

# Package ‘Ularcirc’

May 26, 2026

**Type** Package

**Title** Shiny app for canonical and back splicing analysis (i.e. circular and mRNA analysis)

**Version** 1.31.0

**Description** Ularcirc reads in STAR aligned splice junction files and provides visualisation and analysis tools for splicing analysis. Users can assess backsplice junctions and forward canonical junctions.

**biocViews** DataRepresentation, Visualization, Genetics, Sequencing, Annotation, Coverage, AlternativeSplicing, DifferentialSplicing

**License** file LICENSE

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BSJ_details	<i>BSJ_details This function returns details of a BSJ string and returns a list of coordinates. Can accept two different formats, Ularcirc or generic.</i>
-------------	--

---

### Description

BSJ\_details This function returns details of a BSJ string and returns a list of coordinates. Can accept two different formats, Ularcirc or generic.

### Usage

```
BSJ_details(BSJ)
```

### Arguments

BSJ : backsplice junction as a string. See details below for example formats

### Examples

```
bsj <- 'chr14_99465814_chr14_99458278' # Historic Ularcirc format

bsj <- c("chr14_99465814_chr14_99458278", "chr22_20933778_chr22_20934245",
        "chr12_120155720_chr12_120154969", "chr4_143543508_chr4_143543973",
        "chr10_7285955_chr10_7276891")

BSJ_details(bsj)

bsj <- 'chr10:100923974-100926020:+' # generic format
BSJ_details(bsj)
```

---

```
bsj_fastq_generate    bsj_fastq_generate
```

---

## Description

Takes a circRNA predicted sequence and generates synthetic short sequence reads

## Usage

```
bsj_fastq_generate(  
  circRNA_Sequence,  
  fragmentLength = 300,  
  readLength = 100,  
  variations = 4,  
  headerID = ""  
)
```

## Arguments

```
circRNA_Sequence      : Linear sequence of a circRNA. i.e. the backsplice junction is the first and last  
                        base of this sequence  
fragmentLength       : Is the length the library fragment  
readLength           : The sequence read length  
variations           : Number of sequences returned for each read type. Note each sequence varia-  
                        tion will start at a unique location (where possible)  
headerID             : Character identifier that will be incorporated into sequence header
```

## Value

Returns a list of two DNAstring sets labelled "read1" and "read2" which correspond to forward and reverse read pairs.

## Examples

```
library('Ularcirc')  
  
# Generate a 500nt sequence containing A and which is flanked with GG and CC.  
circRNA_Sequence <- paste(rep('A',500),collapse='')  
circRNA_Sequence <- paste('GG',circRNA_Sequence, 'CC', sep='')  
# The GG and CC ends of sequence represent ends of linear exons that are circularised.  
# Therefore the backsplice junction (BSJ) is GGCC.  
# Generate reads that alternate over this BSJ  
  
fastqReads <- bsj_fastq_generate(circRNA_Sequence, fragmentLength=300, readLength=100,  
  variations = 4, # Four type I , II, III, and IV reads generated  
  headerID='circRNA_example') # Identifier incorporated in name of each sequence  
# The following will indicate 12 sequences are present in each list entry  
length(fastqReads$read1)  
length(fastqReads$read2)
```

```
# Can create fastq file as follows
Biostrings::writeXStringSet( fastqReads$read1, "circRNA_Sample_R1.fastq.gz",
                             compress = TRUE, format="fastq")
Biostrings::writeXStringSet( fastqReads$read2, "circRNA_Sample_R2.fastq.gz",
                             compress = TRUE, format="fastq")
```

---

bsj\_to\_circRNA\_sequence

*bsj\_to\_circRNA\_sequence*

---

## Description

Takes one BSJ coordinate and generates a predicted circular RNA sequence.

## Usage

```
bsj_to_circRNA_sequence(
  BSJ,
  geneID = NULL,
  genome,
  TxDb,
  annotationLibrary,
  reduce_candidates = TRUE,
  shiny = FALSE
)
```

## Arguments

**BSJ** : BSJ coordinate in the format of chr\_coordinate\_chr\_coordinate OR chr:coordinate-coorindate:strand.

**geneID** : The gene ID that the BSJ aligns to. Not essential as this can be identified from the BSJ coordinate, however time performance of function improved if this information can be provided.

**genome** : Is the length f the library fragment

**TxDb** : The sequence read length

**annotationLibrary** : annotation database. See details for example.

**reduce\_candidates** : IF multiple exon entries align to a single BSJ then either return longest entry (TRUE) or all entries (FALSE)

**shiny** : If TRUE then will setup shiny progress bars. Default is FALSE where a standard text progress bar is used.

## Details

Backsplice junction coordinates are typically reported as a character string. Two formats are recognised, ":" delimited (eg circExplorer, CIRI) or "\_" delimited (Ularcirc). The BSJ genomic coordinates are compared against the supplied gene model and exonic sequences from matching splice junctions are concatenated. This means the BSJ is the first and last nucleotide of the returned sequence. The current implementation will automatically check 0 or 1 base coordinates and any match is returned.

In some cases one BSJ will match multiple exon combinations. The default setting is to return the longest sequence. Alternatively all possibilities can be returned by setting `reduce_candidates` to `FALSE`. BSJ candidates that align to multiple exon combinations are added to duplicated list.

BSJ that do not align to any canonical junctions are returned as failed.

### Value

Returns a DNAstring object.

### Examples

```
library('Ularcirc')
TxDb <- TxDb.Hsapiens.UCSC.hg38.knownGene::TxDb.Hsapiens.UCSC.hg38.knownGene
genome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
annotationLibrary <- org.Hs.eg.db::org.Hs.eg.db

# Define BSJ. Following two formats are accepted
BSJ <- 'chr2:40430305-40428472:-' # SLC8A1
BSJ <- 'chr2_40430305_chr2_40428472' # SLC8A1

circRNA_sequence <- bsj_to_circRNA_sequence(BSJ, "SLC8A1", genome, TxDb, annotationLibrary)

# You can also retrieve sequence without passing gene annotation - but this is slower
# circRNA_sequence <- bsj_to_circRNA_sequence(BSJ, NULL, genome, TxDb, annotationLibrary)

TxDb <- TxDb.Hsapiens.UCSC.hg38.knownGene::TxDb.Hsapiens.UCSC.hg38.knownGene
genome <- BSgenome.Hsapiens.UCSC.hg38::BSgenome.Hsapiens.UCSC.hg38
# EXAMPLE1 (3 fail and 2 will produce sequences)
BSJ <- c("chr14_99465814_chr14_99458278", "chr22_20933778_chr22_20934245",
        "chr12_120155720_chr12_120154969", "chr4_143543508_chr4_143543973",
        "chr10_7285955_chr10_7276891")
GeneIDs <- c("SMARCA5", "MSLN", "RNF138", "KIAA0368", "CRKL")
circRNA_sequence <- bsj_to_circRNA_sequence(BSJ, GeneIDs, genome, TxDb, annotationLibrary)

# Returns a list with three items:
# (1) "identified" is a list of DNA strings from BSJ that aligned to FSJ coordinates of the gene model
# (2) "failed" is a character object of BSJ that did not align to FSJ coordinates of gene model. Each entry is
# named with gene ID.
# (3) "duplicates" (not implemented yet) identifies which BSJ returned multiple sequences
```

---

chimericFilters

*chimericFilters*

---

### Description

A wrapper function that prepares a list of filters that can be passed

### Usage

```
chimericFilters(
  BSjuncName = NULL,
  sortDir = "Descending",
  indexNumber = 1,
```

```

displayNumber = 10,
displayRADscore = FALSE,
RADcountThreshold = 10,
applyFSJfilter = FALSE
)

```

### Arguments

BSjuncName : A character string that represents a backsplice junction ID. Set when needing to extract a specific junction. Default NULL.

sortDir : Specifies how data is sorted, either "Descending" (default) or "Ascending".

indexNumber : Filter data according to this file index

displayNumber : Number of records to display in an shiny app

displayRADscore : Boolean. If TRUE then will apply/calculate RAD score

RADcountThreshold : Integer. The minimum count threshold required to calculate RAD score. i.e. A default RAD score of -1 will be applied to any BSJ with a count less than this score

applyFSJfilter : Boolean of whether to apply FSJ filter

---

chimericStats

*chimericStats*

---

### Description

Simple function that returns a list of basic stats obtained from a STAR chimeric file

### Usage

```
chimericStats(chimericDT)
```

### Arguments

chimericDT : Data table of chimeric junctions as provided by STAR aligner

### See Also

FilterChimericJuncs

### Examples

```

extdata_path <- system.file("extdata", package = "Ularcirc")
chimeric.file <- paste0(extdata_path, "/SRR444655_subset.Chimeric.out.junction.gz")
chimericDT <- Ularcirc::loadSTAR_chimeric(chimeric.file, returnColIdx = 1:14)
Ularcirc::chimericStats(chimericDT$data_set)
chimericDT$filtered <- Ularcirc::FilterChimericJuncs(chimericDT$data_set, canonicalJuncs = TRUE)
Ularcirc::chimericStats(chimericDT$filtered)

```

---

circSequence	<i>circSequence</i> This function returns pre-defined full length circRNA sequences.
--------------	--

---

**Description**

circSequence This function returns pre-defined full length circRNA sequences.

**Usage**

```
circSequence(circRNA_ID = "slc8a1")
```

**Arguments**

circRNA\_ID : Character string of a circRNA\_ID

**Examples**

```
circSequence(circRNA_ID="slc8a1")
```

---

Compatible\_Annotation\_DBs

*Compatible\_Annotation\_DBs*

---

**Description**

Interrogates Bioconductor databases and identifies those that are compatible with Ularcirc. Builds a list of commands that the user can copy to install the required database on their local computer. Once installed the databases are immediately available to Ularcirc upon re-starting the shiny app. This function requires connection to the internet.

**Usage**

```
Compatible_Annotation_DBs(search_term = "")
```

**Arguments**

search\_term : character string of a full or part name of a database. Will return only those entries that contain this search term. Not case sensitive.

**Value**

Returns a list of compatible annotation databases

**Examples**

```

# Get all Bioconductor annotation databases that are compatible with Ularcirc
library('BSgenome')
library('httpuv')
library('AnnotationHub')
# Prepare a dataframe of all compatible annotation databases
## Not run: compatible_DBs_human <- Compatible_Annotation_DBs("Hsapiens")

# Example of how to find a relevant database and load the relevant databases:
# This example find hg38 databases
idx <- grep(pattern="hg38", x= compatible_DBs_human[, "genome"])

if (!requireNamespace("BiocManager", quietly=TRUE))
  install.packages("BiocManager")
BiocManager::install(c(compatible_DBs[idx,]))

## End(Not run)

```

---

FilterChimericJuncs     *FilterChimericJuncs*

---

**Description**

A generic function that filters STAR chimeric junction files on certain genomic criteria (eg strand, same chromosome etc). Useful filter to remove the most obvious false positives. The default filter settings are suitable for circRNA discovery in humans / mice data sets.

**Usage**

```

FilterChimericJuncs(
  All_junctions,
  chromFilter = TRUE,
  strandFilter = TRUE,
  genomicDistance = c(200, 1e+05),
  canonicalJuncs = TRUE,
  fileID = c(-1),
  chrM_Filter = TRUE,
  invertReads = FALSE
)

```

**Arguments**

**All\_junctions** : data.table of chimeric reads from STAR aligner

**chromFilter** : when TRUE (default) both chimera parts have to align to same chromosome

**strandFilter** : when TRUE (default) both chimera parts have to align to same strand

**genomicDistance** : minimum and maximum distance filters of chimeric reads on chromosome. Only is applied if ChromFilter is TRUE and StrandFilter is TRUE

**canonicalJuncs** : Will include any canonical junctions (default TRUE). Note STAR keeps canonical junctions that do not conform to aligner rules.

fileID : Specify a file index. Useful if planing to concatenating all data sets into a single table.

chrM\_Filter : Filter out mitochondrial chimeric reads (default TRUE)

invertReads : Boolean that specifies in read strand should be inverted (default FALSE).

### See Also

SelectUniqueJunctions, loadSTAR\_chimeric

### Examples

```
extdata_path <- system.file("extdata",package = "Ularcirc")
chimeric.file <- paste0(extdata_path,"/SRR444655_subset.Chimeric.out.junction.gz")
chimericsDT <- Ularcirc::loadSTAR_chimeric(chimeric.file,returnColIdx = 1:14)
chimericsDT$filtered <- Ularcirc::FilterChimericJuncs(chimericsDT$data_set, canonicalJuncs = TRUE)
```

---

### FilterChimeric\_Ularcirc

*Wrapper function for Ularcirc shiny app which expects a list of objects to be returned*

---

### Description

NEED to ensure that unstranded boolean value is passed to this function Not tested via shiny app yet.

### Usage

```
FilterChimeric_Ularcirc(
  All_junctions,
  chromFilter = TRUE,
  strandFilter = TRUE,
  genomicDistance = c(200, 1e+05),
  canonicalJuncs = TRUE,
  fileID = c(-1),
  chrM_Filter = TRUE,
  invertReads = FALSE,
  unstranded = FALSE,
  summaryNumber = 50
)
```

### Arguments

All\_junctions : data.table of chimeric reads from STAR aligner

chromFilter : when TRUE (default) both chimera parts have to align to same chromosome

strandFilter : when TRUE (default) both chimera parts have to align to same strand

genomicDistance : minimum and maximum distance filters of chimeric reads on chromosome. Only is applied if ChromFilter is TRUE and StrandFilter is TRUE

canonicalJuncs : Will include any canonical junctions (default TRUE). Note STAR keeps canonical junctions that do not conform to aligner rules.

fileID : Specify a file index. Useful if planing to concatenating all data sets into a single table.

chrM\_Filter : Filter out mitochondrial chimeric reads (default TRUE)

invertReads : Boolean that specifies in read strand should be inverted (default FALSE).

unstranded : Boolean for if reads are unstranded

summaryNumber : Number (Integer) of records to display in shiny app

---

Junction\_Sequence\_from\_Genome  
*Grab\_BS\_Junc\_Sequence*

---

### Description

This function extracts genomic sequence that is likely to capture BSJ. Function does not cross validate to gene models.

### Usage

```
Junction_Sequence_from_Genome(SelectUniqueJunct_Value, GeneList)
```

### Arguments

GeneList : GeneList

SelectUniqueJunct\_value : a dataframe with columns names startDonor, strandDonor, startAcceptor

---

loadSTAR\_chimeric      *loadSTAR\_chimeric*

---

### Description

Loads chimeric output file from the STAR aligner and returns a list containing three items (a data table, alignment stats and command line).

### Usage

```
loadSTAR_chimeric(filename = NULL, ID_index = 0, returnColIdx = 1:21)
```

### Arguments

filename : filename of the STAR chimeric output file. Can be gzipped

ID\_index : An index (single integer) which will be added as a separate column in the returned data table. Useful when collating multiple files into one large matrix like object.

returnColIdx : Numeric index of columns to return. Default 1:15

**Details**

:

Reads in a text or gzipped chimeric output file generated by the STAR aligner. Function automatically detects if the last two lines contains meta-data (produced from STAR 2.7) onwards.

Returns a list of containing three items: (1) data\_set (2) alignmentStats and (3) commandLine.

The column names of data\_set are defined as c("chromDonor", "startDonor", "strandDonor", "chromAcceptor", "startAcceptor", "strandAcceptor", "JuncType", "RepeatLength\_L", "RepeatLength\_R", "ReadName", "FirstBase\_1stSeq", "CIGAR\_1stSeq", "FirstBase\_2ndSeq", "CIGAR\_2ndSeq", "Multimapping")

If ID\_index is set to a value greater than 0 then an additional column called "DataSet" is created.

Columns can be subsetted by defining returnColIdx with an integer value that correspond to order of column names listed above.

---

load_miRBase_fasta	<i>load_miRBase_fasta This function extracts species specific mirbase sequences as a Biostrings RNAstringset.</i>
--------------------	---

---

**Description**

load\_miRBase\_fasta This function extracts species specific mirbase sequences as a Biostrings RNAstringset.

**Usage**

```
load_miRBase_fasta(
  mirbaselocalDir = tempdir(),
  speciesCode = "hsa",
  filename = "hairpin.fa"
)
```

**Arguments**

mirbaselocalDir	: Directory where miRbase files are located. If files are not detected will download requested files from mirBase
speciesCode	: Three letter code of species (eg human = hsa). Used to subset species miRbase entries
filename	: The miRBase filename. Expects this to be a fasta formatted file. If file does not exist will download from mirBase.

---

```
miR_binding_site_Analysis
```

*miR\_binding\_site\_Analysis* This function analyses a genomic (RNA or DNA) sequence for matching miRNA seed sequences.

---

## Description

`miR_binding_site_Analysis` This function analyses a genomic (RNA or DNA) sequence for matching miRNA seed sequences.

## Usage

```
miR_binding_site_Analysis(  
  Sequence_to_examine,  
  species_code,  
  seed_length = 6,  
  seed_start = 2,  
  selected_miRs = NULL,  
  rev_comp = TRUE,  
  mirbaselocalDir = tempdir()  
)
```

## Arguments

`Sequence_to_examine` : RNA/DNA Sequence to be analysed for miRNA binding sites. Format can be a character string or Biostrings formatted object.

`seed_length` : Length of seed sequence of a miRNA (default 6).

`seed_start` : Starting position of seed sequence relative to mature miRNA sequence (default 2).

`selected_miRs` : names of miRNA.

`rev_comp` : Should function perform Reverse complement of `Sequence_to_examine` (default TRUE).

`mirbaselocalDir` : Directory where miRbase files are located. If files are not detected will download requested files from mirBase

`speciesCode` : Three letter code of species (eg human = hsa). Used to subset species miRbase entries

## Details

The function will convert T to U and scan for matching miRNA seed sequences. Will return a list containing the following outputs: `SeedMatchResult` - counts of all seeds. Value of -1 indicates seed not detected. `Total_miR_number` - the total number of miRNA scanned against. `miR_Seed_Lookup` - data frame of all miRNA seeds scanned against.

**Examples**

```
Seq <- circSequence()
output <- miR_binding_site_Analysis(Sequence_to_examine = Seq, species_code = "hsa", rev_comp = TRUE)

# display frequency of seed sequences.
# Values of -1 do not occur
head(output$SeedMatchResult)
```

---

plot\_AllJunctions      *plot\_AllJunctions*

---

**Description**

Plots a BSJ, FSJ and transcripts for a nominated gene. Output is combined onto a single page. This function effectively wraps plotting functions from plotgardener

**Usage**

```
plot_AllJunctions(
  assembly = "hg38",
  chrom,
  chromstart,
  chromend,
  BSJData,
  BSJ_colors = "black",
  FSJData,
  FSJ_colors = "black",
  geneSymbol
)
```

**Arguments**

assembly	: Genome assembly
chrom	: chromosome
chromstart	: Starting position of chromosome
chromend	: End position of chromosome
BSJData	: Backsplice junction data table
BSJ_colors	: Backsplice junction assigned colours
FSJData	: Forward junction data table
FSJ_colors	: Forward junction assigned colours
geneSymbol	: Gene symbol

**Value**

Returns a list of two DNAstring sets labelled "read1" and "read2" which correspond to forward and reverse read pairs.

**Examples**

```

library('Ularcirc')
# BSJ data.table
BSJ_data <- data.table::data.table(chrom1="chr2",
                                  start1=c(40139400, 40160764, 40428472, 40428472),
                                  end1=c(40139400, 40160764,40428472, 40428472),
                                  chrom2="chr2", start2=c(40178494,40178494,40430302,40430305),
                                  end2=c(40178494,40178494,40430302,40430305),
                                  score=c(13,20,360,1751))

# FSJ
FSJstarts1 <- c(40115630,40139677,40160865,40164985,40170350,40174721,
               40174843,40175282,40278771,40430302,40430305)
FSJstarts2 <- c(40139400,40160764,40164853,40170280,40174705,40174824,
               40175260,40178386,40428472,40453160,40512348)
FSJ_data <- data.table::data.table(chrom1="chr2", start1=FSJstarts1, end1=FSJstarts1,
                                   chrom2="chr2", start2=FSJstarts2, end2=FSJstarts2,
                                   score=c(225,825,685,666,633,596,517,542,685,101,171))

plot_AllJunctions(assembly="hg38", chrom="chr2",
                  chromstart=40096769, chromend=40611554,
                  BSJData=BSJ_data, FSJData=FSJ_data, geneSymbol="SLC8A1")

```

---

RAD\_score

*RAD\_score*


---

**Description**

Theoretically the position of backsplice junctions should be distributed randomly across a amplicon. This function calculates the read alignment distribution (RAD) of backsplice junctions between forward and reverse read pairs. The RAD score is calculated from CIGAR strings which can be used to identify type II and type III alignments.

**Usage**

```

RAD_score(
  CIGAR_1stSeg = NULL,
  CIGAR_2ndSeg = NULL,
  RADcountThreshold = 10,
  digits = 2
)

```

**Arguments**

CIGAR\_1stSeg : CIGAR string of the first segment.  
CIGAR\_2ndSeg : CIGAR string of the second segment  
RADcountThreshold : Minimum count threshold required to apply RAD score. If there are less than this many entries in CIGAR list then -1 is returned.  
digits : rounding of the RAD score to this many digits (default 2)

---

 SelectUniqueJunctions *SelectUniqueJunctions*


---

**Description**

Builds a summary table from chimeric data obtained from the STAR aligner. Assembles table with the requested number of top entries. Populates with RAD score and FSJ score.

**Usage**

```
SelectUniqueJunctions(
  BSJ_junctions,
  filterlist = chimericFilters(),
  unstranded = FALSE,
  FSJ_Junctions = NULL,
  shinyapp = FALSE
)
```

**Arguments**

BSJ\_junctions : Junction to display  
 filterlist : filterlist  
 unstranded : If TRUE will match reads from both strands.  
 FSJ\_Junctions : Junction to display.  
 shinyapp : Boolean. If true used to setup control status bars in shiny app.

**Details**

This is the workhorse for collated BSJ junctions from the input data. It will return selected rows of data (annotated) that will enable enhanced browsing of raw data on the fly.

Filter options: Junction abundance. Sort

---

 sequence\_from\_exon\_coords  
*sequence\_from\_exon\_coords*


---

**Description**

sequence\_from\_exon\_coords

**Usage**

```
sequence_from_exon_coords(genome, exon_df)
```

**Arguments**

genome : genome object  
 exon\_df : data frame of exons. Must have column with names "chrom", "start", "stop", "strand"

---

Ularcirc

*Ularcirc*

---

**Description**

When the function is invoked the Ularcirc shiny app is started. The starting screen has quickstart instructions on how to use the software. Please refer to the Ularcirc vignette for a more detailed workflow.

**Usage**

```
Ularcirc()
```

**Value**

Does not return anything

**Examples**

```
# The following commands will load the shiny app either through an RStudio session or  
# through your internet browser
```

```
library("Ularcirc")  
## Not run: Ularcirc()
```

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