

Package ‘uncoverappLib’

April 10, 2022

Title Interactive graphical application for clinical assessment of sequence coverage at the base-pair level

Version 1.4.0

Imports markdown, shiny, shinyjs, shinyBS,
shinyWidgets,shinycssloaders, DT, Gviz, Homo.sapiens, openxlsx,
condformat, stringr, org.Hs.eg.db,
TxDb.Hsapiens.UCSC.hg38.knownGene, BiocFileCache,rappdirs,
TxDb.Hsapiens.UCSC.hg19.knownGene, rlist, utils,
EnsDb.Hsapiens.v75, EnsDb.Hsapiens.v86, OrganismDbi,
BSgenome.Hsapiens.UCSC.hg19, processx, Rsamtools, GenomicRanges

Description

a Shiny application containing a suite of graphical and statistical tools to support clinical assessment of low coverage regions.It displays three web pages each providing a different analysis module: Coverage analysis, calculate AF by allele frequency app and binomial distribution.

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Encoding UTF-8

LazyData true

RoxygenNote 7.1.0

URL <https://github.com/Manuelaio/uncoverappLib>

BugReports <https://github.com/Manuelaio/uncoverappLib/issues>

VignetteBuilder knitr

Suggests BiocStyle, knitr, testthat, rmarkdown, dplyr

biocViews Software, Visualization, Annotation, Coverage

NeedsCompilation no

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

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.get_cache	<i>wrapper function for getting BiocFileCache associated with uncoverapp package wrapper function for getting BiocFileCache associated with uncoverapp package</i>
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Description

wrapper function for getting BiocFileCache associated with uncoverapp package wrapper function for getting BiocFileCache associated with uncoverapp package

Usage

.get_cache()

Value

BiocFileCache object associated with uncoverappLib

buildInput	<i>Build input file</i>
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Description

Function to build input file for unCOVERAPP when the number of genes to analyze is > 50.

Usage

```
buildInput(
  geneList,
  genome,
  type_bam,
  bamList,
  outDir,
  MAPQ.min = 1,
  base.quality = 1
)
```

Arguments

geneList	a text file, named with .txt extension, containing HGNC official gene name(s) one per row.
genome	(char) reference genome, hg19 or hg38
type_bam	(char) chromosome notation of their BAM file(s). Use "number" or "chr". In the BAM file, the number option refers to 1, 2, ..., X,.M chromosome notation, while the chr option refers to chr1, chr2, ... chrX, chrM chromosome notation.
bamList	a text file, named with .list extension, containing the absolute paths to BAM file(s) one per row.
outDir	(char) directory where pileup output will be stored
MAPQ.min	(integer) minimum MAPQ value for an alignment to be included in output file.
base.quality	(integer) minimum QUAL value for each nucleotide in an alignment.

Value

A file.bed containing tab-separated specifications of genomic coordinates (chromosome, start position, end position), the coverage value, and the reference:alternate allele counts for each position.

Examples

```
gene.list<- system.file("extdata", "mygene.txt", package = "uncoverappLib")

bam_example <- system.file("extdata", "example_POLG.bam",
  package = "uncoverappLib")
cat(bam_example, file = "bam.list", sep = "\n")
temp_dir=tempdir()
buildInput(geneList= gene.list, genome= "hg19", type_bam= "chr",
  bamList= "bam.list", outDir= temp_dir)
```

getAnnotationFiles	<i>download and rename sorted.bed.gz and sorted.bed.gz.tbi files for annotation of low-coverage positions.</i>
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Description

download and rename sorted.bed.gz and sorted.bed.gz.tbi files for annotation of low-coverage positions.

Usage

```
getAnnotationFiles(verbose = FALSE, vignette = FALSE)
```

Arguments

verbose	(logical) print messages
vignette	(logical) download example annotation-file in vignette mode

Value

(char) Path to local cached file or initial download is required

Examples

```
getAnnotationFiles(verbose = TRUE, vignette= TRUE)
```

run.uncoverapp	<i>run.uncoverapp</i>
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Description

This function launches unCOVERApp, a Shiny application for clinical assessment of sequence coverage.

Usage

```
run.uncoverapp()
```

Value

This return a Shiny App. The is no value

Author(s)

Emanuela Iovino

Examples

```
file.name='../path/sorted.bed.gz'
tbi='../path/sorted.bed.gz.tbi'
run.uncoverapp()

## Only run this example in interactive R sessions

if (interactive()) {
  run.uncoverapp()
}

#After running `run.uncoverapp()` the shiny app appears in your browser
```

uncoverappLib	<i>uncoverappLib: Interactive graphical application for clinical assessment of sequence coverage at the base-pair level</i>
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Author(s)

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Authors:

- Tommaso Pippucci <tommaso.pippucci@unibo.it>

See Also

Useful links:

- <https://github.com/Manuelaio/uncoverappLib>
- Report bugs at <https://github.com/Manuelaio/uncoverappLib/issues>

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