

Package ‘PGA’

April 23, 2016

Type Package

Title An package for identification of novel peptides by customized database derived from RNA-Seq

Description This package provides functions for construction of customized protein databases based on RNA-Seq data, database searching, post-processing and report generation. This kind of customized protein database includes both the reference database (such as Refseq or ENSEMBL) and the novel peptide sequences form RNA-Seq data.

Version 1.0.0

Date 2014-12-31

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Depends R (>= 3.0.1), IRanges, GenomicRanges, Biostrings (>= 2.26.3),
data.table, rTANDEM

Imports Rsamtools (>= 1.10.2), GenomicFeatures (>= 1.19.8), biomaRt
(>= 2.17.1), stringr, RCurl, Nozzle.R1, VariantAnnotation (>= 1.7.28), rtracklayer, RSQLite, ggplot2, AnnotationDbi,
customProDB (>= 1.7.0), pheatmap

Suggests BSgenome.Hsapiens.UCSC.hg19, RUnit, BiocGenerics, BiocStyle,
knitr, R.utils

VignetteBuilder knitr

License GPL-2

biocViews Proteomics, MassSpectrometry, Software, ReportWriting,
RNASeq, Sequencing

NeedsCompilation no

R topics documented:

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Index**11****dbCreator***Create customized protein database from RNA-Seq data***Description**

The main function to create customized protein database from RNA-Seq data

Usage

```
dbCreator(gtffFile = NULL, vcfFile = NULL, bedFile = NULL,
annotation_path = NULL, outdir, outfile_name, lablersid = FALSE,
COSMIC = FALSE, bool_get_longest = TRUE, organism = "Homo sapiens",
make_decoy = TRUE, genome = NULL, ...)
```

Arguments

| | |
|------------------|---|
| gtffFile | A GTF format file containing novel transcripts information |
| vcfFile | A VCF format file containing SNV and INDEL information |
| bedFile | A BED format file containing juction information |
| annotation_path | This directory contains numerous pieces of genome annotation information which can be downloaded by PrepareAnnotationEnsembl2 or PrepareAnnotationRefseq2 . |
| outdir | Output directory |
| outfile_name | Output file name |
| lablersid | A logical indicating whether to do the SNV annotation(dbSNP) |
| COSMIC | A logical indicating whether to do the SNV annotation(COSMIC) |
| bool_get_longest | When it's set as TRUE, the longest sequences will be retained after the DNA sequences are six-frame translated into protein sequences. Otherwise, the protein sequences more than 30 aa are retained. |
| organism | What is the Genus and species of this organism.Please use proper scientific nomenclature for example: "Homo sapiens" and not "human", default is "Homo sapiens". |
| make_decoy | A logical indicating whether to add the decoy sequences |
| genome | Genome information. This is a BSgenome object(e.g. Hsapiens). Default is NULL. |
| ... | Additional arguments |

Value

The database file

Examples

```
vcffile <- system.file("extdata/input", "PGA.vcf", package="PGA")
bedfile <- system.file("extdata/input", "junctions.bed", package="PGA")
gtffile <- system.file("extdata/input", "transcripts.gtf", package="PGA")
annotation <- system.file("extdata", "annotation", package="PGA")
outfile_path<- "db/"
outfile_name<-"test"
library(BSgenome.Hsapiens.UCSC.hg19)
dbCreator(gtffile=gtffile,vcffile=vcffile,bedfile=bedfile,
           annotation_path=annotation,outfile_name=outfile_name,
           genome=Hsapiens,outdir=outfile_path)
```

easyRun

easyRun

Description

This function is used to automate the peptide identification based on searching the customized database derived from RNA-Seq data.

Usage

```
easyRun(gtffile = NULL, vcfFile = NULL, bedFile = NULL, spectra = NULL,
        annotation_path = NULL, outdir = "pga_dir", outPrefix = "pga",
        lablersid = FALSE, COSMIC = FALSE, bool_get_longest = TRUE,
        organism = "Homo sapiens", genome = NULL, enzyme = "[KR][X]",
        tol = 10, tolu = "ppm", itol = 0.6, itolu = "Daltons",
        varmod = NULL, fixmod = NULL, miss = 2, maxCharge = 8, ti = FALSE,
        cpu = 0, alignment = 1, xmx = 2, ...)
```

Arguments

| | |
|-----------------|---|
| gtffile | A GTF format file containing novel transcripts information |
| vcfFile | A VCF format file containing SNV and INDEL information |
| bedFile | A BED format file containing juction information |
| spectra | MS/MS peak list file |
| annotation_path | This directory contains numerous pieces of genome annotation information which can be downloaded by PrepareAnnotationEnsembl2 or PrepareAnnotationRefseq2 . |
| outdir | Output directory. |
| outPrefix | The prefix of output file. |
| lablersid | A logical indicating whether to do the SNV annotation(dbSNP) |

| | |
|------------------|---|
| COSMIC | A logical indicating whether to do the SNV annotation(COSMIC) |
| bool_get_longest | When it's set as TRUE, the longest sequences will be retained after the DNA sequences are six-frame translated into protein sequences. Otherwise, the protein sequences more than 30 aa are retained. |
| organism | What is the Genus and species of this organism. Please use proper scientific nomenclature for example: "Homo sapiens" and not "human", default is "Homo sapiens". |
| genome | Genome information. This is a BSgenome object(e.g. Hsapiens). |
| enzyme | Specification of specific protein cleavage sites. Default is "[KR][X]". |
| tol | Parent ion mass tolerance (monoisotopic mass). |
| tolu | Parent ion M+H mass tolerance window units. |
| itol | Fragment ion mass tolerance (monoisotopic mass). |
| itolu | Unit for fragment ion mass tolerance (monoisotopic mass). |
| varmod | Specification of potential modifications of residues. |
| fixmod | Specification of modifications of residues. |
| miss | The number of missed cleavage sites. Default is 2. |
| maxCharge | The Maximum parent charge, default is 8 |
| ti | anticipate carbon isotope parent ion assignment errors. Default is false. |
| cpu | The number of CPU used for X!Tandem search. Default is 1. |
| alignment | 0 or 1 to determine if peptide should be alignment or not. Default is 0. |
| xmx | The maximum Java heap size. The unit is "G". |
| ... | Additional arguments |

Value

none

Examples

```

vcffile <- system.file("extdata/input", "PGA.vcf", package="PGA")
bedfile <- system.file("extdata/input", "junctions.bed", package="PGA")
gtffile <- system.file("extdata/input", "transcripts.gtf", package="PGA")
annotation <- system.file("extdata", "annotation", package="PGA")
library(BSgenome.Hsapiens.UCSC.hg19)
msfile <- system.file("extdata/input", "pga.mgf", package="PGA")
easyRun(gtffile=gtffile, vcffile=vcffile, bedFile=bedfile, spectra=msfile,
        annotation_path=annotation, genome=Hsapiens, cpu = 6,
        enzyme = "[KR][X]", varmod = "15.994915@M", itol = 0.05,
        fixmod = "57.021464@C", tol = 10, tolu = "ppm", itolu = "Daltons",
        miss = 2, maxCharge = 8, ti = FALSE, xmx=1)

```

Description

This function is mainly for q-value calculation, protein inference and novel peptides spectra annotation.

Usage

```
parserGear(file = NULL, db = NULL, outdir = "parser_outdir",
           prefix = "pga", novelPrefix = "VAR", decoyPrefix = "###REV###",
           alignment = 1, xmx = NULL, thread = 1)
```

Arguments

| | |
|-------------|--|
| file | MS/MS search file. Currently, only XML format file of X!Tandem and DAT result of Mascot are supported. |
| db | A FASTA format database file used for MS/MS searching. Usually, it is from the output of the function dbCreator. |
| outdir | Output directory. |
| prefix | The prefix of output file. |
| novelPrefix | The prefix of novel protein ID. Default is "VAR". "VAR" is the prefix which used by function dbCreator. |
| decoyPrefix | The prefix of decoy sequences ID. Default is "###REV###". "###REV###" is the prefix which used by function dbCreator. |
| alignment | 0 or 1 to determine if peptide should be alignment or not. Default is 1. |
| xmx | The maximum Java heap size. The unit is "G". |
| thread | This parameter is used to specify the number of threads. "0" represents that all of the available threads are used; "1" represents one thread is used; "2" represents two threads are used, and so on. Default is 1. |

Value

none

Examples

```
vcffile <- system.file("extdata/input", "PGA.vcf", package="PGA")
bedfile <- system.file("extdata/input", "junctions.bed", package="PGA")
gtffile <- system.file("extdata/input", "transcripts.gtf", package="PGA")
annotation <- system.file("extdata", "annotation", package="PGA")
outfile_path<- "db/"
outfile_name<- "test"
library(BSgenome.Hsapiens.UCSC.hg19)
```

```

dbfile <- dbCreator(gtffile=gtffile,vcffile=vcffile,bedFile=bedfile,
                     annotation_path=annotation,outfile_name=outfile_name,
                     genome=Hsapieins,outdir=outfile_path)

msfile <- system.file("extdata/input", "pga.mgf", package="PGA")
idfile <- runTandem(spectra = msfile, fasta = dbfile, outdir = "./", cpu = 6,
                     enzyme = "[KR][X]", varmod = "15.994915@M", itol = 0.05,
                     fixmod = "57.021464@C", tol = 10, tolu = "ppm",
                     itolu = "Daltons", miss = 2, maxCharge = 8, ti = FALSE)
parserGear(file = idfile, db = dbfile, decoyPrefix="#REV#", xmx=1, thread=8,
           outdir = "parser_outdir")

```

PrepareAnnotationEnsembl2*Prepare annotation from ENSEMBL***Description**

Prepare the annotation from ENSEMBL through biomaRt. This function is modified from the function [PrepareAnnotationEnsembl](#) in **customProDB**.

Usage

```
PrepareAnnotationEnsembl2(mart, annotation_path, splice_matrix = FALSE,
                          dbsnp = NULL, transcript_ids = NULL, COSMIC = FALSE, ...)
```

Arguments

- `mart` See detail in function [PrepareAnnotationEnsembl](#).
- `annotation_path` See detail in function [PrepareAnnotationEnsembl](#).
- `splice_matrix` See detail in function [PrepareAnnotationEnsembl](#).
- `dbsnp` See detail in function [PrepareAnnotationEnsembl](#).
- `transcript_ids` See detail in function [PrepareAnnotationEnsembl](#).
- `COSMIC` See detail in function [PrepareAnnotationEnsembl](#).
- `...` Additional arguments

Value

Several .RData file containing annotations needed for following analysis.

See Also

[PrepareAnnotationRefseq2](#).

Examples

```
ensembl <- biomaRt::useMart("ENSEMBL_MART_ENSEMBL", dataset="hsapiens_gene_ensembl",
                             host="feb2012.archive.ensembl.org", path="/biomart/martservice",
                             archive=FALSE)

annotation_path <- tempdir()
transcript_ids <- c("ENST00000234420", "ENST00000269305", "ENST00000445888")

PrepareAnnotationEnsembl2(mart=ensembl, annotation_path=annotation_path,
                          splice_matrix=FALSE, dbsnp=NULL, transcript_ids=transcript_ids,
                          COSMIC=FALSE)
```

PrepareAnnotationRefseq2

Prepare annotation from Refseq

Description

Prepare the annotation for Refseq through UCSC table browser. This function is modified from the function [PrepareAnnotationRefseq](#) in **customProDB**.

Usage

```
PrepareAnnotationRefseq2(genome = "hg19", CDSfasta, pepfasta, annotation_path,
                        dbsnp = NULL, transcript_ids = NULL, splice_matrix = FALSE,
                        COSMIC = FALSE, ...)
```

Arguments

| | |
|-----------------|--|
| genome | See detail in function PrepareAnnotationRefseq . |
| CDSfasta | See detail in function PrepareAnnotationRefseq . |
| pepfasta | See detail in function PrepareAnnotationRefseq . |
| annotation_path | See detail in function PrepareAnnotationRefseq . |
| dbsnp | See detail in function PrepareAnnotationRefseq . |
| transcript_ids | See detail in function PrepareAnnotationRefseq . |
| splice_matrix | See detail in function PrepareAnnotationRefseq . |
| COSMIC | See detail in function PrepareAnnotationRefseq . |
| ... | Additional arguments |

Value

Several .RData file containing annotations needed for following analysis.

See Also

PrepareAnnotationEnsembl2.

Examples

reportGear

The main function for report generation

Description

The main function for report generation

Usage

```
reportGear(parser_dir, tab_dir, report_dir)
```

Arguments

| | |
|------------|---|
| parser_dir | The directory which contains the peptide identification results |
| tab_dir | The directory which contains the database annotation files |
| report_dir | The report output directory |

Value

none

Examples

```

vcffile <- system.file("extdata/input", "PGA.vcf", package="PGA")
bedfile <- system.file("extdata/input", "junctions.bed", package="PGA")
gtffile <- system.file("extdata/input", "transcripts.gtf", package="PGA")
annotation <- system.file("extdata", "annotation", package="PGA")
outfile_path<-"db/"
outfile_name<-"test"
library(BSgenome.Hsapiens.UCSC.hg19)
dbfile <- dbCreator(gtffFile=gtffile, vcffile=vcffile, bedFile=bedfile,
                     annotation_path=annotation, outfile_name=outfile_name,
                     genome=Hsapiens, outdir=outfile_path)

```

```

msfile <- system.file("extdata/input", "pga.mgf", package="PGA")
idfile <- runTandem(spectra = msfile, fasta = dbfile, outdir = "./", cpu = 6,
                     enzyme = "[KR][X]", varmod = "15.994915@M", itol = 0.05,
                     fixmod = "57.021464@C", tol = 10, tolu = "ppm",
                     itolu = "Daltons", miss = 2, maxCharge = 8, ti = FALSE)
parserGear(file = idfile, db = dbfile, decoyPrefix="#REV#", xmx=1, thread=8,
           outdir = "parser_outdir")
reportGear(parser_dir = "parser_outdir", tab_dir = outfile_path,
           report_dir = "report")

```

runTandem

*run X!Tandem***Description**

run X!Tandem

Usage

```
runTandem(spectra = "", fasta = "", outdir = ".", cpu = 1,
          enzyme = "[KR][X]", tol = 10, tolu = "ppm", itol = 0.6,
          itolu = "Daltons", varmod = NULL, fixmod = NULL, miss = 2,
          maxCharge = 8, ti = FALSE)
```

Arguments

| | |
|-----------|---|
| spectra | MS/MS peak list file |
| fasta | Protein database file for searching. |
| outdir | The output directory. |
| cpu | The number of CPU used for X!Tandem search. Default is 1. |
| enzyme | Specification of specific protein cleavage sites. Default is "[KR][X]". |
| tol | Parent ion mass tolerance (monoisotopic mass). |
| tolu | Parent ion M+H mass tolerance window units. |
| itol | Fragment ion mass tolerance (monoisotopic mass). |
| itolu | Unit for fragment ion mass tolerance (monoisotopic mass). |
| varmod | Specification of potential modifications of residues. |
| fixmod | Specification of modifications of residues. |
| miss | The number of missed cleavage sites. Default is 2. |
| maxCharge | The Maximum parent charge, default is 8 |
| ti | anticipate carbon isotope parent ion assignment errors. Default is false. |

Value

The search result file path

Examples

```
vcffile <- system.file("extdata/input", "PGA.vcf", package="PGA")
bedfile <- system.file("extdata/input", "junctions.bed", package="PGA")
gtffile <- system.file("extdata/input", "transcripts.gtf", package="PGA")
annotation <- system.file("extdata", "annotation", package="PGA")
outfile_path<- "db/"
outfile_name<-"test"
library(BSgenome.Hsapiens.UCSC.hg19)
dbfile <- dbCreator(gtfFile=gtffile,vcffile=vcffile,bedFile=bedfile,
                     annotation_path=annotation,outfile_name=outfile_name,
                     genome=Hsapiens,outdir=outfile_path)

msfile <- system.file("extdata/input", "pga.mgf", package="PGA")
runTandem(spectra = msfile, fasta = dbfile, outdir = "./", cpu = 6,
           enzyme = "[KR][X]", varmod = "15.994915@M",
           fixmod = "57.021464@C", tol = 10, tolu = "ppm", itol = 0.05,
           itolu = "Daltons", miss = 2, maxCharge = 8, ti = FALSE)
```

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