

Package ‘ProActive’

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Title Detect Elevations and Gaps in Mapped Sequencing Read Coverage

Version 0.1.0

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Description Automate the detection of gaps and elevations in mapped sequencing read coverage using a 2D pattern-matching algorithm. 'ProActive' detects, characterizes and visualizes read coverage patterns in both genomes and metagenomes. Optionally, users may provide gene annotations associated with their genome or metagenome in the form of a .gff file. In this case, 'ProActive' will generate an additional output table containing the gene annotations found within the detected regions of gapped and elevated read coverage. Additionally, users can search for gene annotations of interest in the output read coverage plots.

License GPL-2

Encoding UTF-8

LazyData true

RoxygenNote 7.3.2

URL <https://github.com/jlmaier12/ProActive>,
<https://jlmaier12.github.io/ProActive/>

BugReports <https://github.com/jlmaier12/ProActive/issues>

Imports utils, stats, dplyr, ggplot2, stringr

Suggests knitr, rmarkdown, testthat (>= 3.0.0), kableExtra

VignetteBuilder knitr

Depends R (>= 4.2.0)

Config/testthat/edition 3

NeedsCompilation no

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Contents

| | |
|--------------------------------|----------|
| geneAnnotationSearch | 2 |
| plotProActiveResults | 3 |
| ProActiveDetect | 4 |
| Index | 6 |

geneAnnotationSearch *Search for gene annotations on classified contigs/chunks*

Description

Search contigs classified with ProActive for gene-annotations that match a provided key-word(s). Outputs read coverage plots for contigs/chunks with matching annotations.

Usage

```
geneAnnotationSearch(
  ProActiveResults,
  pileup,
  gffTSV,
  geneOrProduct,
  keyWords,
  inGapOrElev = FALSE,
  bpRange = 0,
  elevFilter,
  saveFilesTo,
  verbose = TRUE
)
```

Arguments

| | |
|------------------|---|
| ProActiveResults | The output from 'ProActive()'. |
| pileup | A .txt file containing mapped sequencing read coverages averaged over 100 bp windows/bins. |
| gffTSV | A .gff file (TSV) containing gene predictions associated with the .fasta file used to generate the pileup. |
| geneOrProduct | "gene" or "product". Search for keyWords associated with genes or gene products. |
| keyWords | The keyWord(s) to search for. Case independent. Searches will return the string that contains the matching keyWord. KeyWord(s) must be in quotes, comma-separated, and surrounded by c() i.e(c("antibiotic", "resistance", "drug")) |
| inGapOrElev | TRUE or FALSE. If TRUE, only search for gene-annotations in the gap/elevation region of the pattern-match. Default is FALSE (i.e search the entire contig/chunk for the gene annotation key-words) |

ProActiveDetect *Detect elevations and gaps in mapped read coverage patterns.*

Description

Performs read coverage pattern-matching and summarizes the results into a list. The first list item summarizes the pattern-matching results. The second list item is the 'cleaned' version of the summary table with all the 'noPattern' classifications removed. (i.e were not filtered out). The third list item contains the pattern-match information needed for pattern-match visualization with 'plot-ProActiveResults()'. The fourth list item is a table containing all the contigs that were filtered out prior to pattern-matching. The fifth list item contains arguments used during pattern-matching (windowSize, mode, chunkSize, chunkContigs). If the user provides a gffTSV files, then the last list is a table consisting of ORFs found within the detected gaps and elevations in read coverage.

Usage

```
ProActiveDetect(
  pileup,
  mode,
  gffTSV,
  windowSize = 1000,
  chunkContigs = FALSE,
  minSize = 10000,
  maxSize = Inf,
  minContigLength = 30000,
  chunkSize = 1e+05,
  IncludeNoPatterns = FALSE,
  verbose = TRUE,
  saveFilesTo
)
```

Arguments

| | |
|--------------|---|
| pileup | A .txt file containing mapped sequencing read coverages averaged over 100 bp windows/bins. |
| mode | Either "genome" or "metagenome" |
| gffTSV | Optional, a .gff file (TSV) containing gene predictions associated with the .fasta file used to generate the pileup. |
| windowSize | The number of basepairs to average read coverage values over. Options are 100, 200, 500, 1000 ONLY. Default is 1000. |
| chunkContigs | TRUE or FALSE, If TRUE and 'mode'="metagenome", contigs longer than the 'chunkSize' will be 'chunked' into smaller subsets and pattern-matching will be performed on each subset. Default is FALSE. |
| minSize | The minimum size (in bp) of elevation or gap patterns. Default is 10000. |
| maxSize | The maximum size (in bp) of elevation or gap patterns. Default is NA (i.e. no maximum). |

| | |
|-------------------|--|
| minContigLength | The minimum contig/chunk size (in bp) to perform pattern-matching on. Default is 25000. |
| chunkSize | If 'mode'="genome" OR if 'mode'="metagenome" and 'chunkContigs'=TRUE, chunk the genome or contigs, respectively, into smaller subsets for pattern-matching. 'chunkSize' determines the size (in bp) of each 'chunk'. Default is 100000. |
| IncludeNoPatterns | TRUE or FALSE, If TRUE the noPattern pattern-matches will be included in the ProActive PatternMatches output list. If you would like to visualize the noPattern pattern-matches in 'plotProActiveResults()', this should be set to TRUE. |
| verbose | TRUE or FALSE. Print progress messages to console. Default is TRUE. |
| saveFilesTo | Optional, Provide a path to the directory you wish to save output to. A folder will be made within the provided directory to store results. |

Value

A list containing 6 objects described in the function description.

Examples

```
metagenome_results <- ProActiveDetect(  
  pileup = sampleMetagenomePileup,  
  mode = "metagenome",  
  gffTSV = sampleMetagenomegffTSV  
)
```

Index

geneAnnotationSearch, [2](#)

plotProActiveResults, [3](#)

ProActiveDetect, [4](#)