upstream PartnerGene object to be the same as the downstream PartnerGene object
upstream_partner_gene(fusion) <- downstream_partner_gene(fusion)
write_fusion_reference

Write fusion junction sequence to a fasta file

Description

This function will write the fusion sequence to a fasta file, using Biostring::writeXStringSet() .

Usage

write_fusion_reference(fusion, filename)

Arguments

- fusion: The Fusion object we want to create a fasta file from.
- filename: The filename to write to.

Value

Writes the fusion junction sequence to the given filename.

Examples

# Import the filtered defuse results
defuse833keFiltered <- system.file("extdata", "defuse_833ke_results.filtered.tsv", package="chimeraviz")
fusions <- import_defuse(defuse833keFiltered, "hg19", 1)
# Get a specific fusion
fusion <- get_fusion_by_id(fusions, 5267)
# Create temporary file to hold the fusion sequence
fastaFileOut <- tempfile(pattern = "fusionSequence", tmpdir = tempdir())
# Write fusion sequence to file
write_fusion_reference(fusion, fastaFileOut)
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