# Package 'kmeRtone'

March 8, 2024

```
Type Package
Version 1.0
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Title kmeRtone
Description Multi-purpose and transferrable k-meric enrichment/depletion analysis software.
SystemRequirements GNU make
Imports data.table (>= 1.15.0),
     R6 (>= 2.5.1),
     Rcpp (>= 1.0.12),
     R.utils (>= 2.12.3),
     openxlsx (>= 4.2.5.2),
     png (>= 0.1-8),
     RcppSimdJson (>= 0.1.11),
     venneuler (>= 1.1-4),
     stringi,
     curl,
     future,
     future.apply,
     isonlite,
     progressr,
     Biostrings,
     seqLogo
Depends R (>= 4.2)
Roxygen list(markdown = TRUE)
RoxygenNote 7.3.1
LinkingTo Rcpp, stringi, Rhtslib, zlibbioc
URL https://github.com/SahakyanLab/kmeRtone
BugReports https://github.com/SahakyanLab/kmeRtone/issues
Encoding UTF-8
License GPL-3 + file LICENSE
LazyData true
Suggests knitr,
     rmarkdown,
     testthat (>= 3.0.0)
Config/testthat/edition 3
VignetteBuilder knitr
```

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# Description

Add transparency to color.

# Usage

addAlphaCol(cols, alpha)

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### **Arguments**

cols Colors in hex format or R color code e.g. "red", "black", etc.

alpha Alpha value.

#### Value

Colors with alpha value in hex format.

bedToCoor

Convert a BED file to chromosome-separated csv files.

### Description

Convert a BED file to chromosome-separated csv files.

### Usage

```
bedToCoor(bed.path, output.path = "coordinate/", compress = TRUE)
```

### **Arguments**

bed.path A path to a BED file.

output.path Output directory path. It should be an empty directory. Default to coordinate/

compress Logical. If TRUE, compress the output CSV files. Default to TRUE.

### Value

None

buildControl

Build control regions

### Description

Build control regions

### Usage

```
buildControl(
  case,
  ctrl.rel.pos,
  genome,
  output.path = "control/",
  verbose = TRUE
)
```

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### **Arguments**

case Case in Coordinate class object format.

ctrl.rel.pos Control relative position.
genome Genome class object.

output .path Output directory path to save control coordinate.

verbose Boolean. Default is TRUE and will print progress updates.

#### Value

Control in Coordinate class object format.

buildKmerTable Count k-mers from given sequence(s) and build a data.table of k-mer

counts.

## Description

Only existed k-mers are returned in data.table object.

### Usage

```
buildKmerTable(dna.seqs, k, method = "auto", remove.N = TRUE)
```

### **Arguments**

dna.seqs String of sequence(s).

k Size of kmer.

method K-mer counting method: "Biostrings", "sliding", or "auto". Default is "auto";

For k > 8, sliding method is used.

remove.N Remove unknown base? Default is TRUE.

#### Value

A data.table object with column kmer and N.

buildRangedKmerTable

buildMidPatternKmerTable

Count k-mers with specified middle pattern from given sequence(s) and build a data.table of k-mer counts.

### **Description**

Only existed k-mers are returned in data.table object.

### Usage

```
buildMidPatternKmerTable(dna.seqs, k, mid.patterns, remove.N = TRUE)
```

### Arguments

```
dna.seqs String of sequence(s).

k Size of kmer.

mid.patterns Middle patterns.

remove.N Remove unknown base? Default is TRUE.
```

#### Value

A data.table object with column kmer and N.

buildRangedKmerTable Count kmers from a sequence in given ranges and build a data.table of k-mer counts.

### **Description**

Count kmers from a sequence in given ranges and build a data.table of k-mer counts.

### Usage

```
buildRangedKmerTable(
  dna.seq,
  starts,
  ends,
  k,
  method = "sliding",
  chopping.method = "auto",
  remove.N = TRUE
)
```

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### **Arguments**

dna.seq String of sequence.

starts Start positions.
ends End positions.
k Size of kmer.

method Method options: "sliding" or "chopping". Chopping consumes a lot of memory

for extremely long sequence using "substring" method. Using "Biostrings" for

k > 12 is memory consuming. Default is "sliding".

chopping.method

Chopping method: "Biostrings" or "substring". Default is "auto".

remove.N Remove unknown base N? Default is TRUE.

#### Value

A data.table object with column kmer and N.

buildRESTurl Function constructs a URL for a REST API call by appending query

parameters.

# Description

Function constructs a URL for a REST API call by appending query parameters.

# Usage

```
buildRESTurl(url, .list = list(), ...)
```

### **Arguments**

url Base URL of the REST API.

.list A list of named query parameters.

... additional optional arguments

### Value

string of the full REST API URL.

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calKmerSkew	Function calculates the skew of k-mers based on their occurrence in positive and negative strands.

#### **Description**

Function calculates the skew of k-mers based on their occurrence in positive and negative strands.

#### Usage

```
calKmerSkew(kmer.table)
```

#### **Arguments**

kmer.table data.table wit

data.table with columns: kmer, pos\_strand, neg\_strand.

#### Value

data.table with the kmer\_skew column.

calPWM Calculate position weight matrix of overlapping sequences. Simulation of human population is based on single nucleotide variation.

### **Description**

Calculate position weight matrix of overlapping sequences. Simulation of human population is based on single nucleotide variation.

### Usage

```
calPWM(
  kmers,
  pseudo.num = 0,
  bg.prop = c(a = 0.295, c = 0.205, g = 0.205, t = 0.295),
  output = "PWM"
)
```

#### **Arguments**

kmers A vector of k-mers to overlap.

pseudo.num Pseudo-number to avoid numerical instability due to lack of base at a position.

Default is zero i.e. no pseudo-number.

bg.prop Background proportion of bases. Default is c(a = 0.295, c = 0.205, g = 0.205, t

= 0.295) which is observed in human genome.

output Output matrix type. Options are PCM, PPM, and PWM which refer to position

count/probability/weight matrix. Default is PWM.

#### Value

A position count/probability/weight matrix.

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catHeader

Function prints a given message in a formatted header with borders.

#### **Description**

Function prints a given message in a formatted header with borders.

#### **Usage**

catHeader(msg)

#### **Arguments**

msg

message to be printed within the header.

Coordinate

Function constructs an R6 object for handling coordinate data. The object includes methods for loading, manipulating, and analyzing coordinate data.

### **Description**

Function constructs an R6 object for handling coordinate data. The object includes methods for loading, manipulating, and analyzing coordinate data.

Function constructs an R6 object for handling coordinate data. The object includes methods for loading, manipulating, and analyzing coordinate data.

#### **Public fields**

root\_path A path to a directory containing coordinate files.

single\_len Single case length e.g. damage length. Default is NULL.

is\_strand\_sensitive Coordinate strand polarity. Default is TRUE.

merge\_replicate Merge coordinate from different replicates. Default is TRUE.

rm\_dup Remove duplicate entry in the coordinate table. Default is TRUE.

add\_col\_rep If add\_col\_rep is TRUE, column replicate is added to the coordinate table. Default is TRUE.

paths Individual coordinate files.

rep\_names Replicate names determined from coordinate subdirectory.

chr\_names Chromosome names determined from filenames.

coor Chromosome-named list of coordinate data.table.

 $is\_kmer\ A\ data.table\ of\ is\_kmer\ status.$  The first column is original is\\_kmer\ status.

k K-mer size when is\_kmer is TRUE. When is\_kmer is FALSE, k is NA.

ori\_first\_index Original chromosome-separated table first index is either starting from zero or one.

load\_limit Maximum coordinate table loaded.

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#### Methods

```
Public methods:
  • Coordinate$new()
  • Coordinate$[()
  • Coordinate$mark_overlap()
  • Coordinate$print()
  • Coordinate$map_sequence()
  • Coordinate$clone()
Method new(): Create a new Coordinate class
 Usage:
 Coordinate$new(
    root.path,
    single.len,
    is.strand.sensitive,
    merge.replicate,
    rm.dup,
    add.col.rep,
    is.kmer,
   ori.first.index,
   load.limit
 )
 Arguments:
 root.path A path to a directory containing either: (1) chromosome-separated coordinate files
     (assume replicates for subdirectories) OR (2) bedfile. (assume replicates for bedfiles)
 single.len Single case length e.g. damage length. Default is NULL
 is.strand.sensitive A boolean whether strand polarity matters. Default is TRUE.
 merge.replicate Merge coordinate from different replicates. Default is TRUE. If not merg-
     ing, duplicates will give weight to the kmer counting. If add_col_rep, merged coordinate
     will contain column replicate e.g. "rep1&rep2".
 rm.dup Remove duplicates in each replicate. Default is FALSE Default is FALSE
 add.col.rep Add column replicate to coordinate table.
 is.kmer Is the coordinate refers to k-mer i.e. expanded case? Default is FALSE.
 k Length of k-mer if is_kmer is TRUE.
 ori.first.index Zero- or one-based index. Default is 1.
 load.limit Maximum coordinate data.table loaded. Default is 1.
 Returns: A new Coordinate object.
```

Method [(): Calling coordinate table by loading on demand. Maximum load is determine by load\_limit field.

```
Usage:
Coordinate $[(
  chr.name,
  state = "current",
 k,
  reload = FALSE,
  rm.other.cols = TRUE
```

countBaseComposition 11

Arguments:

chr. name Chromosome name. It can be a vector of chromosomes.

state Coordinate state: "current", "case", "kmer". The coordinate state is changed automatically on demand. Default is "current".

k K-mer size. If state is "kmer", k is needed to expand the coordinate.

reload Reload the coordinate table from the root.path. Default is TRUE.

rm.other.cols Remove unnecessary columns for kmeRtone operation.

Returns: A single or list of data.table coordinate of requested chromosome.

**Method** mark\_overlap(): Mark overlapping regions in the coordinate table. A column name is\_overlap is added.

Usage:

Coordinate\$mark\_overlap()

Arguments:

chr.names Chromosome names

Returns: New column is\_overlap is added.

Method print(): Print Coordinate object parameters.

Usage:

Coordinate\$print()

Returns: Message of Coordinate object parameters.

**Method** map\_sequence(): Get corresponding sequence from the loaded coordinate.

Usage:

Coordinate\$map\_sequence(genome)

Arguments:

genome Genome object or vector of named chromosome sequences.

Returns: New column seq.

**Method** clone(): The objects of this class are cloneable with this method.

Usage:

Coordinate\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

countBaseComposition

Function performs an analysis of base composition including sequence frequency, PWM calculations, and G/C content at various window sizes.

### **Description**

Function performs an analysis of base composition including sequence frequency, PWM calculations, and G/C content at various window sizes.

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#### Usage

```
countBaseComposition(case, genome, case.pattern, output.path = "./")
```

### **Arguments**

case A Coordinate class object or similar structure.

genome Genome class object or similar structure.

case.pattern String patterns to consider in the analysis.

output.path Output path for saving the analysis results.

countChoppedKmers Function chops k-mers within specified ranges of a sequence and

counts them. It uses either a substring method or functionalities from

the Biostrings package.

# Description

Function chops k-mers within specified ranges of a sequence and counts them. It uses either a substring method or functionalities from the Biostrings package.

## Usage

```
countChoppedKmers(dna.seq, starts, ends, k, method = "auto")
```

### **Arguments**

dna.seq A string of sequence.

starts Start positions.
ends End positions.
k Size of kmer.

method Method: "Biostrings" or "substring". Default is Biostrings.

### Value

A k-mer-named vector of counts.

countDistribution 13

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Function performs an analysis of the distribution of genomic cases, including: A) Check case distribution in:

- 1. replicates
- 2. chromosomes
- 3. strands B) Check case base composition and filter out other case.patterns
- 4. draw seqLogo in 100 or 101 base context
- 5. calculate G+C percentage Then, it generates various plots like bar plots and Venn/Euler diagrams.

### Description

Function performs an analysis of the distribution of genomic cases, including: A) Check case distribution in:

- 1. replicates
- 2. chromosomes
- 3. strands B) Check case base composition and filter out other case.patterns
- 4. draw seqLogo in 100 or 101 base context
- 5. calculate G+C percentage Then, it generates various plots like bar plots and Venn/Euler diagrams.

#### Usage

```
countDistribution(case, genome, case.pattern, output.path = "./")
```

#### Arguments

case A Coordinate class object or similar structure for genomic data.

genome Genome class object or similar structure.

case.pattern String patterns to consider in the analysis.

output.path Output path for saving the analysis results.

countKmers

Count k-mers from string(s) using a simple hash table.

# Description

Count only observed k-mers. Biostrings::oligonucleotideFrequency reports all possible k-mers. For k > 12, the memory for creating empty k-mer counts spiked and crashed the R session.

#### Usage

```
countKmers(sequences, k)
```

#### **Arguments**

sequences Sequence strings. k Size of k-mer.

#### Value

A vector of k-mer counts. The counts of multiple sequences are combined, similar to Biostrings::oligonucleotideFrequences implify.as "collapsed".

countMidPatternContext

Locate a middle sequence pattern and count its sequence context.

### **Description**

Locate a middle sequence pattern and count its sequence context.

#### Usage

countMidPatternContext(sequence, mid\_pattern, window, context\_pattern)

#### **Arguments**

sequence A sequence to slide.

mid\_pattern A middle pattern to search for.
window Size of a surrounding window.

context\_pattern

A context pattern to search for.

#### Value

A numeric vector of count.

countMidPatternContext2

Locate a middle sequence pattern and count its sequence context.

#### **Description**

This function searches for a specified middle pattern within a given sequence. It then counts the occurrences of specific context patterns within a defined window size around the middle pattern. The function returns a map where keys are the counts of context patterns found and values are the frequencies of these counts.

#### Usage

countMidPatternContext2(sequence, mid\_pattern, window, context\_patterns)

countMidPatternKmers 15

#### **Arguments**

sequence A string representing the sequence to be analyzed.

mid\_pattern A string representing the middle pattern to search for within the sequence.

window An integer specifying the size of the surrounding window around the middle

pattern.

context\_patterns

A vector of strings representing the context patterns to search for within the

window.

#### Value

A std::unordered\_map<int,int> where keys are the counts of context patterns found and values are the frequencies of these counts.

### **Examples**

```
sequence <- "ATCGATCGA"
mid_pattern <- "CG"
window <- 5
context_patterns <- c("AT", "GA")
countMidPatternContext2(sequence, mid_pattern, window, context_patterns)</pre>
```

# Description

This function scans through each sequence in the provided vector, locating a specified middle pattern. For each occurrence of the middle pattern, the function extracts and counts the surrounding k-mers. The k-mers are identified based on the given k-mer size and centered around the middle pattern.

# Usage

```
countMidPatternKmers(sequences, k, mid_pattern)
```

### **Arguments**

sequences A vector of strings, each representing a sequence to be analyzed.

k An integer specifying the size of the k-mers to be extracted and counted.

mid\_pattern A string representing the middle pattern to search for within each sequence.

### Value

A std::unordered\_map with k-mers as keys and their counts as values.

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### **Examples**

```
sequences <- c("ATCGATCGA", "GCGCATGCA") 
 k <- 5 
 mid_pattern <- "CG" 
 countMidPatternKmers(sequences, k, mid_pattern)
```

countMidPatternRangedKmers

Count k-mers in given ranges of a sequence.

### **Description**

Slide and update the cumulated table count.

### Usage

```
countMidPatternRangedKmers(sequence, starts, ends, k, mid_pattern)
```

### **Arguments**

sequence A sequence to count.
starts Start positions.
ends End positions.
k K-mer size.

mid\_pattern A middle pattern to search for.

#### Value

A k-mer-named vector of count.

countPointContext

Ccount sequence context of given point positions.

### **Description**

Count sequence context of given point positions.

### Usage

```
countPointContext(sequence, points, len, window, context_pattern)
```

### **Arguments**

sequence A sequence to slide.
points Middle point positions.

len Length of the middle point. Default to 1.

window Size of a surrounding window.

context\_pattern

A context pattern to search for.

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#### Value

A numeric vector of count.

countPointContext2

Ccount sequence context of given point positions.

### Description

Ccount sequence context of given point positions.

### Usage

```
countPointContext2(sequence, points, len, window, context_patterns)
```

### Arguments

sequence A sequence to slide.

points Middle point positions.

len Length of the middle point.

window Size of a surrounding window.

context\_patterns

Context patterns to search for.

### Value

A named vector of frequency of counts.

countRangedKmers

Count k-mers in given ranges of a sequence.

# Description

Slide and update the cumulated table count.

# Usage

```
countRangedKmers(sequence, starts, ends, k)
```

# Arguments

sequence A sequence to count.
starts Start positions.
ends End positions.
k K-mer size.

#### Value

A k-mer-named vector of count.

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countRevCompKmers

Count reverse complement sequence from its opposite strand. Build for k-mer table generated from initKmerTable function but applicable to others with the same format.

### Description

Count reverse complement sequence from its opposite strand. Build for k-mer table generated from initKmerTable function but applicable to others with the same format.

### Usage

countRevCompKmers(kmer.table)

### **Arguments**

kmer.table

A data.table of k-mer with at least 3 columns: kmer, pos\_strand, and neg\_strand. Splitted k-mer columns: kmer\_part1 and kmer\_part2 is supported.

#### Value

Updated k-mer table.

countSlidingWindow

Count sequence content in a sliding window of a sequence.

### Description

Count sequence content in a sliding window of a sequence.

#### Usage

countSlidingWindow(sequence, window, pattern)

# Arguments

sequence A sequence to slide.
window Size of a window.

pattern A pattern to search for.

# Value

A numeric vector of count.

countSlidingWindow2 19

countSlidingWindow2

Count sequence content in a sliding window of a sequence.

### **Description**

Count sequence content in a sliding window of a sequence.

#### Usage

countSlidingWindow2(sequence, window, patterns)

#### **Arguments**

sequence A sequence to slide. window Size of a window.

patterns Patterns of the same size to search for.

#### Value

Named vector of frequency of count.

count\_substring\_fixed Count sequence content in a given sequence.

### **Description**

stringi has no function that search within substring without memory copy it. This function has two versions. One without the need to memory copy denoted as \*\*\*. The only downside to this is std::string::find cannot stop searching past end of substring. I manage to at least stop it as soon as possible. If the pattern is long and rare, it won't stop until it find post-substring pattern. The other version is memory copy substring but as this operation is in the loop, the memory is still within comfortable range. c++17 has std::string\_view that solve this but still new and not widely available. Use count\_substring\_regex to avoid memory copy.

## Usage

```
count_substring_fixed(sequence, start, end, pattern)
```

#### **Arguments**

sequence A sequence to map.
start Start positions.
end End positions.

pattern A pattern to search for.

#### Value

A numeric vector of count.

20 downloadNCBIGenomes

```
count_substring_regex Count sequence content in a given sequence.
```

### **Description**

stringi has no function that search within substring without memory creating it. This function solve that. Unlike count\_substring\_fixed, this function does not need to memory copy substring.

#### Usage

```
count_substring_regex(sequence, start, end, pattern)
```

### Arguments

sequence A sequence to map. start Start positions. end End positions.

pattern A regex pattern to search for within start and end positions.

#### Value

A numeric vector of count.

downloadNCBIGenomes Function downloads genome fasta files from the NCBI FTP database.

Users can provide either organism names or an assembly summary

data table.

### **Description**

Supports options for splitting multi-header fasta files and overwriting existing files.

# Usage

```
downloadNCBIGenomes(
  asm,
  species,
  db,
  output.dir = "./",
  split.fasta = FALSE,
  overwrite = FALSE
)
```

### **Arguments**

asm NCBI assembly summary data.table

species Species names.

db Database record to use: refseq or genbank

output.dir Output directory path. Default is current directory.

split.fasta NCBI fasta files are multi-header. Split them? Default is FALSE.

overwrite Overwrite any existed genome file? Default is FALSE to skip the download.

downloadUCSCgenome

#### Value

Genome fasta file(s) named according to the FTP database convention.

downloadUCSCgenome

Function downloads chromosome-separated fasta genome sequences from the UCSC database. Users can specify a genome name, an output folder, and a specific chromosome or chromosomes. There's an option to choose the download method as well.

### **Description**

Function downloads chromosome-separated fasta genome sequences from the UCSC database. Users can specify a genome name, an output folder, and a specific chromosome or chromosomes. There's an option to choose the download method as well.

### Usage

```
downloadUCSCgenome(genome.name, output.path, chr.name, method = "curl")
```

### **Arguments**

genome.name Genome name (e.g., hg19, hg38, mm19).

output.path Output folder for the downloaded sequences.

chr. name Specific chromosome to download; defaults to all if unspecified.

method Download method for the download.file function.

### Value

An output folder containing chromosome-separated fasta files.

example\_genome\_coor

Example genome coordinate file

### Description

Below is an example code that generates random genomic coordinates.

library(data.table) library(kmeRtone)

1. Randomly generate genomic positions and save results dir.create("./data", showWarnings = FALSE)

```
set.seed (1234) \ for (chr \ in \ 1) \ genomic\_coor <- \ data.table:: data.table( seqnames = paste0("chr", chr), start = sample( \ x = 10000:10000000, size = 100000, replace = FALSE), width = 2)
```

```
data.table::fwrite(
    genomic_coor,
    paste0("./data/chr", chr, ".csv")
)
```

#### Usage

```
example_genome_coor
```

#### **Format**

A data frame with 1001 rows and 3 columns

**seqnames** Chromosome number of the recorded biological event, e.g. DNA strand breaks **start** 5' start position of the recorded biological event **width** Sequence width of the recorded biological event, e.g. 2 for a DNA strand break

```
example_kmeRtone_score
```

Example 2-mer enrichment/depletion scores

#### **Description**

Below is an example code that generates random genomic coordinates and runs the default kmeRtone SCORE function to quantify the k-meric enrichment and depletion.

library(data.table) library(kmeRtone)

1. Randomly generate genomic positions and save results dir.create("./data", showWarnings = FALSE)

```
set.seed(1234) \ for(chr \ in \ 1:22) \ genomic\_coor <- \ data.table::data.table( \ seqnames = paste0("chr", chr), start = sample( \ x = 10000:10000000, size = 100000, replace = FALSE ), width = 2 )
```

```
data.table::fwrite(
    genomic_coor,
    paste0("./data/chr", chr, ".csv")
)
```

#' 2. Run kmeRtone score function kmeRtone::kmeRtone( case.coor.path="./data", genome.name="hg19", strand.sensitive=FALSE, k=2, ctrl.rel.pos=c(80, 500), case.pattern=NULL, single.case.len=2, out-put.dir="output", module="score", rm.case.kmer.overlaps=FALSE, merge.replicate=TRUE, kmer.table=NULL, verbose=TRUE)

### Usage

```
example_kmeRtone_score
```

#### **Format**

```
A data frame with 1001 rows and 3 columns
```

```
case Case k-mers, e.g. damage k-mer counts
```

case\_skew Case k-mers skews, e.g. skew of the damage k-mers counts

control control k-mers, e.g. damage k-mer counts

control\_skew control k-mers skews, e.g. skew of the damage k-mers counts

kmer K-meric sequence

z Intrinsic susceptibility z-score for each k-mer

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### **Source**

https://github.com/SahakyanLab/kmeRtone/tree/master/README.md

**EXPLORE** 

Function generates various exploratory analyses.

### **Description**

Function generates various exploratory analyses.

### Usage

```
EXPLORE(
  case.coor.path,
  genome.name,
  strand.sensitive,
  k,
  case.pattern,
  output.path,
  case,
  genome,
  control,
  genome.path,
  single.case.len,
  rm.dup,
  case.coor.1st.idx,
  coor.load.limit,
  genome.load.limit,
  genome.fasta.style,
  genome.ncbi.db,
  use.UCSC.chr.name,
  verbose
)
```

#### **Arguments**

```
case.coor.path Path to case coordinates.
                  Genome name (e.g., hg19, hg38).
genome.name
strand.sensitive
                  Boolean indicating if strand sensitivity is considered.
k
                  K-mer size.
                  String patterns to consider in the analysis.
case.pattern
                  Output directory path for exploration plots.
output.path
                  Coordinate class object or similar structure for case data.
case
genome
                  Genome class object or similar structure.
                  Control class object or similar structure.
control
genome.path
                  Path to genome fasta files.
```

24 extractKmers

```
single.case.len
                 Length of single cases.
                 Boolean indicating if duplicates should be removed.
rm.dup
case.coor.1st.idx
                 Indexing of case coordinates.
coor.load.limit
                 Maximum number of coordinates to load.
genome.load.limit
                 Maximum number of genome data to load.
genome.fasta.style
                 Fasta file style for genome data.
genome.ncbi.db NCBI database for genome data.
use.UCSC.chr.name
                 Boolean indicating if UCSC chromosome naming is used.
                 Boolean indicating if verbose output is enabled.
verbose
```

#### Value

Output directory containing exploration plots.

extractKmers

Extract k-mers from a given Coordinate object and Genome objects

#### **Description**

A k-mer table is initialized and updated in every chromosome-loop operation. There are 3 modes of extraction. (1) When k is smaller than 9 or k is larger than 15, the k-mer is extracted in a standard way. A k-mer table with every possible k-mers is created and updated. (2) For k between 9 and 13, the k-mer sequence is split to half to reduce memory usage significantly. e.g. ACGTACGTA will become ACGT ACGTA. (3) When k is larger than 14, k-mers are extracted the same way as (1) but the k-mer table is grown or expanded for every new k-mer found.

# Usage

```
extractKmers(
  coor,
  genome,
  k,
  central.pattern = NULL,
  rm.overlap.region = TRUE,
  verbose = TRUE
)
```

### **Arguments**

coor Coordinate class object.
genome Genome class object.
k Length of k-mer.

```
central.pattern
Central pattern of the k-mer, if applicable.

rm.overlap.region
Boolean indicating if overlapping regions should be removed. Default is TRUE.

verbose
Boolean indicating if verbose output is enabled.
```

#### Value

A k-mer table with counts for each k-mer.

generateGenicElementCoor

Function processes UCSC genePred tables to generate coordinates for various genic elements like introns, exons, CDS, UTRs, and upstream and downstream regions. It handles these coordinates with consideration for strand sensitivity and genome information.

#### **Description**

All the operations in here are vectorized. If the table is big, expect a spike in memory. Using ncbiRefSeq table and genome hg38, the memory is stable at 4-5 GB. I can utilise data.table package to process by chunk if needed. Original table is zero-based open-end index. The indexing system is changed temporarily to follow Rs system. The output coordinate table is one-based close-end index. Critical information based on UCSC Genome website: Column Explanation bin Indexing field to speed chromosome range queries. (Only relevant to UCSC program) name Name of gene (usually transcript\_id from GTF) chrom Reference sequence chromosome or scaffold strand + or - for strand txStart Transcription start position (or end position for minus strand item) txEnd Transcription end position (or start position for minus strand item) cdsStart Coding region start (or end position for minus strand item) cdsEnd Coding region end (or start position for minus strand item) exonCount Number of exons exonEnds Exon end positions (or start positions for minus strand item) exonStart Exon start positions (or end positions for minus strand item) name2 Alternate name (e.g. gene\_id from GTF) cdsStartStat Status of CDS start annotation (none, unknown, incomplete, or complete) = ('none','unk','incmpl','cmpl') cdsEndStat Status of CDS end annotation (none, unknown, incomplete, or complete) exonFrames Exon frame 0,1,2, or -1 if no frame for exon (Related to codon. Number represents extra bases (modulus of 3) from previous exon block brought to a current exon block.) If cdsStart == cdsEnd, that means non-coding sequence.

• maybe cdsStartStat and cdsEndStat == "none" mean the same thing. maybe exonFrames == "-1," means the same thing.

#### Usage

```
generateGenicElementCoor(
  genepred,
  element.names = "all",
  upstream = NULL,
  downstream = NULL,
  genome.name = NULL,
  genome = NULL,
  return.coor.obj = FALSE
)
```

#### **Arguments**

genepred UCSC genome name (e.g., hg19, mm39). Types of genic elements to output: "all", "intron", "exon", "CDS", or "UTR". element.names Default is "all". upstream Length of upstream sequence (can overlap other genes). downstream Length of downstream sequence (can overlap other genes). genome.name UCSC genome name for trimming overflowing coordinates. Genome object for coordinate resolution. genome

return.coor.obj

Whether to return a Coordinate object (default: FALSE).

#### Value

Genic element coordinates in a data. table or Coordinate object.

generateIntergenicCoor

Resolve and generate genic element coordinates from UCSC genePred table.

#### **Description**

Function generates intergenic coordinates from a UCSC genePred table. It allows users to specify the genePred data source, the relative position and minimum length for intergenic regions, and whether to return the results as a Coordinate object or a data.table.

### Usage

```
generateIntergenicCoor(
 genepred,
  genome.name,
  igr.rel.pos = c(5000, 7500),
 igr.min.length = 150,
  return.coor.obj = FALSE
)
```

#### **Arguments**

genepred UCSC genePred table or database name ("refseq" or "gencode"). UCSC genome name (e.g., hg38, mm39). genome.name Intergenic relative position, defaults to c(5000, 7500). igr.rel.pos igr.min.length Minimum length for intergenic regions, default is 150. return.coor.obj

Return results as a Coordinate object? Default FALSE.

#### Value

Intergenic coordinates as a data. table or Coordinate object.

getCOSMICauthURL 27

getCOSMICauthURL	Get COSMIC authenticated URL.
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### **Description**

To access the data for non-commercial usage, you must register with the COSMIC. This function fetch the authenticated URL from the public URL given by the COSMIC website.

### Usage

```
getCOSMICauthURL(email, password, url)
```

### **Arguments**

email Email registered with COSMIC.

password Password associated with the registered email.

url Public URL provided by the COSMIC website for data access.

### Value

Authenticated URL valid for 1-hour access to COSMIC data.

```
{\tt getCOSMIC} cancer {\tt GeneCensus}
```

Get Cancer Gene Census (CGC) from COSMIC database.

### Description

To access the data for non-commercial usage, you must register with the COSMIC. This function fetch the latest CGC.

# Usage

```
getCOSMICcancerGeneCensus(email, password)
```

# Arguments

email Email registered with COSMIC.

password Password associated with the registered email.

### Value

A data. table containing the Cancer Gene Census data.

getCOSMIClatestVersion

Function retrieves the latest version information of the COSMIC database and the associated genome version by scraping data from the COSMIC website.

### Description

Function retrieves the latest version information of the COSMIC database and the associated genome version by scraping data from the COSMIC website.

### Usage

```
getCOSMIClatestVersion()
```

#### Value

A named vector containing the latest COSMIC version (cosmic) and genome version (genome).

getCOSMICmutantExport Function downloads the latest Cosmic Mutant Export data from the COSMIC database. It requires the user to be registered with COSMIC for non-commercial use. The function constructs the URL for the latest mutant export file, authenticates the URL, and then downloads the data.

### **Description**

Function downloads the latest Cosmic Mutant Export data from the COSMIC database. It requires the user to be registered with COSMIC for non-commercial use. The function constructs the URL for the latest mutant export file, authenticates the URL, and then downloads the data.

# Usage

```
getCOSMICmutantExport(email, password)
```

# **Arguments**

Email registered with COSMIC for accessing data. email

Password for the COSMIC account. password

#### Value

A data. table containing the Cosmic Mutant Export data.

getEnsemblData 29

getEnsemblData	A generic function to get Ensembl data persistently from a URL. This is an internal function used by other getEnsemblXXX functions.

#### **Description**

Error is handled based on their rule as set out at https://github.com/Ensembl/ensembl-rest/wiki/HTTP-Response-Codes

### Usage

```
getEnsemblData(url, handle, max.attempt = 5)
```

#### **Arguments**

url Pre-built Ensembl REST API URL.

handle curl handle object configured for the Ensembl REST API.

max.attempt Maximum number of attempts to fetch the data, default is 5.

#### Value

Parsed JSON data, which could be in the form of a data.frame or a list of lists, depending on the API response.

```
getEnsemblRegionFeatures
```

Get features of a given region.

#### **Description**

Function fetches various genomic features for a specified region from the Ensembl database. It allows specifying the species, chromosome, region range, and types of features to query.

### Usage

```
getEnsemblRegionFeatures(species, chromosome, start, end, features)
```

### **Arguments**

species Species name or alias (e.g., homo\_sapiens, human).

chromosome Chromosome name in Ensembl format (without 'chr' prefix).

start Start position of the region. end End position of the region.

features List of region features to retrieve from Ensembl. Valid options include "band",

"gene", "transcript", "cds", "exon", "repeat", "simple", "misc", "variation", "somatic\_variation", "structural\_variation", "somatic\_structural\_variation", "constrained", "regulatory", "motif", "peak", "other\_regulatory", "array\_probe", "mane".

### Value

A data.table containing the requested Ensembl features.

```
getEnsemblVariantFeatures
```

Get features of given variant IDs.

### **Description**

Function retrieves features for given variant IDs from the Ensembl database. It handles requests asynchronously in batches due to server limits and includes options to fetch additional variant information. Error handling for different HTTP response statuses is implemented to manage request failures.

### Usage

```
getEnsemblVariantFeatures(
   species,
   variant.ids,
   include.genotypes = FALSE,
   include.phenotypes = FALSE,
   include.allele.frequencies = FALSE,
   include.genotype.frequencies = FALSE,
   curl.max.con = 100,
   verbose = 1
)
```

### **Arguments**

```
species
                 Species name or alias (e.g., homo_sapiens, human).
variant.ids
                 A vector of variant IDs (e.g., rs56116432, COSM476).
include.genotypes
                 Include genotypes in the response? Default FALSE.
include.phenotypes
                 Include phenotypes in the response? Default FALSE.
include.allele.frequencies
                 Include allele frequencies? Default FALSE.
include.genotype.frequencies
                 Include genotype frequencies? Default FALSE.
curl.max.con
                 Maximum number of concurrent connections for curl requests. Default is 100.
verbose
                  Verbosity level: 1 for error only, 2 for all requests. Default 1.
```

## Value

A variant-named list containing lists of variant features.

```
getEnsemblVariantFeatures_serial
```

Get features of given variant IDs.

#### **Description**

Function fetches variant features from the Ensembl database for a set of variant IDs. It handles variant IDs in batches to comply with server limits and can include additional information like genotypes, phenotypes, allele frequencies, and genotype frequencies.

### Usage

```
getEnsemblVariantFeatures_serial(
  species,
  variant.ids,
  include.genotypes = FALSE,
  include.phenotypes = FALSE,
  include.allele.frequencies = FALSE,
  include.genotype.frequencies = FALSE)
```

#### **Arguments**

```
species Species name or alias (e.g., homo_sapiens, human).

variant.ids A vector of variant IDs (e.g., rs56116432, COSM476).

include.genotypes

Include genotypes in the response? Default FALSE.

include.phenotypes

Include phenotypes in the response? Default FALSE.

include.allele.frequencies

Include allele frequencies? Default FALSE.

include.genotype.frequencies

Include genotype frequencies? Default FALSE.
```

#### Value

A list, named by variant IDs, containing lists of variant features.

getGnomADvariants

Get gnomAD VCF file using tabix.

# Description

Function retrieves variant data from gnomAD VCF files using tabix for a specified set of genomic regions. It allows users to select the gnomAD version and server location (Google, Amazon, or Microsoft) for fetching the data.

#### Usage

```
getGnomADvariants(
  chr.names,
  starts,
  ends,
  INFO.filter = NULL,
  version = "3.1.2",
  server = "random"
)
```

### Arguments

chr.names Chromosome names.

starts Start positions.
ends End positions.

INFO. filter Parse only filtered INFO ID. Default is to parse all IDs.

version The gnomAD version. Default to latest version 3.1.2.

server Server locations: "google", "amazon", or "microsoft". Default is random.

#### Value

A data.table of VCF.

```
getICTVvirusMetadataResource
```

Get Virus Metadata Resource (VMR) from International Committee on Taxonomy of Viruses (ICTV)

# Description

Always get the latest VMR table, so no argument.

### Usage

```
getICTVvirusMetadataResource()
```

### Value

Virus Metadata Resource data.table.

getNCBIassemblySummary

Get NCBI assembly summary.

### **Description**

Retrieves the assembly summary from NCBI for a specified taxonomic group. This function allows users to obtain genome assembly information from either RefSeq or GenBank databases for various taxonomic groups.

### Usage

```
getNCBIassemblySummary(organism.group, db = "refseq")
```

### **Arguments**

organism.group A string specifying the taxonomic group for which the assembly summary is requested. Options include 'archaea', 'bacteria', 'fungi', 'invertebrate', 'plant', 'protozoa', 'vertebrate\_mammalian', 'vertebrate\_other', 'viral', or 'all'.

db A string specifying the database to use, either 'refseq' or 'genbank'.

### Value

A data.table containing the assembly summary for the specified taxonomic group.

getNCBItaxonomy

Get all taxonomy classification from NCBI

# Description

NCBI provides taxonomy for all life domains in rankedlineage.dmp file. This function download and store the file in kmeRtone\_data path.

#### Usage

```
getNCBItaxonomy()
```

#### Value

A data.table of taxonomy classification.

getScores	Function calculates scores for k-mers based on case and control k-mer
	counts.

#### **Description**

Function calculates scores for k-mers based on case and control k-mer counts.

#### Usage

```
getScores(case.kmers, control.kmers)
```

#### **Arguments**

case.kmers A data.table containing k-mer counts in case samples. control.kmers A data.table containing k-mer counts in control samples.

#### Value

A data.table containing scores for each k-mer.

getUCSCgenePredTable Retrieve Gene Prediction Table from UCSC for a Given Genome

#### **Description**

This function retrieves the gene prediction table from the UCSC genome database for a specified genome. It can fetch data from either the RefSeq or GENCODE databases.

#### Usage

```
getUCSCgenePredTable(genome.name, db)
```

### **Arguments**

genome.name A string specifying the UCSC genome name for which the gene prediction table

is to be retrieved, e.g., 'hg38', 'mm39'.

db A string specifying the database used by UCSC to generate the table. Options

are 'refseq' or 'gencode'.

### Value

A data. table containing the gene prediction table from the specified UCSC genome and database.

#### **Examples**

```
## Not run:
    Example usage:
    Retrieve the RefSeq gene prediction table for the human genome (hg38)
    genePredTable <- getUCSCgenePredTable(genome.name = "hg38", db = "refseq")
## End(Not run)</pre>
```

getVCFmetainfo 35

getVCFmetainfo	Read VCF metainfo file using tabix.

### **Description**

Require tabix in PATH VCF manual is referred from https://samtools.github.io/hts-specs/VCFv4.3.pdf

#### Usage

```
getVCFmetainfo(vcf.file)
```

### **Arguments**

vcf.file

A path to a local or remote tabix-indexed VCF file.

#### Value

VCF metainfo.

GET\_OPTIMUM\_K

Function aims to determine the optimum k-mer size for genomic analysis based on the provided case data and genome. It may involve analyzing central patterns within the genomic sequences.

### Description

Function aims to determine the optimum k-mer size for genomic analysis based on the provided case data and genome. It may involve analyzing central patterns within the genomic sequences.

# Usage

```
GET_OPTIMUM_K(case, genome, central.pattern)
```

# **Arguments**

case Coordinate class object or similar structure for case data.

genome Genome class object or similar structure.

central.pattern

Central pattern of the k-mer.

#### Value

Optimum k-mer size for the given genomic data.

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initKmerTable

Initialise k-mer table with all possible k-mers

### **Description**

Initialise k-mer table with the following columns: kmer, pos\_strand, and neg\_strand

#### Usage

```
initKmerTable(k, central.pattern = NULL, split.kmer = FALSE)
```

#### **Arguments**

k K-mer size. Limit to 15 because vector size is limited to .Machine\$integer.max. For 9- to 15-mer, the kmer sequence is separated to two columns (kmer\_part1 and kmer\_part2) to reduce memory significantly. central.pattern

Central pattern(s) of the k-mer. Default is NULL.

split.kmer

Whether to split the k-mer sequence into two parts for large k values. Default is FALSE.

#### Value

data.table with 3 columns: kmer, pos\_strand, neg\_strand

kmeRtone

kmeRtone all-in-one user interface

### Description

This function serves as an all-in-one interface for various genomic data analyses leveraging k-mer based techniques.

### Usage

```
kmeRtone(
  case.coor.path,
  genome.name,
  strand.sensitive,
  ctrl.rel.pos = c(80, 500),
  case.pattern,
  output.dir = "output/",
  case,
  genome,
  control,
  control.path,
  genome.path,
  rm.case.kmer.overlaps,
```

kmeRtone 37

```
single.case.len,
merge.replicates,
kmer.table,
module = "score",
rm.dup = TRUE,
case.coor.1st.idx = 1,
ctrl.coor.1st.idx = 1,
coor.load.limit = 1,
genome.load.limit = 1,
genome.fasta.style = "UCSC",
genome.ncbi.db = "refseq",
use.UCSC.chr.name = FALSE,
verbose = TRUE,
kmer.cutoff = 5,
selected.extremophiles,
other.extremophiles,
cosmic.username,
cosmic.password,
tumour.type.regex = NULL,
tumour.type.exact = NULL,
cell.type = "somatic",
genic.elements.counts.dt,
population.size = 1e+06,
selected.genes,
add.to.existing.population = FALSE,
population.snv.dt = NULL,
pop.plot = TRUE,
pop.loop.chr = FALSE
```

#### **Arguments**

)

case.coor.path Path to a folder containing chromosome-separated coordinate files or bedfiles.

Assumed replicates for subfolder or bedfiles.

 $\label{eq:condition} \mbox{genome.name} \quad \mbox{Name of the genome (e.g., "hg19", "hg38"). Default is "unknown". strand.sensitive$ 

Logical value indicating whether strand polarity matters. Default is TRUE.

k Length of k-mer to be investigated. Recommended values are 7 or 8.

ctrl.rel.pos A vector of two integers specifying the relative range positions of control re-

gions.

case.pattern Regular expression pattern for identifying case regions. Default is NULL.

output.dir Directory path for saving output files. Default is "output/".

case Optional pre-built Coordinate object.
genome Optional pre-built Genome object.

control Optional pre-built control Coordinate object.
control.path Path for pre-built control Coordinate object.

genome.path Path to a directory of user-provided genome FASTA files.

rm.case.kmer.overlaps

Logical indicating whether to remove overlapping k-mers in case regions. Default is FALSE.

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single.case.len

Integer indicating uniform length of case regions.

merge.replicates

Logical indicating whether to merge replicates. Default is TRUE.

kmer.table Pre-calculated k-mer score table.

module Selected kmeRtone module to run. Possible values include "score", "explore",

"tune", among others.

rm. dup Logical indicating whether to remove duplicate coordinates. Default is TRUE.

case.coor.1st.idx

Integer specifying indexing format for case coordinates.

ctrl.coor.1st.idx

Integer specifying indexing format for control coordinates.

coor.load.limit

Maximum number of coordinates to load. Default is 1.

genome.load.limit

Maximum number of genome sequences to load. Default is 1.

genome.fasta.style

String specifying the style of the genome FASTA. Possible values are "UCSC", "NCBI". Default is "UCSC".

genome.ncbi.db String specifying the NCBI database to use. Possible values are "refseq", "genbank". Default is "refseq".

use.UCSC.chr.name

Logical indicating whether to use UCSC chromosome names.

verbose Logical indicating whether to display progress messages. Default is TRUE.

 $\label{lem:cutoff} \text{Cutoff percentage for $k$-mer selection in case studies. Default is 5.}$ 

selected.extremophiles

Vector of selected extremophile species for study.

other.extremophiles

Vector of other extremophile species for control.

cosmic.username

COSMIC username for accessing the cancer gene census.

cosmic.password

COSMIC password for accessing the cancer gene census.

tumour.type.regex

Regular expression pattern for filtering tumour types.

tumour.type.exact

Exact tumour type to be included in the cancer gene census.

cell. type Cell type to be included in the cancer gene census. Default is "somatic".

genic.elements.counts.dt

Data table of susceptible k-mer counts in genic elements.

population.size

Size of the population for cross-population studies. Default is 1 million.

selected genes Selected genes for mutation in cross-population studies.

add.to.existing.population

Logical indicating whether to add to the existing simulated population. Default is FALSE.

Kmer\_Table 39

population.snv.dt

Data table of single nucleotide variants used in population simulations.

pop.plot Logical indicating whether to plot the outcome of the cross-population study.

Default is TRUE.

pop. loop. chr Logical indicating whether to loop based on chromosome name in cross-population

studies. Default is FALSE.

#### Value

Depends on the selected module.

Kmer\_Table

A R6 class wrapper for data.table

### **Description**

A R6 class wrapper for data.table

A R6 class wrapper for data.table

#### **Details**

A way to grow data.table in different environment but still retaining access to it. A temporary fix until data.table developer develop update row by reference.

#### **Public fields**

DT data.table of k-mers

### Methods

### **Public methods:**

```
• Kmer_Table$new()
```

- Kmer\_Table\$print()
- Kmer\_Table\$remove\_N()
- Kmer\_Table\$filter\_central\_pattern()
- Kmer\_Table\$update\_count()
- Kmer\_Table\$update\_row()
- Kmer\_Table\$clone()

Method new(): initialize empty data.table of k-mers

Usage:

Kmer\_Table\$new()

Method print(): Print method.

Usage:

Kmer\_Table\$print()

Method remove\_N(): Remove unknown base N.

Usage:

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```
Kmer_Table$remove_N()
Method filter_central_pattern(): Filter out k-mers without defined central patterns.
 Usage:
 Kmer_Table$filter_central_pattern(central.pattern, k)
 Arguments:
 central.pattern Central pattern.
 k Length of k-mer.
 Returns: None.
Method update_count(): Update count for existed k-mers in the table.
 Kmer_Table$update_count(kmers, is.strand.sensitive, strand)
 Arguments:
 kmers K-mer table with new count to be added to the main table.
 is.strand.sensitive Does strand polarity matter?
 strand If yes, what is the strand refers to? "+" or "-".
 Returns: None.
Method update_row(): Add new rows for new k-mers with their respective counts that is not
existed yet in the main table.
 Usage:
 Kmer_Table$update_row(kmers, is.strand.sensitive, strand)
 kmers K-mer table with new k-mers to be added to the main table.
 is.strand.sensitive Does strand polarity matter?
 strand If yes, what is the strand refers to? "+" or "-".
 Returns: None.
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 Kmer_Table$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

 ${\tt loadCoordinate}$ 

Build Coordinate object.

# Description

The Coordinate object is capable of loading genomic coordinates on demand. Chromosome-specific coordinates can be called in a bracket. The coordinates can also be expanded to k-mer size equally on both flanks

loadGenome 41

#### Usage

```
loadCoordinate(
  root.path = NULL,
  single.len = NULL,
  is.strand.sensitive = TRUE,
  merge.replicates = TRUE,
  rm.dup = TRUE,
  add.col.rep = FALSE,
  is.kmer = FALSE,
  k = NA,
  ori.first.index = 1,
  load.limit = 1
)
```

### **Arguments**

root.path A path to a directory containing either: (1) chromosome-separated coordinate

files (multiple replicates is assumed for sub-folder) or (2) bedfile (multiple repli-

cates is assumed for separate bedfiles).

single.len Single case length relevant when all coordinates have the same length. This is

for memory optimization. Default is NULL.

is.strand.sensitive

A boolean whether strand polarity matters. Default is TRUE.

merge.replicates

Merge coordinate from different replicates. Default is TRUE. If not merging, duplicates will give weight to the k-mer counting. If add.col.rep, merged coordinates will give weight to the k-mer counting.

dinate will contain column replicate e.g. "rep1&rep2".

rm. dup Remove duplicates in each replicate. Default is TRUE.

add.col.rep Add column replicate to the coordinate table.

is.kmer Is the coordinate refers to k-mer i.e. expanded case? Default is FALSE.

k Length of k-mer relevant only when is.kmer is TRUE.

ori.first.index

Indexing format of the coordinate: 0 for zero-based (start, end) and 1 for one-

based (start, end). Default is 1.

load.limit Maximum number of coordinate data.table loaded on RAM. Default is 1.

### Value

Coordinate object.

loadGenome

Build Genome object.

# Description

The Genome object is capable of loading chromosome sequence on demand. UCSC Genomes are included in this kmeRtone package. Their specific chromosome sequence will be downloaded on demand once.

42 loadGenomicContents

#### Usage

```
loadGenome(
  genome.name,
  fasta.style,
  mask = "none",
  fasta.path,
  ncbi.db,
  ncbi.asm,
  use.UCSC.name = FALSE,
  load.limit = 1
)
```

### **Arguments**

A genome name. UCSC and NCBI genome is included with kmeRtone. Input genome.name their name e.g. hg19 or GRCh37. FASTA version: "UCSC" or "NCBI". fasta.style Genome mask: "none", "soft", or "hard". Default is "none". mask fasta.path Path to the fasta file as a character vector. ncbi.db NCBI database: "refseq" or "genbank". ncbi.asm NCBI assembly table. use.UCSC.name For NCBI Genome, use UCSC-style chromosome name? Default is FALSE. load.limit Maximum chromosome sequences loaded. Default is 1.

### Value

A UCSC\_Genome or NCBI\_Genome object.

### **Examples**

```
## Not run:
    genome <- loadGenome(...)
    genome["chr1"]
    genome[c("chr1", "chr2")]
## End(Not run)</pre>
```

 ${\tt loadGenomicContents}$ 

Function calculates various genomic content metrics based on the provided genome object.

# Description

Function calculates various genomic content metrics based on the provided genome object.

### Usage

loadGenomicContents(genome)

mapKmers 43

### **Arguments**

genome An object of class 'NCBI\_Genome' containing genomic information.

# Value

A data.table containing calculated genomic content metrics.

mapKmers	Map k-mers of a given sequence and coordinate	

# Description

This function maps k-mers within a specified sequence based on provided start and end coordinates, or based on a fixed length.

# Usage

```
mapKmers(seq, start, end = NULL, len = NULL, k, rm.trunc.kmer = TRUE)
```

# Arguments

seq	A single sequence string in which k-mers are to be mapped.
start	A vector of start coordinates for mapping k-mers. If only start positions are provided, exact k-mer extraction is performed.
end	A vector of end coordinates corresponding to the start positions. If NULL, all regions are assumed to have the same length. Used for varied region lengths to perform a sliding window.
len	An integer specifying the fixed length of regions. Used when regions have a uniform length greater than k. End coordinates are assumed NULL in this case.
k	An integer specifying the length of k-mers to be mapped.
rm.trunc.kmer	Logical indicating whether to remove truncated k-mers resulting from out-of-bound regions. Default is TRUE.

# Value

A vector of mapped k-mers.

44 mixColors

 ${\tt mergeCoordinate}$ 

Merge overlapping or continuous regions.

# Description

Table must have start and end columns. The output is exactly similar to the reduce function from GenomicRanges.

# Usage

```
mergeCoordinate(coor)
```

# Arguments

coor

Coordinate data. table.

### Value

Merged coordinate data.table.

 ${\tt mixColors}$ 

Mix color

# Description

This is useful to get overlayed colors.

# Usage

```
mixColors(cols, alpha)
```

# Arguments

cols Colors in hex format or R color code e.g. "red", "black", etc.

alpha Add alpha transparency value.

# Value

New mixed colors in hex format.

NCBI\_Genome 45

NCBI\_Genome

Class constructor - build NCBI Genome object

### **Description**

Class constructor - build NCBI Genome object Class constructor - build NCBI Genome object

#### **Details**

NCBI FASTA file contain nucleotide accession number at the headers, followed by some information about the sequence whether they are chromosome, plasmid, or mictochondria, their assembly status, etc.

#### **Public fields**

```
fasta_file A path to FASTA file. fasta files.

genome_name A genome name.

db NCBI database: "refseq" or "genbank"

seq A chromosome-named list of sequences.

seq_len A chromosome-named vector of sequence length.

load_limit Maximum chromosome sequences loaded.

mask Genome mask status: "hard", "soft", or "none".

use_UCSC_name Use UCSC style chromosome name? Default to FALSE.

headers A chromosome-named vector of headers.

avail_seqs Available chromosome sequences in the fasta file.

asm Assembly summary.
```

### Methods

### **Public methods:**

```
NCBI_Genome$new()NCBI_Genome$[()NCBI_Genome$print()NCBI_Genome$get_assembly_report()NCBI_Genome$clone()
```

### Method new(): Create a new NCBI Genome class

```
Usage:
NCBI_Genome$new(
  genome.name,
  db,
  fasta.file,
  asm,
  mask,
  use.UCSC.name,
  load.limit
```

46 partitionCoordinate

```
Arguments:
 genome.name A genome name. NCBI genome is included with kmeRtone.
 db NCBI database: "refseq" or "genbank".
 fasta.file A path to the NCBI-style fasta files. This is for user's own FASTA file.
 asm NCBI assembly summary.
 mask Genome mask status: "hard", "soft", or "none". Default is "none".
 use.UCSC.name Use UCSC style chromosome name? Default to FALSE.
 load.limit Maximum chromosome sequences loaded. Default is 1.
 Returns: A new NCBI Genome object.
Method [(): Calling chromosome sequence by loading on demand. Maximum load is determine
by load_limit field.
 Usage:
 NCBI_Genome$[(chr.names, reload = FALSE)
 Arguments:
 chr. names Chromosome name. It can be a vector of chromosomes.
 reload Reload the sequence from the fasta_file. Default is FALSE.
 Returns: A single or list of sequence of requested chromosome.
Method print(): Print summary of Genome object.
 Usage:
 NCBI_Genome$print()
 Returns: Message of Genome object summary.
Method get_assembly_report(): Get NCBI assembly report for the genome.
 Usage:
 NCBI_Genome$get_assembly_report()
 Returns: Message of Genome object summary.
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 NCBI_Genome$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

 $partition {\tt Coordinate} \qquad \textit{Partition overlapping or continuous regions}.$ 

### **Description**

Table must have start and end columns. The mechanism is similar to the disjoin function from GenomicRanges but the end coordinate is different.

### Usage

```
partitionCoordinate(coor)
```

persistentDownload 47

### **Arguments**

coor Coordinate data.table.

### Value

Partitioned coordinate data. table.

persistentDownload

Download file until successful

# Description

If download failed, it will be repeated until max attempt reached.

### Usage

```
persistentDownload(
  file.url,
  output.name,
  max.attempt = 5,
  user.invoke = TRUE,
  header
)
```

# Arguments

file.url File uniform resource locator.

output.name Output name.

max.attempt Maximum number of attempt. Default is 5.

user.invoke If number of attempt is reached, ask user whether to keep trying. Default is

TRUE to invoke the prompt.

header A named list or vector of curl header.

### Value

A downloaded file.

48 readBED

readBCF

Read BCF files and extract chromosome information

# Description

This function reads BCF (Binary Variant Call Format) files and extracts chromosome information.

### Usage

```
readBCF(fname, regions, is_file)
```

### **Arguments**

fname The file name or URL of the BCF file.

regions A string specifying the regions to be read from the BCF file.

is\_file Boolean indicating if the regions parameter is a file.

### Value

A list containing chromosome information.

readBED

Read a BED file. One-based indexing is enforced.

# Description

Read a BED file. One-based indexing is enforced.

# Usage

```
readBED(bed.path)
```

# Arguments

bed.path

A path to a BED file.

### Value

data.table.

readFASTA 49

readFASTA

Read FASTA files.

# Description

Read FASTA files.

# Usage

```
readFASTA(fasta.file)
```

# Arguments

fasta.file A path to a FASTA file.

# Value

A sequence vector with header names

readSingleFASTA

Read fast single-header FASTA file.

# Description

Read fast single-header FASTA file.

# Usage

```
readSingleFASTA(file_path, mask = "none")
```

# Arguments

file\_path A path to FASTA file.

mask Capitalize all base letters?

# Value

A single sequence string without header.

50 readVCF2

	D INCECT : . I.	
readVCF	Read VCF file using tabix.	

### **Description**

Require tabix in PATH VCF manual is referred from https://samtools.github.io/hts-specs/VCFv4.3.pdf

#### Usage

```
readVCF(vcf.file, chr.names, starts, ends, INFO.filter = NULL)
```

### **Arguments**

vcf.file A path to a local or remote tabix-indexed VCF file.

chr.names Chromosome names.

starts Start positions. ends End positions.

INFO.filter Parse only filtered INFO ID. Default is to parse all IDs.

### Value

A data.table of VCF.

readVCF2	Read VCF file using tabix.	

### **Description**

Require tabix in PATH VCF manual is referred from https://samtools.github.io/hts-specs/VCFv4.3.pdf

### Usage

```
readVCF2(vcf.file, chr.names, starts, ends, INFO.filter = NULL)
```

### **Arguments**

vcf. file A path to a local or remote tabix-indexed VCF file.

chr.names Chromosome names.
starts Start positions.
ends End positions.

INFO. filter Parse only filtered INFO ID. Default is to parse all IDs.

### Value

A data.table of VCF.

removeRegion 51

removeRegion	Remove overlapping region in coordinate data.table.
--------------	---

### **Description**

Any "coor" that overlap within the "region" will be removed e.g. region = 10-20 and coor = 1-30 The results will be: coor = 1-10, 20-30 The coor still overlap one base at the terminal. This is done to produce exact result as the previous MPhil research.

### Usage

```
removeRegion(coor, region)
```

#### **Arguments**

coor Coordinate data. table.

region A data. table of region coordinate to be removed.

### Value

New coordinate data. table with the regions removed.

reverseComplement Get reverse complement sequence of DNA

### Description

Get reverse complement sequence of DNA

### Usage

```
reverseComplement(DNA.sequence, form = "string")
```

### **Arguments**

DNA sequence an be in a form of character vector or string. Multiple sequences

are accepted.

form Specify the form: "string" of "vector". Default is "string"

#### Value

Reverse complementary sequence

#### **Examples**

```
## Not run:
    reverseComplement("AAAAA")
    reverseComplement(c("AAAAA", "CCCCC"))
    reverseComplement(c("A", "A", "A", "A"), form = "vector")
## End(Not run)
```

52 **SCORE** 

**SCORE** 

Calculate susceptibility scores for k-mers in case and control regions.

### **Description**

Function calculates susceptibility scores for k-mers in case and control regions. Case regions are defined by genomic coordinates provided in a file or data.table. Control regions can be constructed relative to the case regions or provided directly. The scores are computed based on the occurrence of k-mers in case and control regions.

# Usage

```
SCORE(
  case.coor.path,
  genome.name,
  strand.sensitive,
 k,
 ctrl.rel.pos,
  case.pattern,
 output.path,
  case,
  genome,
  control,
  control.path,
  genome.path,
  rm.case.kmer.overlaps,
  single.case.len,
 merge.replicates,
 rm.dup,
  case.coor.1st.idx,
  ctrl.coor.1st.idx,
  coor.load.limit,
  genome.load.limit,
  genome.fasta.style,
  genome.ncbi.db,
 use.UCSC.chr.name,
  verbose
)
```

# **Arguments**

case.pattern

```
case.coor.path Path to the file containing genomic coordinates of case regions.
genome.name
                  Name of the genome to be used.
strand.sensitive
                  Logical indicating whether strand information should be considered.
                  Length of the k-mers to be extracted.
ctrl.rel.pos
                  Relative positions of control regions with respect to case regions. It should be a
                  vector of two integers indicating the upstream and downstream distances from
                  the case regions.
                  Regular expression pattern to identify the central sequence in case regions.
```

scoreKmers 53

output.path Directory path where the output files will be saved.

case Data.table containing the genomic coordinates of case regions.
genome Genome data.table containing the genomic sequence information.

control Data.table containing the genomic coordinates of control regions.

control.path Path to the file containing genomic coordinates of control regions (optional).

genome.path Path to the genome FASTA file.

rm.case.kmer.overlaps

Logical indicating whether overlapping k-mers within case regions should be

removed.

single.case.len

Single case length.

merge.replicates

Logical indicating whether replicates should be merged.

rm. dup

Logical indicating whether duplicate k-mers should be removed.

case.coor.1st.idx

First index in the case coordinate file.

ctrl.coor.1st.idx

First index in the control coordinate file.

coor.load.limit

Maximum number of coordinates to load.

genome.load.limit

Maximum number of genome sequences to load.

genome.fasta.style

FASTA style.

genome.ncbi.db NCBI database.

use.UCSC.chr.name

Logical indicating whether to use UCSC chromosome names.

verbose Logical indicating whether to display progress messages.

### Value

Data.table containing susceptibility scores for k-mers.

scoreKmers	Function calculates the Z-score for each k-mer based on the observed
	case counts and expected case counts under the null hypothesis.

# Description

Function calculates the Z-score for each k-mer based on the observed case counts and expected case counts under the null hypothesis.

### Usage

```
scoreKmers(kmer.table)
```

### **Arguments**

kmer.table

A data.table containing k-mer counts, where each row represents a k-mer and columns "case" and "control" represent the counts in case and control samples respectively.

#### Value

A modified version of the input kmer. table with an additional column "z" containing the calculated Z-scores for each k-mer.

 ${\tt selectGenomesForCrossSpeciesStudy}$ 

Select genomes for cross-species studies.

### **Description**

The following filters are applied:

- 1. assembly\_level: "Complete Genome" or "Chromosome"
- 2. genome\_rep: "Full"
- 3. Unique species\_taxid (single representative species)
- 4. refseq\_category of "reference genome" is prioritised over "representative genome"

# Usage

```
selectGenomesForCrossSpeciesStudy(organism.group = "bacteria", db = "refseq")
```

# Arguments

organism.group Species group: archaea, bacteria, fungi, invertebrate, plant, protozoa, vertebrate\_mammalian, vertebrate\_other, or viral.

db Database record to use: refseq or genbank

# Value

NCBI assembly summary with added column organism.group.

```
{\tt selectRepresentativeFromASM}
```

Select the best representative species from the NCBI assembly summary.

# Description

sort.idx is a weight to sort where heavier will be preffered. Any tie weight will be further sorted by organism\_name. Only the top unique species\_taxid will be retained in the final assembly summary.

# Usage

```
selectRepresentativeFromASM(asm)
```

### **Arguments**

asm

NCBI assembly summary.

### Value

Trimmed NCBI assembly summary.

 ${\it simulate Population}$ 

Simulate a population given ranges of chromosome sequence to mutate.

# Description

Simulate a population given ranges of chromosome sequence to mutate.

# Usage

```
simulatePopulation(
  chrom_seq,
  starts,
  ends,
  strand,
  snv_df,
  pop_size,
  top_kmers,
  central_pattern,
  k
)
```

56 splitFASTA

### **Arguments**

chrom\_seq A chromosome sequence.

starts Start positions.
ends End positions.

strand Strand type: "+" or "-".

snv\_df A table of SNV frequency. Columns: position, base, count.

pop\_size Size of population.

central\_pattern

K-mer central pattern.

k K-mer size.

#### Value

A count matrix with 4 rows for total top k-mers and susceptible k-mers in sense and antisense. Columns correspond to population individuals.

splitFASTA Split a FASTA file by header.

# Description

The first non-space word in the header will be used as the file name.

### Usage

```
splitFASTA(fasta.file, output.dir = "./")
```

### **Arguments**

fasta.file A path to a FASTA file.

output.dir A path to save the output results. Default is current working directory.

### **Details**

data.table::fread is not built for reading in chunks. The developers expect skip and nrow arguments to be in a small number. Large number slows the reading a bit.

### Value

A splitted fasta files by its headers.

splitFASTA2 57

splitFASTA2

Read fast single-header FASTA file.

### **Description**

Slower than data.table::fread

# Usage

```
splitFASTA2(file_path, output_dir)
```

# **Arguments**

```
file_path A path to FASTA file.

output_dir A path to save the output results.
```

#### Value

A single sequence string without header.

```
STUDY_ACROSS_POPULATIONS
```

Study k-mer composition of selected COSMIC causal cancer genes across human populations worldwide.

# Description

Simulation of human population is based on single nucleotide variantion.

# Usage

```
STUDY_ACROSS_POPULATIONS(
   kmer.table,
   kmer.cutoff = 5,
   genome.name,
   k,
   db = "refseq",
   central.pattern = NULL,
   population.size = 1e+06,
   selected.genes,
   add.to.existing.population = FALSE,
   output.dir = "study_across_populations/",
   population.snv.dt = NULL,
   loop.chr = TRUE,
   plot = FALSE
)
```

#### **Arguments**

kmer.table A data.table of kmer table.

kmer.cutoff Percentage of extreme kmers to study. Default to 5.

genome.name UCSC genome name.

k K-mer size.

db Database used by UCSC to generate gene prediction: "refseq" or "gencode".

Default is "refseq".

central.pattern

K-mer's central patterns. Default is NULL.

population.size

Size of population to simulate. Default is 1 million.

selected.genes Set of genes to study e.g. skin cancer genes.

add.to.existing.population

Add counts to counts.csv? Default is FALSE.

output.dir A directory for the outputs. Default to study\_across\_populations.

population.snv.dt

Population SNV table.

loop.chr Loop chromosome?. Default is TRUE. If FALSE, beware of a memory spike

because of VCF content. VCF contains zero counts for every population. Input

pre-computed trimmed-version population.snv.dt.

plot Boolean. Default is FALSE. If TRUE, will plot results.

#### Value

An output directory containing plots.

STUDY\_ACROSS\_SPECIES Study k-mer composition across species.

# Description

Analysis of distribution of highly enriched k-mers across species.

### Usage

```
STUDY_ACROSS_SPECIES(
  kmer.table,
  kmer.cutoff = 5,
  k,
  central.pattern = NULL,
  selected.extremophiles,
  other.extremophiles,
  output.dir = "study_across_species/"
)
```

### **Arguments**

kmer.table A data.table of kmer table or path to it.

kmer.cutoff Percentage of extreme kmers to study. Default to 5 percent.

k K-mer size.

central.pattern

K-mer's central patterns. Default is NULL.

selected.extremophiles

A vector of selected extremophile species. e.g. c("Deinococcus soli", "Deinococcus deserti") The best representative will be selected from the assembly sum-

mary.

other.extremophiles

A vector of other extremophile species. These are used as a control to compare

with the selected extremophiles.

output.dir A directory for the outputs.

#### Value

An output directory containing plots.

STUDY\_CANCER\_GENES

Study k-mer composition of causal cancer genes from COSMIC Cancer Gene Census (CGC) database.

### Description

Detail of Cancer Gene Census can be accessed and read at https://cancer.sanger.ac.uk/census

### Usage

```
STUDY_CANCER_GENES(
  cosmic.username,
  cosmic.password,
  tumour.type.regex = NULL,
  tumour.type.exact = NULL,
  cell.type = "somatic",
  genic.elements.counts.dt,
  output.dir = "study_cancer_genes/"
)
```

### **Arguments**

```
cosmic.username
```

COSMIC username i.e. registered email.

cosmic.password

COSMIC password.

tumour.type.regex

Regular expression for "Tumour Types" column in Cancer Gene Census table. Default is NULL.

```
tumour.type.exact
```

Exact keywords for "Tumour Types" column in Cancer Gene Census table. De-

fault is NULL.

cell.type Type of cell: "somatic" or "germline". Default is "somatic".

genic.elements.counts.dt

Genic element count table generated from STUDY\_GENIC\_ELEMENTS.

output.dir A directory for the outputs.

### Value

An output directory containing plots.

STUDY\_GENIC\_ELEMENTS Study k-mer composition across species.

### **Description**

Study k-mer composition across species.

# Usage

```
STUDY_GENIC_ELEMENTS(
   kmer.table,
   kmer.cutoff = 5,
   k,
   genome.name = "hg38",
   central.pattern = NULL,
   db = "refseq",
   output.dir = "study_genic_elements/")
```

# **Arguments**

kmer.table A data.table of kmer table.

kmer.cutoff Percentage of extreme kmers to study. Default to 5.

k K-mer size.

genome.name UCSC genome name.

central.pattern

K-mer's central patterns. Default is NULL.

db Database used by UCSC to generate gene prediction: "refseq" or "gencode".

Default is "refseq".

 ${\tt output.dir}$  A directory for the outputs.

### Value

An output directory containing plots.

system3 61

system3

A system2 wrapper. If anything happen, just give me error!

### **Description**

Turn warning to error.

#### Usage

```
system3(
  command,
  args = character(),
  stdout = "",
  stderr = "",
  stdin = "",
  input = NULL,
  env = character(),
  wait = TRUE,
  minimized,
  invisible,
  timeout = 0
)
```

### **Arguments**

command the system command to be invoked, as a character string.

args a character vector of arguments to command.

stdout, stderr where output to 'stdout' or 'stderr' should be sent. Possible values are "", to

the R console (the default), NULL or FALSE (discard output), TRUE (capture the

output in a character vector) or a character string naming a file.

stdin should input be diverted? "" means the default, alternatively a character string

naming a file. Ignored if input is supplied.

input if a character vector is supplied, this is copied one string per line to a temporary

file, and the standard input of command is redirected to the file.

env character vector of name=value strings to set environment variables.

wait a logical (not NA) indicating whether the R interpreter should wait for the com-

mand to finish, or run it asynchronously. This will be ignored (and the interpreter will always wait) if stdout = TRUE or stderr = TRUE. When running the command asynchronously, no output will be displayed on the Rgui console in Windows (it

will be dropped, instead).

minimized, invisible

arguments that are accepted on Windows but ignored on this platform, with a

warning.

timeout in seconds, ignored if 0. This is a limit for the elapsed time running

command in a separate process. Fractions of seconds are ignored.

62 UCSC\_Genome

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Trim out-of-bound coordinates

# Description

It operates in two mode: coordinate table with and without chromosome. The former require Genome for the chromosomal sequence length.

### Usage

```
trimCoordinate(coor, seq.len, genome)
```

### **Arguments**

coor Coordinate data. table.

seq. len Sequence length to trim end position.

genome Genome class object.

#### Value

Trimmed coordinate data. table.

UCSC\_Genome

Class constructor - build Genome object

# Description

Class constructor - build Genome object

Class constructor - build Genome object

# **Public fields**

root\_path A path to a directory containing chromosome-separated fasta files.

genome\_name A genome name.

paths Individual chromosome sequence files.

seq A chromosome-named list of sequences.

seq\_len A chromosome-named vector of sequence length.

load\_limit Maximum chromosome sequences loaded.

mask Genome mask status: "hard", "soft", or "none".

info\_file Path to info file with pre-computed values.

chr\_names Chromosome names.

UCSC\_Genome 63

#### Methods

```
Public methods:
```

```
• UCSC_Genome$new()
```

- UCSC\_Genome\$[()
- UCSC\_Genome\$print()
- UCSC\_Genome\$get\_length()
- UCSC\_Genome\$get\_content()
- UCSC\_Genome\$clone()

### Method new(): Create a new Genome class

```
Usage:
```

UCSC\_Genome\$new(genome.name, root.path, mask, load.limit)

Arguments:

genome.name A genome name. UCSC genome is included with kmeRtone.

root.path A path to a directory containing chromosome-separated fasta files.

mask Genome mask status: "hard", "soft", or "none". Default is "none".

load.limit Maximum chromosome sequences loaded. Default is 1.

Returns: A new Genome object.

**Method** [(): Calling chromosome sequence by loading on demand. Maximum load is determine by load\_limit field.

```
Usage:
```

UCSC\_Genome\$[(chr.names, reload = FALSE)

Arguments:

chr. names Chromosome name. It can be a vector of chromosomes.

reload Reload the sequence from the root\_path. Default is FALSE.

Returns: A single or list of sequence of requested chromosome.

Method print(): Print summary of Genome object.

Usage:

UCSC\_Genome\$print()

Returns: Message of Genome object summary.

**Method** get\_length(): Get chromosome length from pre-calculated length

Usage:

UCSC\_Genome\$get\_length(chr.names, recalculate = FALSE)

Arguments:

chr. names Chromosome name. It can be a vector of chromosomes.

recalculate Recalculate the pre-calculated length.

Returns: A chromosome-named vector of length value.

**Method** get\_content(): Get pre-calculated sequence content e.g. G+C content

Usage:

UCSC\_Genome\$get\_content(chr.names, seq, recalculate = FALSE)

Arguments:

64 writeFASTA

chr. names Chromosome name. It can be a vector of chromosomes.

seq Sequence to count. e.g. c("G", "C")

recalculate Recalculate the pre-calculated length.

Returns: A chromosome-named vector of sequence content.

**Method** clone(): The objects of this class are cloneable with this method.

Usage:

UCSC\_Genome\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

writeBED

Write a BED file. Zero-based indexing is enforced.

## **Description**

Write a BED file. Zero-based indexing is enforced.

### Usage

```
writeBED(bed, output.filename)
```

### **Arguments**

bed A BED data.table.

 $\verb"output.filename"$ 

An output BED filename.

writeFASTA

Write FASTA files.

### **Description**

Write FASTA files.

### Usage

```
writeFASTA(seqs, fasta.path, append = FALSE)
```

### **Arguments**

seqs A vector or list of sequences with header name. If it is a list, it must only contain

one single sequence string for every element e.g. list(chr1 = "NNNNNNN")

not list(chr1 = c("NNNNNN", "AAAAAA"))

fasta.path A path to a FASTA file.

append Boolean. Default is FALSE. If TRUE, will append the results to existing file.

### Value

None

writeVCF 65

writeVCF	Write VCF file and compress using bgzip.

# Description

Require bgzip in PATH VCF manual is referred from https://samtools.github.io/hts-specs/VCFv4.3.pdf

# Usage

```
writeVCF(vcf, output.vcf.gz, append = FALSE, tabix = FALSE)
```

# **Arguments**

vcf A VCF object.

output.vcf.gz Output filename including vcf.gz extension.

append To append or not? Default is FALSE.

tabix To tabix or not? Default is FALSE.

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