

Package ‘sapFinder’

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Type Package

Title A package for variant peptides detection and visualization in shotgun proteomics.

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Imports pheatmap,Rcpp (>= 0.10.6),graphics,grDevices,stats, utils

biocViews MassSpectrometry, Proteomics, SNP, RNASeq, Visualization, ReportWriting

Description sapFinder is developed to automate

- (1) variation-associated database construction,
- (2) database searching,
- (3) post-processing,
- (4) HTML-based report generation in shotgun proteomics.

License GPL-2

LazyLoad yes

LinkingTo Rcpp

NeedsCompilation yes

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dbCreator

dbCreator

Description

An integrated function to generate variation-associated database based on sample-specific NGS data or public SNV data.

Usage

```
dbCreator(vcf = NULL, annotation = NULL, refseq = NULL, outdir = "./",
prefix = "test", xmx = NULL, xref = "noxref")
```

Arguments

<code>vcf</code>	Input VCF file name. This file contains the information of gene sequence variations.
<code>annotation</code>	Input annotation file name. It contains the gene annotation information and can be downloaded from UCSC Genome Browser. Currently it supports RefSeq genes and ENSEMBL genes annotation file.
<code>refseq</code>	Input mRNA sequences file with FASTA format. It can be downloaded from UCSC Genome Browser.
<code>outdir</code>	Output directory.
<code>prefix</code>	The prefix of output file.
<code>xmx</code>	The maximum Java heap size. The unit is "G".
<code>xref</code>	Optional external cross-reference file, generally it's downloaded through BioMart. If this file is provided, the final html report will present some relevant protein id or description.

Value

A vector containing two file names. One is a FASTA format file contains the mutated peptides, the normal protein sequences and their reverse versions, and the other is a tab-delimited file contains detailed variation information.

Examples

```
vcf      <- system.file("extdata/sapFinder_test.vcf",
                      package="sapFinder")
annotation <- system.file("extdata/sapFinder_test_ensGene.txt",
                          package="sapFinder")
refseq    <- system.file("extdata/sapFinder_test_ensGeneMrna.fa",
                      package="sapFinder")
xref      <- system.file("extdata/sapFinder_test_BioMart.Xref.txt",
                      package="sapFinder")
outdir    <- "db_dir"
```

```
prefix      <- "sapFinder_test"
db.files <- dbCreator(vcf=vcf, annotation=annotation,
                      refseq=refseq, outdir=outdir,
                      prefix=prefix, xref=xref)
```

easyRun***easyRun***

Description

This function is used to automate the variation-associated database construction, MS/MS searching, post-processing and HTML-based report generation.

Usage

```
easyRun(vcf = NULL, annotation = NULL, refseq = NULL, outdir = "./",
         prefix = "sapFinder_test", spectra = "", cpu = 1, enzyme = "[KR][X]",
         tol = 10, tolu = "ppm", itol = 0.6, itolu = "Daltons",
         varmod = NULL, fixmod = NULL, miss = 2, maxCharge = 8, ti = FALSE,
         alignment = 1, xref = "noxref", xmx = NULL, ...)
```

Arguments

vcf	Input VCF file name. This file contains the information of gene sequence variations.
annotation	Input annotation file name. It contains the gene annotation information and can be downloaded from UCSC Genome Browser. Currently it supports RefSeq genes and ENSEMBL genes annotation file.
refseq	Input mRNA sequences file with FASTA format. It can be downloaded from UCSC Genome Browser.
outdir	Output directory.
prefix	The prefix of output file.
spectra	MS/MS peak list file
cpu	The number of CPU used for X!Tandem search. Default is 1.
enzyme	Specification of specific protein cleavage sites. Default is "[KR][X]".
varmod	Specificiation of potential modifications of residues.
fixmod	Specification of modifications of residues.
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).
miss	The number of missed cleavage sites. Default is 2.
maxCharge	The Maximum parent charge, default is 8

<code>ti</code>	anticipate carbon isotope parent ion assignment errors. Default is false.
<code>alignment</code>	0 or 1 to determine if peptide should be alignment or not. Default is 0.
<code>xmx</code>	The maximum Java heap size. The unit is "G".
<code>xref</code>	Optional external cross-reference file, generally it's downloaded through BioMart. If this file is provided, the final html report will present some relevant protein id or description.
<code>...</code>	Additional arguments

Examples

```

vcf      <- system.file("extdata/sapFinder_test.vcf",
                      package="sapFinder")
annotation <- system.file("extdata/sapFinder_test_ensGene.txt",
                          package="sapFinder")
refseq    <- system.file("extdata/sapFinder_test_ensGeneMrna.fa",
                          package="sapFinder")
mgf.path  <- system.file("extdata/sapFinder_test.mgf",
                          package="sapFinder")
xref      <- system.file("extdata/sapFinder_test_BioMart.Xref.txt",
                          package="sapFinder")
easyRun(vcf=vcf, annotation=annotation, refseq=refseq, outdir="test",
prefix="sapFinder_test", spectra=mgf.path, cpu=0, tol=10, tolu="ppm", itol=0.1,
itolu="Daltons", alignment=1, xref=xref)

```

Description

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

Usage

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

Arguments

<code>file</code>	MS/MS search file. Currently, only XML format file of X!Tandem and DAT result of Mascot are supported.
<code>db</code>	A FASTA format database file used for MS/MS searching. Usually, it is from the output of the function dbCreator.
<code>outdir</code>	Output directory.
<code>prefix</code>	The prefix of output file.

mutPrefix	The prefix of variant peptides 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.
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.
xmx	The maximum Java heap size. The unit is "G".

Examples

```

## Step 1. Variation-associated database construction
vcf      <- system.file("extdata/sapFinder_test.vcf",
                        package="sapFinder")
annotation <- system.file("extdata/sapFinder_test_ensGene.txt",
                           package="sapFinder")
refseq    <- system.file("extdata/sapFinder_test_ensGeneMrna.fa",
                           package="sapFinder")
xref      <- system.file("extdata/sapFinder_test_BioMart.Xref.txt",
                           package="sapFinder")
outdir    <- "db_dir"
prefix    <- "sapFinder_test"
db.files <- dbCreator(vcf=vcf, annotation=annotation,
                      refseq=refseq, outdir=outdir,
                      prefix=prefix, xref=xref)

## Step 2. MS/MS searching
mgf.path  <- system.file("extdata/sapFinder_test.mgf",
                        package="sapFinder")
fasta.path <- db.files[1]
xml.path   <- runTandem(spectra=mgf.path, fasta=fasta.path, outdir=".",
                        tol=10, tolu="ppm", itolu=0.1, itolu="Daltons")

## Step 3. Post-processing
parserGear(file=xml.path, db=fasta.path, prefix=prefix,
            outdir="parser_outdir", alignment=1)

```

Description

This function is used for HTML-based report writing

Usage

```
reportCreator(indir = ".", outdir = .REPORT.DIR, db = NULL,
              prefix = NULL, varInfor = NULL)
```

Arguments

<code>indir</code>	The directory of output files of function <code>parserGear</code> .
<code>outdir</code>	Output directory for this report
<code>db</code>	A FASTA format database file used for MS/MS searching. Usually, it is from the output of the function <code>dbCreator</code> .
<code>prefix</code>	It must be set the same with the parameter of "prefix" in function <code>parserGear</code> .
<code>varInfor</code>	It is a tab-delimited file contains detailed variation information and is from the output of the function <code>dbCreator</code> .

Examples

```

## Step 1. Variation-associated database construction
vcf      <- system.file("extdata/sapFinder_test.vcf",
                        package="sapFinder")
annotation <- system.file("extdata/sapFinder_test_ensGene.txt",
                           package="sapFinder")
refseq    <- system.file("extdata/sapFinder_test_ensGeneMrna.fa",
                        package="sapFinder")
xref      <- system.file("extdata/sapFinder_test_BioMart.Xref.txt",
                        package="sapFinder")
outdir    <- "db_dir"
prefix    <- "sapFinder_test"
db.files <- dbCreator(vcf=vcf, annotation=annotation,
                      refseq=refseq, outdir=outdir,
                      prefix=prefix,xref=xref)

## Step 2. MS/MS searching
mgf.path  <- system.file("extdata/sapFinder_test.mgf",
                        package="sapFinder")
fasta.path <- db.files[1]
xml.path   <- runTandem(spectra=mgf.path, fasta=fasta.path, outdir=".",
                        tol=10, tolu="ppm", itol=0.1, itolu="Daltons")

## Step 3. Post-processing
parserGear(file=xml.path, db=fasta.path, prefix=prefix,
            outdir="parser_outdir")

## Step 4. HTML-based report generation
reportCreator(indir="parser_outdir", outdir="report", db=fasta.path,
               prefix=prefix, varInfor=db.files[2])

```

`runTandem`

run xtandem

Description

`run xtandem`

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]".
varmod	Specificiation of potential modifications of residues.
fixmod	Specification of modifications of residues.
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).
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

```
# Variation-associated database construction
vcf      <- system.file("extdata/sapFinder_test.vcf",
                        package="sapFinder")
annotation <- system.file("extdata/sapFinder_test_ensGene.txt",
                           package="sapFinder")
refseq    <- system.file("extdata/sapFinder_test_ensGeneMrna.fa",
                        package="sapFinder")
xref      <- system.file("extdata/sapFinder_test_BioMart.Xref.txt",
                        package="sapFinder")
outdir    <- "db_dir"
prefix    <- "sapFinder_test"
db.files <- dbCreator(vcf=vcf, annotation=annotation,
                      refseq=refseq, outdir=outdir,
                      prefix=prefix,xref=xref)

# MS/MS searching
mgf.path  <- system.file("extdata/sapFinder_test.mgf",
```

runTandem

```
    package="sapFinder")
runTandem(spectra=mgf.path,fasta=db.files[1],
          tol=10,tolu="ppm",itol=0.1,itolu="Daltons")
```

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