

Package ‘iscream’

February 2, 2026

Title Make fast and memory efficient BED file queries, summaries and matrices

Version 1.1.1

Description BED files store ranged genomic data that can be queried even when the files are compressed. iscream can query data from BED files and return them in multiple formats: parsed records or their summary statistics as data frames or GenomicRanges objects, and matrices as matrix, GenomicRanges, or SummarizedExperiment objects. iscream also provides specialized support for importing methylation data.

URL <https://huishenlab.github.io/iscream/>,
<https://github.com/huishenlab/iscream/>

BugReports <https://github.com/huishenlab/iscream/issues>

License MIT + file LICENSE

Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 7.3.3

Depends R (>= 4.5)

LinkingTo Rcpp, RcppArmadillo, RcppProgress, RcppSpdlog, Rhtslib,
stringfish

Imports Rcpp, Matrix, data.table, methods, pbapply, parallelly,
stringfish,

Suggests BiocFileCache, BiocStyle, bsseq, ggplot2, ggridges, knitr,
microbenchmark, rmarkdown, GenomicRanges, IRanges, Rsamtools,
SummarizedExperiment, S4Vectors, testthat (>= 3.0.0)

VignetteBuilder knitr

Config/testthat/edition 3

NeedsCompilation yes

SystemRequirements htslib: htslib-devel (rpm) or libhts-dev (deb) &
tabix: htslib-tools (rpm) or tabix (deb) & GNU make

biocViews DataImport, Software, Sequencing, SingleCell, DNAMethylation

git_url <https://git.bioconductor.org/packages/iscream>

git_branch devel

git_last_commit b1bda1c

git_last_commit_date 2025-10-29

Repository Bioconductor 3.23

Date/Publication 2026-02-01

Author James Eapen [aut, cre] (ORCID: <<https://orcid.org/0000-0001-6016-3598>>),
 Jacob Morrison [aut] (ORCID: <<https://orcid.org/0000-0001-8592-4744>>),
 Nathan Spix [ctb],
 Hui Shen [aut, ths, fnd] (ORCID:
 <<https://orcid.org/0000-0001-9767-4084>>)

Maintainer James Eapen <james.eapen@vai.org>

Contents

| | |
|----------------------------------|----|
| check_files_exist | 3 |
| check_thread_count | 3 |
| Cpp_query_all | 4 |
| Cpp_set_log_level | 4 |
| Cpp_summarize_regions | 5 |
| get_df_string | 6 |
| get_granges_string | 7 |
| get_threads | 7 |
| htslib_version | 8 |
| make_mat | 8 |
| make_mat_bsseq | 10 |
| query_chroms | 11 |
| set_log_level | 12 |
| set_threads | 13 |
| summarize_meth_regions | 14 |
| summarize_regions | 16 |
| tabix | 18 |
| validate_log_level | 20 |
| verify_aligner_or_stop | 20 |
| verify_files_or_stop | 21 |
| verify_filetype | 21 |

| | |
|-------------------|-------------------------------|
| check_files_exist | <i>Check that files exist</i> |
|-------------------|-------------------------------|

Description

Check that files exist

Usage

```
check_files_exist(files_vec, error_file_prefix = "Bedfile")
```

Arguments

| | |
|-------------------|--|
| files_vec | A vector of file paths |
| error_file_prefix | Error message prefix for 'Bedfile' vs 'Tabix file' |

Value

TRUE if all input BED files have an associated tabix index file. FALSE if not

| | |
|--------------------|--|
| check_thread_count | <i>Check that the required threads are available</i> |
|--------------------|--|

Description

Check that the required threads are available

Usage

```
check_thread_count(  
  n_threads,  
  avail_threads = availableCores(),  
  opt_set = FALSE  
)
```

Arguments

| | |
|---------------|---|
| n_threads | The number of threads to check availability for |
| avail_threads | The number of threads that are available on the system. Defaults to <code>parallelly::availableCores()</code> |
| opt_set | Whether the <code>iscream.threads</code> options is set |

Value

`n_threads` if the requested number of threads are available and stops if not

Cpp_query_all*Query all methylation info into M and coverage matrices*

Description

Query all methylation info into M and coverage matrices

Usage

```
Cpp_query_all(
  bedfiles,
  regions,
  aligner,
  valInd,
  merged,
  sparse,
  prealloc,
  nthreads
)
```

Arguments

| | |
|----------|---|
| bedfiles | A vector of BED files |
| regions | A vector of regions |
| aligner | The aligner used to make the WGBS BED files, only for <code>make_mat_bsseq</code> |
| valInd | The index of the data column needed for the matrix, for <code>make_mat</code> |
| merged | Whether the input strands have been merged/collapsed |
| prealloc | The number of rows to initialize the matrices with |
| nthreads | Set number of threads to use overriding the "iscream.threads" option. See <code>?set_threads</code> for more information. |

Value

A list of one or two matrices, chromosome, position, and filename vectors

Cpp_set_log_level*spdlog Logging Lever Setter*

Description

A helper function to turn a logging level given as string into the current logging level

Usage

```
Cpp_set_log_level(name)
```

Arguments

| | |
|------|---|
| name | A string with the logging level. Value understood are, in decreasing verbosity 'trace', 'debug', 'info', 'warning', 'error', 'critical', and 'off'. Unrecognised names are equivalent to 'off'. |
|------|---|

Value

Nothing is returned.

Cpp_summarize_regions *Apply a function over BED file records within genomic features*

Description

This function should be called from `summarize_regions()` since there are few sanity checks on the C++ side.

Usage

```
Cpp_summarize_regions(
  bedfiles,
  regions,
  fun_vec,
  col_indices,
  col_names,
  aligner,
  mval = FALSE,
  region_rownames = FALSE,
  nthreads = 1L
)
```

Arguments

| | |
|-------------|--|
| bedfiles | A vector of BED file paths |
| regions | A vector of genomic regions |
| fun_vec | Vector of the armadillo-supported stats functions to apply over the CpGs in the 'regions': "sum", "mean", "median", "stddev", "variance" "count", "min", "max", and "range". |
| col_indices | A vector of genomic regions |
| col_names | A vector of genomic regions |
| mval | Calculates M values when TRUE, use beta values when FALSE |

region_rownames

Whether to set rownames to the regions strings. Not necessary if your regions vector is unnamed. If its names, then the "feature" column is set to the names and the rownames are set to the regions string

nthreads

Number of cores to use. See details.

Details

The optimal number of threads depends on the number of bedfiles, but is set to half the available OpenMP cores. See ?get_threads for more details. It can be manauully set with set_threads().

Value

A summary data.frame

get_df_string

DataFrame to region strings

Description

Convert DataFrame to a vector of strings. Set feature names in a "name" column

Usage

```
get_df_string(regions_df, feature_col = NULL)
```

Arguments

regions_df A data frame with "chr", "start" and "end" columns

feature_col The data frame column to use as the names of the output string vector

Value

A character vector

Examples

```
(df <- data.frame(chr = c("chr1", "chr2"), start = c(1, 5), end = c(4, 10)))
get_df_string(df)
```

get_granges_string *GRanges to region strings*

Description

Coerces GenomicRanges to chr:start-end strings with as.character. If any regions have the same start and end, as.character returns chr:start strings which are invalid for the htseq API. These are corrected to chr:start-start.

Usage

```
get_granges_string(gr)
```

Arguments

gr A GRanges object

Value

A character vector

Examples

```
if (requireNamespace("GenomicRanges", quietly = TRUE)) {  
  get_granges_string(GenomicRanges::GRanges(c("chr1:1-10", "chr2:15-20")))  
}
```

get_threads *Get the number of available threads*

Description

Gets the number of threads iscream is currently set to use, whether the "iscream.threads" option is set and how many threads are available for use. To set the number of threads use `set_threads()` or set the `iscream.threads` option in your `~/.Rprofile`. See `?set_threads` for more information.

Usage

```
get_threads()
```

Value

A named vector:

- `use_threads` = the number of threads iscream will use
- `opt_set` = whether the option was set by the user
- `avail_threads` = The number of available threads as reported by `parallelly::availableCores`

Examples

```
get_threads()
```

| | |
|----------------|--|
| htslib_version | <i>Get htslib version and available features</i> |
|----------------|--|

Description

Returns the version of htslib being used by iscream and whether features such as libdeflate support are available. This information may not always correspond to the htslib version used during iscream's installation if a different htslib version is available for linking at runtime.

Usage

```
htslib_version()
```

Value

None

Examples

```
htslib_version()
```

| | |
|----------|---|
| make_mat | <i>Make a matrix from a numeric column of BED files</i> |
|----------|---|

Description

Queries the provided regions and produces a matrix along with genomic positions as a named list (make_mat()), a RangedSummarizedExperiment (make_mat_se()), GRanges (make_mat_gr()). Parallelized across files using threads from the "iscream.threads" option.

Usage

```
make_mat(
  bedfiles,
  regions,
  column,
  mat_name = "value",
  sparse = FALSE,
  prealloc = 10000,
  nthreads = NULL
)
```

```

make_mat_se(
  bedfiles,
  regions,
  column,
  mat_name = "value",
  sparse = FALSE,
  prealloc = 10000,
  nthreads = NULL
)

make_mat_gr(
  bedfiles,
  regions,
  column,
  mat_name = "value",
  prealloc = 10000,
  nthreads = NULL
)

```

Arguments

| | |
|----------|--|
| bedfiles | A vector of BED file paths |
| regions | A vector, data frame or GenomicRanges of genomic regions. See details. |
| column | The index of the data column needed for the matrix |
| mat_name | What to name the matrix in the returned object |
| sparse | Whether to return a sparse matrix |
| prealloc | The number of rows to initialize the matrices with. If the number of loci are approximately known, this can reduce runtime as fewer resizes need to be made. |
| nthreads | Set the number of threads to use. Overrides the "iscream.threads" option. See <code>?set_threads</code> for more information. |

Details

The input regions may be string vector in the form "chr:start-end" or a GRanges object. If a data frame is provided, they must have "chr", "start", and "end" columns.

Value

- `make_mat()`: A named list of
 - the matrix with the value of interest
 - a character vector of chromosomes and numeric vector of base positions
 - a character vector of the input sample BED file names
- `make_mat_gr()`: if GenomicRanges is available, a GRanges
- `make_mat_se()`: if SummarizedExperiment is available, a RangedSummarizedExperiment

Examples

```
bedfiles <- system.file("extdata", package = "iscream") |>
  list.files(pattern = "[a|b|c|d].bed.gz$", full.names = TRUE)
# examine the bedfiles
colnames <- c("chr", "start", "end", "beta", "coverage")
lapply(bedfiles, function(i) knitr::kable(read.table(i, col.names = colnames)))

# make a vector of regions
regions <- c("chr1:1-6", "chr1:7-10", "chr1:11-14")
# make matrix of beta values
make_mat(bedfiles, regions, column = 4)
```

make_mat_bsseq

Make M/beta and coverage matrices from WGBS BED files

Description

Queries the CpG/CpH loci from provided regions and produces M/beta and coverage matrices with their genomic positions. Parallelized across files using threads from the "iscream.threads" option. The output of `make_mat_bsseq` may be used to create a BSseq object: `do.call(BSseq, make_mat_bsseq(...))`.

Usage

```
make_mat_bsseq(
  bedfiles,
  regions,
  aligner = "biscuit",
  mval = TRUE,
  merged = TRUE,
  sparse = FALSE,
  prealloc = 10000,
  nthreads = NULL
)
```

Arguments

| | |
|-----------------------|--|
| <code>bedfiles</code> | A vector of BED file paths |
| <code>regions</code> | A vector, data frame or GenomicRanges of genomic regions. See details. |
| <code>aligner</code> | The aligner used to produce the BED files - one of "biscuit", "bismark", "bsbolt". |
| <code>mval</code> | Whether to return M-values or beta-values with the coverage matrix. Defaults to M-value. Set <code>mval=FALSE</code> to get beta value matrix. |
| <code>merged</code> | Whether the input strands have been merged/collapsed |
| <code>sparse</code> | Whether to return a sparse matrix |
| <code>prealloc</code> | The number of rows to initialize the matrices with. If the number of loci are approximately known, this can reduce runtime as fewer resizes need to be made. |
| <code>nthreads</code> | Set the number of threads to use. Overrides the "iscream.threads" option. See <code>?set_threads</code> for more information. |

Details

The input regions may be string vector in the form "chr:start-end" or a GRanges object. If a data frame is provided, they must have "chr", "start", and "end" columns.

Value

A named list of

- coverage and either a beta- or M-value matrix
- a character vector of chromosomes and numeric vector of corresponding CpG base positions
- a character vector of the input sample names

Bitpacking limits

`make_mat_bsseq()` makes two matrices: M-value (or beta-value) and coverage. For speed and memory efficiency these two values are bitpacked during matrix creation so that only one matrix needs to be populated and resized. This matrix is unpacked into the two required matrices only after the matrix dimensions are known after querying all input files. The two values are packed using the INT16 type, which has an upper limit of 32,767, into one INT32. If the coverage values exceed 32,767, the upper limit of a 16-bit signed integer, it will be capped at the limit. Beta values will also be capped similarly, but any such beta values would indicate a bug in the aligner that produced the data.

Examples

```
bedfiles <- system.file("extdata", package = "iscream") |>
  list.files(pattern = "[a|b|c|d].bed.gz$", full.names = TRUE)
# examine the BED files
colnames <- c("chr", "start", "end", "beta", "coverage")
lapply(bedfiles, function(i) knitr::kable(read.table(i, col.names = colnames)))

# make a vector of regions
regions <- c("chr1:1-6", "chr1:7-10", "chr1:11-14")
mat <- make_mat_bsseq(bedfiles, regions)
# for BSseq object run
if (requireNamespace("bsseq", quietly = TRUE)) {
  do.call(bsseq::BSseq, mat)
}
```

query_chroms

Query the chromosomes or seqnames from a vector of BED files

Description

Query the chromosomes or seqnames from a vector of BED files

Usage

```
query_chroms(bedfiles, nthreads = NULL)
```

Arguments

bedfiles The vector of BED file paths
nthreads Set number of threads to use overriding the "iscream.threads" option. See ?set_threads for more information.

Value

A vector of seqnames

Examples

```
bedfiles <- system.file("extdata", package = "iscream") |>
  list.files(pattern = "[a|b|c|d].bed.gz$", full.names = TRUE)
query_chroms(bedfiles)
```

set_log_level *Set and get logging level*

Description

Set and get logging level

Usage

```
set_log_level(level = "info")
get_log_level()
```

Arguments

level The logging verbosity level to use

- "info": the default that gives provides basic information about the number of files and regions used in a function
- "debug": more verbose about row allocations, how many CpGs were found in a region, filename parsing etc. This mode cannot be used on more than one thread as R cannot output messages from multiple threads without crashing.
- "off": no logging

Value

- `set_log_level()`: None; sets the log level to the provided level
- `get_log_level()`: The current logging level as a string

Examples

```
set_log_level("info")
get_log_level()
```

set_threads*Set the number of available threads*

Description

Sets the "iscream.threads" option to n_threads. To see how many threads you have available see ?get_threads().

Usage

```
set_threads(n_threads)
```

Arguments

| | |
|-----------|------------------------------|
| n_threads | The number of threads to use |
|-----------|------------------------------|

Details

iscream uses OpenMP to parallelize certain functions. You can use as many threads as are available to you on your system to varying degrees of performance improvements. The get_threads() function uses `parallelly::availableCores()` to report the number of available threads. Although OpenMP can detect the number of available cores, on high performance computers (HPCs) with resource allocating job schedulers like SLURM, OpenMP may detect all available threads across the HPC and not limit itself to the cores that were allocated to you by the scheduler. If your system administrator has not set up any limits, this may result in your job taking resources from other jobs. If there are limits, trying to use more threads than those available will reduce iscream's performance. Job schedulers will typically have an environment variable (e.g. `SLURM_CPUS_ON_NODE` with SLURM) that gives you the actual number of available cores. Further, on hyperthreaded systems, this count may be double that of the available processors. Using hyperthreading does not guarantee any performance improvement - it may be better to set the number of threads to half the reported number. `parallelly::availableCores()` takes HPC scheduler/CRAN/Bioconductor limits into account when reporting the number of available threads but it may not reliably report hyperthreading ('system' or 'nproc'). To set the number of threads without having to call `set_threads()` in every session, put

```
options(iscream.threads = [n_threads])
```

in your .Rprofile See `help('Rprofile')` for information on startup options.

Functions currently using multithreading:

- `tabix()`, `tabix_gr()`, `tabix_raw()`
- `query_chroms()`
- `make_mat()`, `make_mat_se()`, `make_mat_gr()`, `make_mat_bsseq()`
- `summarize_regions()`, `summarize_meth_regions()`

Value

None. Sets the `iscream.threads` option to the requested number of threads if available

Examples

```
(ncores <- parallelly::availableCores()

set_threads(ncores)
```

summarize_meth_regions

Summarize methylation information over genomic regions

Description

Run summarizing functions on the CpG/CpH loci in BED files across genomic regions. Parallelized across files using threads from the `"iscream.threads"` option.

Usage

```
summarize_meth_regions(
  bedfiles,
  regions,
  fun = "all",
  aligner = "biscuit",
  feature_col = NULL,
  mval = TRUE,
  set_region_rownames = FALSE,
  nthreads = NULL
)
```

Arguments

| | |
|--------------------------|---|
| <code>bedfiles</code> | A vector of BED file paths |
| <code>regions</code> | A vector, data frame or GenomicRanges of genomic regions. See details. |
| <code>fun</code> | Function(s) to apply over the region. See details. |
| <code>aligner</code> | The aligner used to produce the BED files - one of "biscuit", "bismark", "bsbolt". |
| <code>feature_col</code> | Column name of the input <code>regions</code> data frame containing a name for each genomic region. Set only if the using a data frame as the input <code>regions</code> format. See details. |
| <code>mval</code> | Whether to calculate the M value ($\text{coverage} \times \beta$) or use the beta value when applying the function. |

set_region_rownames

Use the region strings as the returned data frame's rownames. Can be useful if you have a named regions and want both the regions strings rownames and the feature names. See details.

nthreads

Set number of threads to use overriding the "iscream.threads" option. See ?set_threads for more information.

Value

A data.frame

Supported functions

- Sum: "sum"
- Mean: "mean"
- Median: "median"
- Standard deviation: "stddev"
- Variance: "variance"
- Minimum: "min"
- Maximum: "max"
- Range: "range"
- No. of records in the region: "count"

The summarizing computations are backed by the Armadillo library. See https://arma.sourceforge.net/docs.html#stats_fns for futher details on the supported functions

Using feature identifiers

regions may be string vector in the form "chr:start-end", a GRanges object or a data frame with "chr", "start", and "end" columns. The feature column of the output will contain a "chr:start-end" identifier for each summarized region. To use other identifiers, like a gene name for a region instead of the coordinates, set the names of the vector or GRanges to those identifiers. These names will be used instead of the genomic region string to describe each feature in the output dataframe. If regions is a data frame make an additional column with the identifiers and pass that column name to feature_col. See examples.

Examples

```
# also see examples from ?summarize_regions

bedfiles <- system.file("extdata", package = "iscream") |>
  list.files(pattern = "[a|b|c|d].bed.gz$", full.names = TRUE)

# make a vector of regions
regions <- c("chr1:1-6", "chr1:7-10", "chr1:11-14")
summarize_meth_regions(bedfiles, regions)
names(regions) <- c("A", "B", "C")
summarize_meth_regions(bedfiles, regions, fun = c("mean", "stddev"), mval = FALSE)
summarize_meth_regions(bedfiles, regions, fun = "sum")
```

| | |
|-------------------|---|
| summarize_regions | <i>Summarize information over genomic regions from any BED file</i> |
|-------------------|---|

Description

Run summarizing functions on BED file records across genomic regions. Parallelized across files using threads from the "iscream.threads" option.

Usage

```
summarize_regions(
  bedfiles,
  regions,
  columns,
  col_names = NULL,
  fun = "all",
  feature_col = NULL,
  set_region_rownames = FALSE,
  nthreads = NULL
)
```

Arguments

| | |
|---------------------|--|
| bedfiles | A vector of BED file paths |
| regions | A vector, data frame or GenomicRanges of genomic regions. See details. |
| columns | A vector of indices of the numeric columns to be summarized |
| col_names | A vector of names to use for columns in the output |
| fun | Function(s) to apply over the region. See details. |
| feature_col | Column name of the input regions data frame containing a name for each genomic region. Set only if the using a data frame as the input regions format. See details. |
| set_region_rownames | Use the region strings as the returned data frame's rownames. Can be useful if you have a named regions and want both the regions strings rownames and the feature names. See details. |
| nthreads | Set number of threads to use overriding the "iscream.threads" option. See ?set_threads for more information. |

Value

A data.frame

Supported functions

- Sum: "sum"
- Mean: "mean"
- Median: "median"
- Standard deviation: "stddev"
- Variance: "variance"
- Minimum: "min"
- Maximum: "max"
- Range: "range"
- No. of records in the region: "count"

The summarizing computations are backed by the Armadillo library. See https://arma.sourceforge.net/docs.html#stats_fns for further details on the supported functions

Using feature identifiers

regions may be string vector in the form "chr:start-end", a GRanges object or a data frame with "chr", "start", and "end" columns. The feature column of the output will contain a "chr:start-end" identifier for each summarized region. To use other identifiers, like a gene name for a region instead of the coordinates, set the names of the vector or GRanges to those identifiers. These names will be used instead of the genomic region string to describe each feature in the output data frame. If regions is a data frame make an additional column with the identifiers and pass that column name to feature_col. See examples.

Examples

```
bedfiles <- system.file("extdata", package = "iscream") |>
  list.files(pattern = "[a|b|c|d].bed.gz$", full.names = TRUE)
# examine the bedfiles
colnames <- c("chr", "start", "end", "beta", "coverage")
lapply(bedfiles, function(i) knitr::kable(read.table(i, col.names = colnames)))

# make a vector of regions
regions <- c("chr1:1-6", "chr1:7-10", "chr1:11-14")
summarize_regions(bedfiles, regions, columns = c(4, 5), col_names = c("beta", "cov"))

# select functions
summarize_regions(
  bedfiles,
  regions,
  fun = c("mean", "stddev"),
  columns = c(4, 5),
  col_names = c("beta", "cov")
)

# add names to the regions
names(regions) <- c("A", "B", "C")
summarize_regions(
  bedfiles,
```

```

regions,
fun = "sum",
columns = 5,
col_names = "coverage"
)

# using `feature_col`
library(data.table)

# convert string vector to a data.table
regions_df <- data.table::as.data.table(regions) |>
_[, tstrsplit(regions, ":+-", fixed = FALSE, names = c("chr", "start", "end"))] |>
_[, start := as.integer(start)] |>
_[, feature := LETTERS[.I]][]
regions_df

summarize_regions(
  bedfiles,
  regions_df,
  fun = "sum",
  columns = 5,
  col_names = "coverage",
  feature_col = "feature"
)

```

tabix

Query records from tabixed BED files

Description

Query records from tabixed BED files

Usage

```

tabix(bedfiles, regions, aligner = NULL, col.names = NULL, nthreads = NULL)

tabix_gr(
  bedfiles,
  regions,
  aligner = NULL,
  col.names = NULL,
  zero_based = TRUE,
  nthreads = NULL
)

tabix_raw(bedfiles, regions, nthreads = NULL)

```

Arguments

| | |
|---|---|
| bedfiles | The BED files to be queried |
| regions | A vector, data frame or GenomicRanges of genomic regions. See details. |
| <td>The aligner used to produce the BED files - one of "biscuit", "bismark", "bsbolt". Will set the result data.table's column names based on this argument.</td> | The aligner used to produce the BED files - one of "biscuit", "bismark", "bsbolt". Will set the result data.table's column names based on this argument. |
| col.names | A vector of column names for the data columns of the result.table, not including "chr", "start", and "end". Set if your BED file is not from the supported aligners or is a general BED file. |
| nthreads | Set number of threads to use overriding the "iscream.threads" option. See ?set_threads for more information. |
| zero_based | Whether the input BED file has a zero-based start column - used when converting the result data frame to GenomicRanges. |

Details

Query method:

'iscream' has two methods to query records from BED files:

- the *tabix* shell executable: fast since its output can be redirected to a file (which `data.table::fread()` can then read) instead of having to allocate memory and store it during the query
- *iscream*'s tabix implementation, based on the *tabix* executable using *htslib*, but slower on large queries since it stores the records as they are found instead of writing to a file. However it's able to store each region's records independently instead of in a single file and is used in `make_mat()`, `make_mat_bsseq()`, and `summarize_regions()`.

When *iscream* is attached, it checks that the *tabix* executable is available with `Sys.which()` and, if available, sets `options("tabix.method" = "shell")`. `tabix()` then uses the *tabix* executable to make queries, except for `tabix_raw()`. If *tabix* is not found, *iscream* uses its tabix implementation. To use only *iscream*'s tabix implementation, set `options("tabix.method" = "htslib")`.

Input region formats:

The input regions format may be string vector in the form "chr:start-end", a data frame with "chr", "start" and "end" columns or a GRanges object. Input regions must be 1-based. When using "htslib" as the query method, if the input GRanges object of regions contains any single locus regions where the start and end positions are the same, iscream will notify that such regions were found and fixed as chr:start format strings are invalid for the htslib API (see `?get_granges_string`).

Value

- `tabix()`: A data frame
- `tabix_gr()`: A GRanges object for single files and GRangesList for multiple files. When making GRanges, the 0-based records from BED-files will be converted to 1-based with `GenomicRanges::makeGRangesFromDataFrame()`. Bismark's coverage files will not be converted as they are already 1-based and the ranges slot will be only one position.
- `tabix_raw()`: A named list of raw strings from the regions in the style of `Rsamtools::scanTabix`

Examples

```
bedfiles <- system.file("extdata", package = "iscream") |>
  list.files(pattern = "[a|b|c|d].bed.gz$", full.names = TRUE)
regions <- c("chr1:1-6", "chr1:7-10", "chr1:11-14")
tabix(bedfiles, regions, col.names = c("beta", "coverage"))
if (require("GenomicRanges", quietly = TRUE)) {
  tabix_gr(bedfiles, regions, col.names = c("beta", "coverage"))
}
tabix_raw(bedfiles, regions)
```

`validate_log_level` *Validate provided logging level*

Description

Only "info" and "debug" are currently supported, with "debug" only supported when using 1 thread

Usage

```
validate_log_level(level = get_log_level(), n_threads)
```

Arguments

| | |
|------------------------|--|
| <code>level</code> | The logging level to validate |
| <code>n_threads</code> | The number of threads that the next iscream function call will use |

Value

None; sets the log level to the provide level

Examples

```
set_log_level("info")
```

`verify_aligner_or_stop` *Validate provided aligner*

Description

Only "biscuit", "bismark", and "bsbolt" are currently supported

Usage

```
verify_aligner_or_stop(aligner)
```

Arguments

aligner The input aligner

Value

true; quits if the input is not among supported_aligners

verify_files_or_stop *Verify that BED files are tabixed*

Description

Verify that BED files are tabixed

Usage

```
verify_files_or_stop(bedfiles, verify_tabix = TRUE)
```

Arguments

bedfiles A vector of BED file paths
verify_tabix Whether to verify the presence of tabix files

Value

TRUE if all input BED files have an associated tabix index file. FALSE if not

verify_filetype *Verify that the input BED files are of the type specified by the input aligner*

Description

Verify that the input BED files are of the type specified by the input aligner

Usage

```
verify_filetype(bedfiles, aligner, stop_on_error = FALSE)
```

Arguments

bedfiles A vector of BED file paths
aligner The aligner chosen
stop_on_error Whether to warn or stop on aligner-filename mismatch

Value

TRUE if all input BED files have an associated tabix index file. FALSE if not

Index

* **internal**

- check_files_exist, 3
- check_thread_count, 3
- Cpp_query_all, 4
- Cpp_set_log_level, 4
- Cpp_summarize_regions, 5
- validate_log_level, 20
- verify_aligner_or_stop, 20
- verify_files_or_stop, 21
- verify_filetype, 21

- check_files_exist, 3
- check_thread_count, 3
- Cpp_query_all, 4
- Cpp_set_log_level, 4
- Cpp_summarize_regions, 5

- get_df_string, 6
- get_granges_string, 7
- get_log_level (set_log_level), 12
- get_threads, 7

- htslib_version, 8

- make_mat, 8
- make_mat_bsseq, 10
- make_mat_gr (make_mat), 8
- make_mat_se (make_mat), 8

- query_chroms, 11

- set_log_level, 12
- set_threads, 13
- summarize_meth_regions, 14
- summarize_regions, 16

- tabix, 18
- tabix_gr (tabix), 18
- tabix_raw (tabix), 18

- validate_log_level, 20