# Package 'packFinder'

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Type Package

```
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Version 1.20.0
Description Algorithm and tools for in silico pack-TYPE transposon discovery. Fil-
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      gation of returned matches. Sequences are input in DNAString format, and ranges are re-
      turned as a dataframe (in the format returned by as.dataframe(GRanges)).
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```

## **Contents**

	arabidopsisThalianaRefseq	2
	blastAnalysis	3
	blastAnnotate	5
	collapseSeqs	6
	filterWildcards	7
	getPackSeqs	8
	getPacksFromCsv	10
	getPacksFromFasta	11
	getPacksFromGRanges	12
	getTsds	13
	identifyPotentialPackElements	14
	identifyTirMatches	15
	makeBlastDb	17
	packAlign	18
	packBlast	20
	packClust	22
	packFinder	24
	packMatches	25
	packSearch	26
	packsToCsv	28
	packsToFasta	29
	packsToGRanges	30
	readBlast	31
	readUc	33
	tirClust	35
Index		37

arabidopsisThalianaRefseq

Arabidopsis thaliana Refseq Genome Chromosome 3 Subset

### Description

The chromosome 3 reference sequence for Arabidopsis thaliana as a DNAStringSet. Can be used as a test data set, as in the associated introduction vignette. The DNA sequence between bases 10,500,000 and 14,300,000 was extracted for use in this dataset.

### Usage

data(arabidopsisThalianaRefseq)

#### **Format**

A DNAStringSet object containing a DNAString for Arabidopsis thaliana's chromosome 3 sequence.

blastAnalysis 3

### Author(s)

Jack Gisby

#### Source

The Arabidopsis thaliana genome was downloaded from the NCBI refseq database on 20/SEP/2019, using getGenome, and chromosome 3 was extracted. The genome may also be accessed from the NCBI ftp server: ftp://ftp.ncbi.nlm.nih.gov/genomes.

### See Also

```
getGenome, DNAStringSet, DNAString, packSearch
```

### **Examples**

```
data(arabidopsisThalianaRefseq)

packMatches <- packSearch(
    Biostrings::DNAString("CACTACAA"),
    arabidopsisThalianaRefseq,
    elementLength = c(300, 3500),
    tsdLength = 3
)</pre>
```

blastAnalysis

BLAST Analysis of PackTYPE Elements

### **Description**

Run BLAST against user-specified databases of non-transposon and transposon-related proteins. Can be used to classify transposons based on their internal sequences.

```
blastAnalysis(
  packMatches,
  Genome,
  blastPath,
  protDb = NULL,
  autoDb = NULL,
  minE = 0.001,
  blastTask = "blastn-short",
  maxHits = 100,
  threads = 1,
  saveFolder = NULL,
  tirCutoff = 0
)
```

4 blastAnalysis

#### **Arguments**

packMatches A dataframe of potential Pack-TYPE transposable elements, in the format given

by packSearch. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-

object to a dataframe: data.frame(GRanges). Will be saved as a FASTA file

for VSEARCH.

Genome A DNAStringSet object containing sequences referred to in packMatches (the

object originally used to predict the transposons packSearch).

blastPath Path to the BLAST+ executable, or name of the BLAST+ application for Linux/MacOS

users.

protDb For assigning Pack-TYPE elements. Path to the blast database containing nu-

cleotide or protein sequences to be matched against internal transposon sequences.

Can be generated using BLAST+, or with link{makeBlastDb}.

autoDb For assigning autonomous elements. Path to the blast database containing nu-

cleotide or protein sequences to be matched against internal transposon sequences.

Can be generated using BLAST+, or with link{makeBlastDb}.

minE Blast results with e values greater than the specified cutoff will be ignored.

blastTask Type of BLAST+ task, defaults to "blastn-short".

maxHits Maximum hits returned by BLAST+ per query.

threads Allowable number of threads to be utilised by BLAST+.

saveFolder Directory to save BLAST+ results in; defaults to the working directory.

tirCutoff How many bases to ignore at the terminal ends of the transposons to prevent hits

to TIR sequences.

#### Value

No return value; executes BLAST+ to generate hits which are stored in a .blast file in the chosen directory.

### Author(s)

Jack Gisby

### References

For further information, see the NCBI BLAST+ application documentation and help pages (https://www.ncbi.nlm.nih.gov/pul

### See Also

blastAnnotate, readBlast, packBlast

```
## Not run:
packMatches <- data(packMatches)
Genome <- data(arabidopsisThalianaRefseq)</pre>
```

blastAnnotate 5

```
blastAnalysis(packMatches, Genome,
    protDb = "C:/data/TAIR10_CDS",
    autoDb = "C:/data/TAIR10_transposons",
    blastPath = "C:/blast/bin/blastn.exe")
## End(Not run)
```

blastAnnotate

Functional Annotation of PackTYPE Elements

### Description

Uses hits, previously generated using blast, to annotate transposon hits. Transposons with non-redundant transposase hits are classed as autonomous ("auto"), while others are classed as "other" or "pack" based on whether the element has non-redundant hits to other proteins.

### Usage

blastAnnotate(protHits, autoHits, packMatches)

#### **Arguments**

protHits BLAST results for non-transposon related genes or proteins (as a data.frame).

Generated using blastAnalysis.

autoHits BLAST results for transposon related genes or proteins (as a data. frame). Gen-

erated using blastAnalysis.

packMatches A dataframe of potential Pack-TYPE transposable elements, in the format given

by packSearch. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-

object to a dataframe: data.frame(GRanges). Will be saved as a FASTA file

for VSEARCH.

#### Value

Returns the original packMatches dataframe, with the addition of a "classification" column containing one of the following values:

- auto elements that match known transposases or transposon-related proteins are classified as autonomous elements
- pack elements that match other proteins or genic sequences may be classified as Pack-TYPE elements
- other elements that generate no significant hits

#### Note

Requires that the query ids in the protein and autonomous hits match the row names in packMatches.

6 collapseSeqs

### Author(s)

Jack Gisby

#### References

For further information, see the NCBI BLAST+ application documentation and help pages (https://www.ncbi.nlm.nih.gov/pul

#### See Also

blastAnalysis, readBlast, packBlast

### **Examples**

```
data("packMatches")

# read in some protein hits

p <- data.frame(
    query_id = c(2, 3),
    subject_id = c("prot", "hyp")
)

# read in some autonomous hits
a <- data.frame(
    query_id = c(3, 4),
    subject_id = c("transposase", "mutator")
)

blastAnnotate(p, a, packMatches)</pre>
```

collapseSeqs

Collapse Overlapping Sequences

### **Description**

The sequences predicted by packSearch often overlap, which may be due to the presence of closely interspersed elements or false TIR identification. In such cases, these elements can be combined using link[GenomicRanges:GRanges-class]{GRanges} in order to collapse overlapping elements, preventing over-estimation of transposon numbers. Also removes duplicate elements that have been generated in the case of multiple searches.

```
collapseSeqs(packMatches, Genome)
```

filterWildcards 7

### **Arguments**

packMatches

A dataframe containing genomic ranges and names referring to sequences to be

extracted. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-class object to a dataframe: data.frame(GRanges).

Must contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

Genome

A DNAStringSet object containing sequences referred to in packMatches (the object originally used to predict the transposons packSearch).

#### Value

A set of non-overlapping transposon sequences in the format of the input dataframe.

### Author(s)

Jack Gisby

#### See Also

```
packSearch, link[GenomicRanges:GRanges-class]{GRanges}
```

### **Examples**

```
data(packMatches)
data(arabidopsisThalianaRefseq)

packMatches$start <- 1
packMatches$end <- 10

collapseSeqs(packMatches, arabidopsisThalianaRefseq)</pre>
```

filterWildcards

Remove Low Quality Sequences

### Description

Takes transposable elements detected by packSearch and removes those with large numbers of wildcard ("N") bases. Used by packClust and packAlign to remove poor quality sequences that may interfere with the quality of sequence alignments.

```
filterWildcards(packMatches, Genome, maxWildcards = 0.05)
```

8 getPackSeqs

### **Arguments**

packMatches A dataframe containing genomic ranges and names referring to sequences to be

extracted.

Genome The original set of sequences used to generate the transposons detected by packSearch.

maxWildcards The maximal allowable proportion of wildcards in the sequence of each match

(defaults to 0.05).

#### Value

The original dataframe, packMatches, with sequences removed that are found to contain a proportion of wildcards ("N") greater than that specified in maxWildcards.

### Author(s)

Jack Gisby

#### See Also

packClust, packAlign, packMatches, data(arabidopsisThalianaRefseq)

### Examples

```
data(arabidopsisThalianaRefseq)
data(packMatches)

filteredMatches <- filterWildcards(
    packMatches,
    arabidopsisThalianaRefseq,
    maxWildcards = 0.05
)</pre>
```

getPackSeqs

Extract Sequences of Pack-TYPE Elements

### Description

Method to quickly extract the sequences of predicted Pack-TYPE elements (as created by packSearch).

```
getPackSeqs(packMatches, Genome, output = "DNAStringSet")
```

getPackSeqs 9

### **Arguments**

packMatches

A dataframe containing genomic ranges and names referring to sequences to be extracted. This dataframe is in the format produced by coercing a link[GenomicRanges: GRanges-class object to a dataframe: data.frame(GRanges).

Must contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

Genome

A DNAStringSet object containing sequences referred to in packMatches (the object originally used to predict the transposons packSearch).

output

The type of object to be returned:

- output = "DNAStringSet", returns a DNAStringSet object (default).
- output = "character", returns a character vector.

#### Value

transposon sequences extracted from packMatches. At default returns the sequences as a DNAStringSet or, if output is set to "character", returns a character vector.

### Author(s)

Jack Gisby

#### See Also

DNAStringSet, packSearch, DNAString

```
data(arabidopsisThalianaRefseq)

packMatches <- packSearch(
    Biostrings::DNAString("CACTACAA"),
    arabidopsisThalianaRefseq,
    elementLength = c(300, 3500),
    tsdLength = 3
)

packSeqs <- getPackSeqs(packMatches, arabidopsisThalianaRefseq)</pre>
```

10 getPacksFromCsv

getPacksFromCsv

Retrieve Saved packFinder Results (.csv)

### Description

Retrieves a dataframe of potential Pack-TYPE elements, previously saved using packSearch followed by packsToCsv.

### Usage

```
getPacksFromCsv(file)
```

### Arguments

file

File path to predicted transposons in CSV format.

#### Value

Dataframe in the format used by packSearch.

### Author(s)

Jack Gisby

#### See Also

```
packsToCsv, read.table, packSearch
```

```
data(packMatches)

packMatches <- getPacksFromCsv(
    system.file("extdata", "packMatches.csv", package = "packFinder")
)</pre>
```

getPacksFromFasta 11

getPacksFromFasta

Retrieve Saved packFinder Results (.fasta)

### Description

Retrieves a dataframe of potential Pack-TYPE elements, previously saved using packSearch followed by packsToFasta. Parses the .fasta file and title field containing:

- seqnames name of origin sequence
- start transposon base start position on origin sequence
- end transposon base end position on origin sequence
- width width of transposon
- strand direction of transposon ("+", "-" or "\*")
- TSD terminal site duplication (TSD) sequence

### Usage

```
getPacksFromFasta(file)
```

### **Arguments**

file

Path to predicted transposons in FASTA format.

### Value

Dataframe in the format used by packSearch.

### Author(s)

Jack Gisby

### See Also

```
packsToFasta, packSearch
```

```
data(arabidopsisThalianaRefseq)
data(packMatches)

packMatches <- getPacksFromFasta(
    system.file("extdata", "packMatches.fasta", package = "packFinder")
)</pre>
```

getPacksFromGRanges

Retrieve packFinder Results from GRanges Object

### **Description**

A link[GenomicRanges:GRanges-class]{GRanges} object, potentially generated using packSearch and packsToGRanges, can be converted to a dataframe. If a GRanges object is supplied without TSD information, this can be calculated and appended to the final dataframe.

### Usage

```
getPacksFromGRanges(packGRanges, Genome = NULL, tsdLength = NULL)
```

### **Arguments**

packGRanges link[GenomicRanges:GRanges-class]{GRanges} object to be coerced.

Genome (optional) Sequences referred to by packGRanges.

tsdLength (optional) Length of TSD sequences.

### Value

Dataframe in the format used by packSearch. If Genome and tsdLength are supplied, then TSD sequences are retrieved and returned as part of the dataframe.

### Author(s)

Jack Gisby

### See Also

```
packsToGRanges, link[GenomicRanges:GRanges-class]{GRanges}, packSearch
```

```
data(packMatches)

GRangesObject <- packsToGRanges(packMatches)
packMatches <- getPacksFromGRanges(GRangesObject)</pre>
```

getTsds 13

getTsds	Get Flanking Terminal Site Duplication Sequences

#### **Description**

Gets the flanking TSD sequences of TIRs or predicted Pack-TYPE transposable elements. A dataframe of these elements can be in tirMatches.

### Usage

```
getTsds(tirMatches, Genome, tsdLength, strand = "+", output = "character")
```

### **Arguments**

tirMatches A dataframe containing genomic ranges and names referring to TIR sequences

or predicted Pack-TYPE transposable elements. Should be in the format used

by packSearch.

Genome A DNAStringSet object containing sequences referred to in tirMatches.

tsdLength The length of the TSD region to be retrieved (integer).

strand The strand of the TIR; "+" for forward, "-" for reverse. If the TSD sequences

of transposable elements are being predicted, then this parameter can be left as default ("+"); if the TSD sequences of TIRs are being found then the strand

direction must be supplied.

output The type of object to be returned:

• output = "DNAStringSet", returns a DNAStringSet object.

• output = "character", returns a character vector (default).

### **Details**

Called by packSearch. It is recommended to use the general pipeline function packSearch for identification of potential pack elements, which returns TSD sequences as a feature of results, however each stage may be called individually.

#### Value

Flanking TSD sequences as a vector of characters, or if output is specified as "DNAStringSet", TSD sequences will be returned as a DNAStringSet object.

### Author(s)

Jack Gisby

#### See Also

DNAStringSet, packSearch, tirMatches

#### **Examples**

```
data(arabidopsisThalianaRefseq)
data(packMatches)

tsdSeqs <- getTsds(packMatches, arabidopsisThalianaRefseq, 3)</pre>
```

identifyPotentialPackElements

Pack Element Filtering

### Description

Primary filtering stage for the packSearch algorithm. Identifies potential Pack-TYPE transposable elements based on proximity of matching inverted repeats and equality of TSD sequences.

### Usage

```
identifyPotentialPackElements(
  forwardMatches,
  reverseMatches,
  Genome,
  elementLength,
  tsdMismatch = 0
)
```

### **Arguments**

forwardMatches A dataframe containing genomic ranges and names referring to forwards-facing

TIR sequences and their respective TSD sequences.

reverseMatches A dataframe containing genomic ranges and names referring to reverse-facing

TIR sequences and their respective TSD sequences.

Genome A DNAStringSet object containing the matches referred to in forwardMatches

and reverseMatches

elementLength A vector of two integers containing the minimum and maximum transposable

element length.

tsdMi smatch An integer referring to the allowable mismatch (substitutions or indels) between

a transposon's TSD sequences. matchPattern from Biostrings is used for pat-

tern matching.

#### Details

Used by packSearch as a primariy filtering stage. Identifies matches likely to be transposons based on their TIR region, from identifyTirMatches, and their TSD region, from getTsds. It is recommended to use the general pipeline function packSearch for identification of potential pack elements, however each stage may be called individually. Note that only exact TSD matches are considered, so supplying long sequences for TSD elements may lead to false-negative results.

identifyTirMatches 15

### Value

A dataframe, packMatches, containing the locations of potential Pack-TYPE transposable elements in Genome.

### Author(s)

Jack Gisby

#### See Also

packSearch

### **Examples**

```
data(arabidopsisThalianaRefseq)
forwardMatches <- identifyTirMatches(</pre>
   Biostrings::DNAString("CACTACAA"),
   arabidopsisThalianaRefseq,
    tsdLength = 3,
    strand = "+"
)
reverseMatches <- identifyTirMatches(</pre>
    Biostrings::reverseComplement(Biostrings::DNAString("CACTACAA")),
    arabidopsisThalianaRefseq,
    tsdLength = 3,
    strand = "-"
)
packMatches <- identifyPotentialPackElements(</pre>
    forwardMatches,
    {\tt reverseMatches},
   arabidopsisThalianaRefseq,
   c(300, 3500)
)
```

identifyTirMatches

Identify Terminal Inverted Repeat Matches

### Description

Searches a DNAStringSet for potential TIRs based on sequence similarity.

16 identifyTirMatches

### Usage

```
identifyTirMatches(
   tirSeq,
   Genome,
   mismatch = 0,
   strand = "*",
   tsdLength,
   fixed = TRUE
)
```

### **Arguments**

tirSeq A DNAString object to be searched for.

Genome A DNAStringSet object containing the DNAString objects to be searched.

mismatch The allowable mismatch between tirSeq and a given slice of Genome. Includes

indels.

strand The directionality of the search string ("+" or "-"). Note that this does affect the

search for tirSeqs, if you wish to search the reverse strand you should use the

reverse complement of your sequence.

tsdLength Integer referring to the length of the flanking TSD region.

fixed Logical that will be passed to the 'fixed' argument of matchPattern. Deter-

mines the behaviour of IUPAC ambiguity codes when searching for TIR se-

quences.

### **Details**

Called by packSearch. Used by packSearch as an initial filtering stage. matchPattern from Biostrings is used for pattern matching. It is recommended to use the general pipeline function packSearch for identification of potential pack elements, however each stage may be called individually.

#### Value

A dataframe, tirMatches, containing identified matches. The dataframe is in the format generated by packSearch.

### Author(s)

Jack Gisby

### See Also

DNAStringSet, packSearch, matchPattern, DNAString

makeBlastDb 17

### **Examples**

```
data(arabidopsisThalianaRefseq)

forwardMatches <- identifyTirMatches(
    Biostrings::DNAString("CACTACAA"),
    arabidopsisThalianaRefseq,
    tsdLength = 3,
    strand = "+"
)</pre>
```

makeBlastDb

Make Blast Database

### **Description**

Generates a BLAST database to be queried. Required for identifying sequences using the BLAST+ software.

### Usage

```
makeBlastDb(fastaFile, dbPath, blastPath, dbType = "nucl")
```

### **Arguments**

fastaFile FASTA file containing sequences to generate a BLAST database from.

dbPath Path to save the BLAST database to.

blastPath Path/name of BLAST program to use. Name of the application for Linux/MacOS,

absolute path for the executable for windows users.

dbType Type of BLAST database to create, e.g. "nucl" for a nucleotide database.

### Value

No return value; generates a blast database in the chosen directory.

### Author(s)

Jack Gisby

### References

For further information, see the NCBI BLAST+ application documentation and help pages (https://www.ncbi.nlm.nih.gov/pul

### See Also

packSearch

18 packAlign

### **Examples**

```
## Not run:
makeBlastDb("genes.fasta", "blastdb.db", "C:/blast.exe")
## End(Not run)
```

packAlign

Global Alignment with VSEARCH

### **Description**

A global pairwise alignment of pack-TYPE elements by sequence similarity. mIt may be useful to run packClust to identify groups of similar transposable elements, before generating alignments of each group.

### Usage

```
packAlign(
  packMatches,
  Genome,
  identity = 0,
  threads = 1,
  identityDefinition = 2,
  maxWildcards = 0.05,
  saveFolder,
  vSearchPath = "vsearch"
)
```

### **Arguments**

packMatches A dataframe of potential Pack-TYPE transposable elements, in the format given

by packSearch. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-

object to a dataframe: data.frame(GRanges). Will be saved as a FASTA file

for VSEARCH.

Genome A DNAStringSet object containing sequences referred to in packMatches (the

object originally used to predict the transposons packSearch).

identity The sequence identity of two transposable elements in packMatches required to

be grouped into a cluster.

threads The number of threads to be used by VSEARCH.

identityDefinition

The pairwise identity definition used by VSEARCH. Defaults to 2, the standard

VSEARCH definition.

maxWildcards The maximal allowable proportion of wildcards in the sequence of each match

(defaults to 0.05).

packAlign 19

saveFolder The folder to save saveFolder files (uc, blast6out, FASTA)

vSearchPath When the package is run on windows systems, the location of the VSEARCH

executable file must be given; this should be left as default on Linux/MacOS

systems.

#### Value

Saves alignment information, including a uc, blast6out and a pairwise alignment fasta file, to the specified location. Returns the uc summary file generated by the alignment.

#### Note

In order to align sequences using VSEARCH, the executable file must first be installed.

### Author(s)

Jack Gisby

### References

VSEARCH may be downloaded from https://github.com/torognes/vsearch, along with a manual documenting the program's parameters. See https://www.ncbi.nlm.nih.gov/pubmed/27781170 for further information.

### See Also

tirClust, packClust, readBlast, readUc, filterWildcards, packSearch

20 packBlast

packBlast

Pipeline for BLAST/Classification of PackTYPE Elements

### **Description**

Run BLAST against user-specified databases of non-transposon and transposon-relates proteins. Can be used to classify transposons based on their internal sequences.

### Usage

```
packBlast(
  packMatches,
  Genome,
 blastPath,
  protDb,
  autoDb,
 minE = 0.001,
 blastTask = "blastn-short",
 maxHits = 100,
  threads = 1,
  saveFolder = NULL,
  tirCutoff = 100,
  autoCutoff = 1e-05,
  autoLength = 150,
  autoIdentity = 70,
  autoScope = NULL,
  protCutoff = 1e-05,
  protLength = 250,
 protIdentity = 70,
  protScope = 0.3
)
```

### **Arguments**

packMatches	A dataframe of potential Pack-	TYPE transposable elements,	in the format given
-------------	--------------------------------	-----------------------------	---------------------

by packSearch. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-

object to a dataframe: data.frame(GRanges). Will be saved as a FASTA file

for VSEARCH.

Genome A DNAStringSet object containing sequences referred to in packMatches (the

object originally used to predict the transposons packSearch).

blastPath Path to the BLAST+ executable, or name of the BLAST+ application for Linux/MacOS

users.

protDb For assigning Pack-TYPE elements. Path to the blast database containing nu-

cleotide or protein sequences to be matched against internal transposon sequences.

Can be generated using BLAST+, or with link{makeBlastDb}.

packBlast 21

autoDb	For assigning autonomous elements. Path to the blast database containing nucleotide or protein sequences to be matched against internal transposon sequences. Can be generated using BLAST+, or with link{makeBlastDb}.
minE	Blast results with e values greater than the specified cutoff will be ignored. This will be passed to BLASTN and applied to both transposon and non-transposon matches.
blastTask	Type of BLAST+ task, defaults to "blastn-short".
maxHits	Maximum hits returned by BLAST+ per query.
threads	Allowable number of threads to be utilised by BLAST+.
saveFolder	Directory to save BLAST+ results in; defaults to the working directory.
tirCutoff	How many bases to ignore at the terminal ends of the transposons to prevent hits to TIR sequences.
autoCutoff	Blast results for transposon-related elements will be filtered to ignore those with e values above the specified cutoff.
autoLength	Blast results for transposon-related elements containing hits with alignment lengths lower than this value will be ignored
autoIdentity	Blast results for transposon-related elements containing hits with sequence identities lower than this value will be ignored
autoScope	If specified, transposon-related blast results below the specified value will be ignored. Note that the dataframe of transposon matches must also be supplied to calculate scope. Scope is the proportion of the transposon's internal sequence occupied by the BLAST hit.
protCutoff	Blast results for genic/other matches will be filtered to ignore those with e values above the specified cutoff.
protLength	Blast results for genic/other matches containing hits with alignment lengths lower than this value will be ignored
protIdentity	Blast results for genic/other matches containing hits with sequence identities lower than this value will be ignored
protScope	If specified, genic/other blast matches below the specified value will be ignored. Note that the dataframe of transposon matches must also be supplied to calculate scope. Scope is the proportion of the transposon's internal sequence occupied by the BLAST hit.

### Value

Returns the original packMatches dataframe, with the addition of a "classification" column containing one of the following values:

- auto elements that match known transposases or transposon-related proteins are classified as autonomous elements
- pack elements that match other proteins or genic sequences may be classified as Pack-TYPE elements
- other elements that generate no significant hits

22 packClust

### Author(s)

Jack Gisby

#### References

For further information, see the NCBI BLAST+ application documentation and help pages (https://www.ncbi.nlm.nih.gov/pul

### See Also

blastAnalysis, packSearch, readBlast, blastAnnotate

### **Examples**

```
## Not run:
packMatches <- data(packMatches)
Genome <- data(arabidopsisThalianaRefseq)

packBlast(packMatches, Genome,
    protDb = "C:/data/TAIR10_CDS",
    autoDb = "C:/data/TAIR10_transposons",
    blastPath = "C:/blast/bin/blastn.exe")

## End(Not run)</pre>
```

packClust

Cluster Transposons with VSEARCH

### Description

Cluster potential pack-TYPE elements by sequence similarity. Resulting groups may be aligned with packAlign, or the clusters may be analysed with tirClust

```
packClust(
  packMatches,
  Genome,
  identity = 0.6,
  threads = 1,
  identityDefinition = 2,
  maxWildcards = 0.05,
  strand = "both",
  saveFolder = NULL,
  vSearchPath = "vsearch"
)
```

packClust 23

#### **Arguments**

packMatches A dataframe of potential Pack-TYPE transposable elements, in the format given

by packSearch. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-

object to a dataframe: data.frame(GRanges). Will be saved as a FASTA file

for VSEARCH.

Genome A DNAStringSet object containing sequences referred to in packMatches (the

object originally used to predict the transposons packSearch).

identity The sequence identity of two transposable elements in packMatches required to

be grouped into a cluster.

threads The number of threads to be used by VSEARCH.

identityDefinition

The pairwise identity definition used by VSEARCH. Defaults to 2, the standard

VSEARCH definition.

maxWildcards The maximal allowable proportion of wildcards in the sequence of each match

(defaults to 0.05).

strand The strand direction (+, - or \*) to be clustered.

saveFolder The folder to save output files (uc, blast6out, FASTA)

vSearchPath When the package is run on windows systems, the location of the VSEARCH

executable file must be given; this should be left as default on Linux/MacOS

systems.

#### Value

Saves cluster information, including a uc and blast6out file, to the specified location. Returns the given packMatches dataframe with an additional column, cluster, containing cluster IDs.

#### Note

In order to cluster sequences using VSEARCH, the executable file must first be installed.

### Author(s)

Jack Gisby

#### References

VSEARCH may be downloaded from https://github.com/torognes/vsearch. See https://www.ncbi.nlm.nih.gov/pubmed/27781170 for further information.

#### See Also

tirClust, packAlign, readBlast, readUc, filterWildcards, packSearch

24 packFinder

### **Examples**

packFinder

packFinder: a package for the de novo Annotation of Pack-TYPE Transposable Elements

### **Description**

Algorithm and tools for in silico pack-TYPE transposon discovery. Filters a given genome for properties unique to DNA transposons and provides tools for the investigation of returned matches.

### **Main Algorithm**

The goal of packFinder was to implement a simple tool for the prediction of potential Pack-TYPE elements. packFinder uses the following prior knowledge, provided by the user, to detect transposons:

- Terminal Inverted Repeat (TIR) Base Sequence
- Length of Terminal Site Duplication (TSD)
- · Length of the Transposon

These features provide enough information to detect autonomous and pack-TYPE elements. For a transposon to be predicted by packFinder its TSD sequences must be identical to each other, its forward TIR sequence must match the base sequence provided and its reverse TIR sequence must match its reverse complement.

Transposons are therefore predicted by searching a given genome for these characteristics, and further analysis steps can reveal the nature of these elements - while the packFinder tool is sensitive for the detection of transposons, it does not discriminate between autonomous and Pack-TYPE elements. Autonomous elements will contain a transposase gene within the terminal inverted repeats and tend to be larger than their Pack-TYPE counterparts; pack-TYPE elements instead capture sections of host genomes. Following cluster analysis, BLAST can be used to discern which predicted elements are autonomous (transposase-containing) and with are true Pack-TYPE elements.

packMatches 25

### Workflow

An example of a standard workflow can be found using browseVignettes(package = "packFinder"). The primary functions include:

- packSearch the packSearch algorithm uses simple pattern matching to detect DNA transposons.
- packClust VSEARCH is used for clustering elements based on sequence similarity.

Having obtained the sequences of transposable elements in a given genome, it is recommende to carry out a BLAST search for each transposon cluster. This can identify which elements are likely autonomous, and which may be Pack-TYPE.

The packFinder functions report the position of elements in a given genome using a dataframe in the format of packMatches. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-class] object to a dataframe: data.frame(GRanges).

### Author(s)

Jack Gisby

#### See Also

packSearch

packMatches

Sample packFinder Output

### **Description**

A sample output from packSearch with cluster information. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-class]{GRanges} object to a dataframe: data.frame(GRanges).

#### Usage

data(packMatches)

### **Format**

A dataframe of 9 obs. and 7 variables.

26 packSearch

### **Details**

Was obtained from running packSearch on the Arabidopsis thaliana chromosome 3 reference sequence, followed by clustering using packClust. Contains the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

The dataset was generated as in the example below.

#### See Also

```
packSearch, data.frame, arabidopsisThalianaRefseq
```

### **Examples**

```
data(arabidopsisThalianaRefseq)

packMatches <- packSearch(
    Biostrings::DNAString("CACTACAA"),
    arabidopsisThalianaRefseq,
    elementLength = c(300, 3500),
    tsdLength = 3
)</pre>
```

packSearch

packFinder Algorithm Pipeline

### **Description**

General use pipeline function for the Pack-TYPE transposon finding algorithm.

```
packSearch(
   tirSeq,
   Genome,
   mismatch = 0,
   elementLength,
   tsdLength,
   tsdMismatch = 0,
   fixed = TRUE
)
```

packSearch 27

#### **Arguments**

tirSeq A DNAString object containing the TIR sequence to be searched for.

Genome A DNAStringSet object to be searched.

mismatch The maximum edit distance to be considered for TIR matches (indels + substi-

tions). See matchPattern for details.

elementLength The maximum element length to be considered, as a vector of two integers. E.g.

c(300, 3500)

tsdLength Integer referring to the length of the flanking TSD region.

tsdMismatch An integer referring to the allowable mismatch (substitutions or indels) between

a transposon's TSD sequences. matchPattern from Biostrings is used for pat-

tern matching.

fixed Logical that will be passed to the 'fixed' argument of matchPattern. Deter-

mines the behaviour of IUPAC ambiguity codes when searching for TIR se-

quences.

#### **Details**

Finds potential pack-TYPE elements based on:

• Similarity of TIR sequence to tirSeq

· Proximity of potential TIR sequences

· Directionality of TIR sequences

· Similarity of TSD sequences

The algorithm finds potential forward and reverse TIR sequences using identifyTirMatches and their associated TSD sequence via getTsds. The main filtering stage, identifyPotentialPackElements, filters matches to obtain a dataframe of potential PACK elements. Note that this pipeline does not consider the possibility of discovered elements being autonomous elements, so it is recommended to cluster and/or BLAST elements for further analysis. Furthermore, only exact TSD matches are considered, so supplying long sequences for TSD elements may lead to false-negative results.

#### Value

A dataframe, containing elements identified by the algorithm. These may be autonomous or pack-TYPE elements. Will contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.
- width the width of the predicted element.
- strand the strand direction of the transposable element. This will be set to "\*" as the
  packSearch function does not consider transposons to have a direction only TIR sequences.
  Passing the packMatches dataframe to packClust will assign a direction to each predicted
  Pack-TYPE element.

28 packsToCsv

This dataframe is in the format produced by coercing a link[GenomicRanges: GRanges-class]{GRanges} object to a dataframe: data.frame(GRanges). Downstream functions, such as packClust, use this dataframe to manipulate predicted transposable elements.

#### Note

This algorithm does not consider:

- Autonomous elements autonomous elements will be predicted by this algorithm as there
  is no BLAST step. It is recommended that, after clustering elements using packClust, the
  user analyses each group to determine which predicted elements are autonomous and which
  are likely Pack-TYPE elements. Alternatively, databases such as Repbase (https://www.
  girinst.org/repbase/) supply annotations for autonomous transposable elements that can
  be used to filter autonomous matches.
- TSD Mismatches if two TIRs do not have exact matches for their terminal site duplications they will be ignored. Supplying longer TSD sequences will likely lead to a lower false-positive rate, however may also cause a greater rate of false-negative results.

Pattern matching is done via matchPattern.

### Author(s)

Jack Gisby

### See Also

identifyTirMatches, getTsds, identifyPotentialPackElements, packClust, packMatches, DNAStringSet, DNAString, matchPattern

### **Examples**

```
data(arabidopsisThalianaRefseq)

packMatches <- packSearch(
    Biostrings::DNAString("CACTACAA"),
    arabidopsisThalianaRefseq,
    elementLength = c(300, 3500),
    tsdLength = 3
)</pre>
```

packsToCsv

Save packFinder Results in CSV Format (.csv)

#### **Description**

Saves a dataframe of potential Pack-TYPE elements, usually generated via packSearch. May be retrieved using getPacksFromCsv.

packsToFasta 29

### Usage

```
packsToCsv(packMatches, file)
```

### **Arguments**

packMatches

A dataframe containing genomic ranges and names referring to sequences to be extracted. Can be obtained from packSearch or generated from a GRanges object, after conversion to a dataframe. Must contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

file CSV file save path.

### Value

Save location of csv file.

#### Author(s)

Jack Gisby

#### See Also

```
getPacksFromCsv, write.table, packSearch
```

### **Examples**

```
data(packMatches)

packsToCsv(
    packMatches,
    system.file("extdata", "packMatches.csv", package = "packFinder")
)
```

packsToFasta

Save packFinder Results in FASTA Format (.fasta)

### Description

Saves a dataframe of potential Pack-TYPE elements, usually generated via packSearch. May be retrieved using getPacksFromFasta.

```
packsToFasta(packMatches, file, Genome)
```

30 packsToGRanges

### **Arguments**

packMatches

taframe containing genomic ranges and names referring to sequences to be extracted. Can be obtained from packSearch or generated from a GRanges object, after conversion to a dataframe. Must contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

file FASTA file save path.

Genome

A DNAStringSet object containing sequences referred to in packMatches (the object originally used to predict the transposons packSearch).

#### Value

Save location of Fasta file.

### Author(s)

Jack Gisby

#### See Also

getPacksFromFasta, packSearch

### **Examples**

```
data(arabidopsisThalianaRefseq)
data(packMatches)

packsToFasta(
    packMatches,
    system.file("extdata", "packMatches.fasta", package = "packFinder"),
    arabidopsisThalianaRefseq
)
```

packsToGRanges

Export packFinder Results to a GRanges Object

### **Description**

A dataframe containing genomic ranges and names referring to sequences to be extracted, likely obtained from packSearch, can be converted to a GRanges object. Can be converted back to a dataframe using getPacksFromGRanges. Additional features, such as clusters and TSD sequences, will be included in the object as metadata columns.

readBlast 31

### Usage

```
packsToGRanges(packMatches)
```

#### **Arguments**

packMatches

A dataframe containing genomic ranges and names referring to sequences to be extracted. Can be obtained from packSearch or generated from a GRanges object, after conversion to a dataframe. Must contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

#### Value

A GRanges object containing the ranges contained in packMatches and additional metadata columns. May be easily converted between dataframe and GRanges format for use in the packFinder package and link[GenomicRanges:GRanges-class]{GRanges} package. Note that most functions in the packFinder package require sequence ranges to be provided in dataframe format.

#### Author(s)

Jack Gisby

### See Also

```
getPacksFromGRanges, link[GenomicRanges:GRanges-class]{GRanges}
```

### **Examples**

```
data(packMatches)
packGRanges <- packsToGRanges(packMatches)</pre>
```

readBlast

Convert NCBI BLAST+ Files to Dataframe

### **Description**

Reads .blast6out files (NCBI Blast Format) generated by the VSEARCH clustering and alignment algorithms.

32 readBlast

### Usage

```
readBlast(
  file,
  minE = 1,
  length = 0,
  identity = 0,
  removeExactMatches = FALSE,
  scope = NULL,
  packMatches = NULL
)
```

### **Arguments**

file The file path of the blast file.

minE Blast results with e values greater than the specified cutoff will be ignored.

length Blast results alignment lengths lower below this value will be ignored

identity Blast results with target sequence identities below this value will be ignored.

removeExactMatches

If true, matches with 100 be ignored to prevent self-hits.

scope

If specified, blast results below the specified value will be ignored. Note that the dataframe of transposon matches must also be supplied to calculate scope. Scope is the proportion of the transposon's internal sequence occupied by the BLAST hit.

packMatches

taframe containing genomic ranges and names referring to sequences to be extracted. Can be obtained from packSearch or generated from a GRanges object, after conversion to a dataframe. Must contain the following features:

- start the predicted element's start base sequence position.
- end the predicted element's end base sequence position.
- seqnames character string referring to the sequence name in Genome to which start and end refer to.

### **Details**

blast6out file is tab-separated text file compatible with NCBI BLAST m8 and NCBI BLAST+ outfmt 6 formats. One cluster/alignment can be found for each line.

#### Value

A dataframe containing the converted .blast6out file. The file contains the following features:

- · Query sequence ID
- Target sequence ID
- Percenty sequence identity
- Alignment length
- Number of mismatches

readUc 33

- · Number of gaps
- Base position of alignment start in query sequence
- · Base position of alignment end in query sequence
- Base position of alignment start in target sequence
- Base position of alignment end in target sequence
- E-value
- Bit score

### Author(s)

Jack Gisby

### References

For further information, see the NCBI BLAST+ application documentation and help pages (https://www.ncbi.nlm.nih.gov/pub VSEARCH may be downloaded from https://github.com/torognes/vsearch; see https://www.ncbi.nlm.nih.gov/pubmed/27781170 for further information.

### See Also

codeblastAnalysis, codeblastAnnotate, codepackAlign, codereadUc, codepackClust

### **Examples**

```
readBlast(system.file(
   "extdata",
   "packMatches.blast6out",
   package = "packFinder"
))
```

readUc

Convert .uc Files to Dataframe

### **Description**

Reads .uc files (USEARCH Cluster Format) generated by the VSEARCH clustering and alignment algorithms.

```
readUc(file, output = "cluster")
```

34 readUc

### Arguments

file The file path of the .uc file.

output The type of analysis that was carried out to produce the .uc file.

• If output is specified as "cluster", VSEARCH clustering was carried out.

 If output is specified as "alignment", VSEARCH pairwise global alignment was carried out.

Note that clustering produces one "H" record for each sequence, and one "C" record for each cluster, while an alignment produces an "H" record for each alignment (see details).

#### **Details**

USEARCH cluster format is a tab separated text file that contains clustering and/or alignment information for a set of sequences. For each sequence a record type, "H, C or N", is provided providing information about the type of "hit" in the dataframe. These refer to:

- H Hit for alignments, indicates an identified alignment of two supplied sequences. For clustering, indicates the cluster assignment for a query.
- C Cluster record a record for each cluster generated.
- N No hit indicates that no cluster was assigned or no alignment was found with a target sequence. For clustering, a query with no hits becomes the centroid of a new cluster.

Additionally, for each record a "compressed alignment" is generated. This is the alignment represented in a compact format including the letters "M", "D", and "I". Before each letter, the number of consecutive columns of the given letter type is also given. The letter types are as follows:

- "M" Match Identical bases between the query and target sequence
- "D" Deletion A gap in the target sequence
- "I" Insertion A gap in the query sequence

An example of this would be "13M", referring to 13 consecutive matches between the query and target sequence.

#### Value

A dataframe containing the converted .uc file. The fields contained within are as follows:

- Record type "H, C or N", see details for further information.
- Cluster designation (output = "cluster" only)
- Sequence length, or cluster size
- Percent identity to target
- The nucleotide strand (output = "cluster" only)
- A compressed alignment see details for further information.
- ID of query sequence
- ID of target sequence ("H" records only)

tirClust 35

### Author(s)

Jack Gisby

#### References

VSEARCH may be downloaded from https://github.com/torognes/vsearch. See https://www.ncbi.nlm.nih.gov/pubmed/27781170 for further information.

### See Also

codetirClust, codepackAlign, codereadBlast, codepackClust

### **Examples**

```
readUc(system.file(
   "extdata",
   "packMatches.uc",
   package = "packFinder"
))
```

tirClust

Analyse TIR Sequences of Pre-clustered Transposable Elements

### **Description**

Takes transposable elements clustered by VSEARCH, packClust, and produces consensus sequences for the terminal inverted repeats of each. Allows for the visualisation of TIR similarities between clusters for both forward and reverse strands.

#### Usage

```
tirClust(
  packMatches,
  Genome,
  tirLength = 25,
  plot = TRUE,
  plotSavePath = NULL,
  k = 5,
  output = "consensus"
)
```

### Arguments

packMatches

A dataframe containing genomic ranges and names referring to sequences to be extracted. This dataframe is in the format produced by coercing a link[GenomicRanges:GRanges-class object to a dataframe: data.frame(GRanges).

Must contain the following features:

36 tirClust

• start - the predicted element's start base sequence position.

• end - the predicted element's end base sequence position.

• seqnames - character string referring to the sequence name in Genome to

which start and end refer to.

A DNAStringSet object containing sequences referred to in packMatches (the

object originally used to predict the transposons packSearch).

tirLength The TIR size to be considered. Consensus sequences will be generated based on

the first and last tirLength bases of a transposon.

plot Argument specifying whether the TIR consensus sequences should be plottted

as a dendrogram.

plotSavePath File path for the dendrogram plot. If unspecified, the dendrogram plot is not

saved.

k The k-mer size to be used for calculating a distance matrix between TIR con-

sensus sequences. See kdistance. Larger word sizes will not be suitable for longer TIR sequences, due to processing time required. Additionally, k must be

greater than the TIR sequence length.

output Controls the output of tirClust. If output is specified as "consensus", the con-

sensus sequences of each TIR cluster will be returned; else, if output is specified as "dendrogram", a dendrogram object will be returned for creation of customis-

able plots.

#### Value

Genome

If output is specified as "consensus" (default), returns a list of consensus sequences for each cluster specified in packMatches as a DNAStringSet. Else if output is specified as "dendrogram", returns a dendrogram object used to create hierarchical clustering diagrams.

### Author(s)

Jack Gisby

### See Also

codepackClust, codepackAlign, kdistance, DNAStringSet, as.alignment, packSearch

### **Examples**

data(arabidopsisThalianaRefseq)
data(packMatches)

tirClust(packMatches, arabidopsisThalianaRefseq)

# **Index**

```
* datasets
                                                    packsToCsv, 10, 28
    arabidopsis Thaliana Refseq, 2
                                                    packsToFasta, 11, 29
    packMatches, 25
                                                    packsToGRanges, 12, 30
arabidopsisThalianaRefseq, 2, 26
                                                    read.table, 10
as.alignment, 36
                                                    readBlast, 4, 6, 19, 22, 23, 31, 35
                                                    readUc, 19, 23, 33, 33
blastAnalysis, 3, 5, 6, 22, 33
blastAnnotate, 4, 5, 22, 33
                                                    tirClust, 19, 22, 23, 35, 35
collapseSeqs, 6
                                                    write.table, 29
data.frame, 26
DNAString, 2, 3, 9, 16, 27, 28
DNAStringSet, 2, 3, 9, 13, 15, 16, 27, 28, 36
filterWildcards, 7, 19, 23
getGenome, 3
getPackSeqs, 8
getPacksFromCsv, 10, 28, 29
getPacksFromFasta, 11, 29, 30
getPacksFromGRanges, 12, 30, 31
getTsds, 13, 14, 27, 28
GRanges, 29–32
identifyPotentialPackElements, 14, 27,
identifyTirMatches, 14, 15, 27, 28
kdistance, 36
makeBlastDb, 17
matchPattern, 14, 16, 27, 28
packAlign, 7, 8, 18, 22, 23, 33, 35, 36
packBlast, 4, 6, 20
packClust, 7, 8, 18, 19, 22, 26–28, 33, 35, 36
packFinder, 24
packMatches, 25, 25, 28
packSearch, 3-14, 16-20, 22, 23, 25, 26, 26,
         28–32, 36
```