

Package ‘DominoEffect’

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Type Package

Title Identification and Annotation of Protein Hotspot Residues

Version 1.33.0

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Description The functions support identification and annotation of hotspot residues in proteins. These are individual amino acids that accumulate mutations at a much higher rate than their surrounding regions.

License GPL (>= 3)

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LazyData true

Depends R(>= 3.5)

Imports biomaRt, data.table, utils, stats, Biostrings, palign, SummarizedExperiment, VariantAnnotation, AnnotationDbi, Seqinfo, IRanges, GenomicRanges, methods

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align_to_unip	<i>Align protein segment around the hotspot to the UniProt/Swiss-Prot KB sequence.</i>
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Description

This function aligns the Ensembl protein region with a hotspot to the UniProt sequence. The Ensembl region encompasses 15 amino acids where the hotspot is in the middle. If the hotspot was at the protein start or end the region is still 15 amino acids long, but the hotspot position is shifted.

Usage

```
align_to_unip(ens.seq, uni.seq, ensembl_mut_position)
```

Arguments

ens.seq	AAString object with the Ensembl protein sequence corresponding to the representative transcript.
uni.seq	AAString with the UniProt sequence for the identifier matching the Ensembl gene name.
ensembl_mut_position	Numeric vector defining the hotspot position in the Ensembl sequence, i.e. in the ens.seq

Value

Returns a list where the first element is a character vector defining whether there was a significant alignment and the second element provides the hotspot position in the UniProt sequence.

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Examples

```
library(Biostrings)

ens.seq <- AAString("MDLSALREEVQNVINAMQKILECPICLELIKEPVSTKCDHIFCKFCMLK")
uni.seq <- AAString("MDLSALRVEEVQNVINAMQKILECPICLELIKEPVSTKCDHIFCKFCMLA")
ensembl_mut_position <- 25

align_to_unip(ens.seq, uni.seq, ensembl_mut_position)
```

calculate_boundary *calculate_boundary*

Description

The function calculates boundaries of sequence windows around the mutation. It is possible to define up to two window lengths. If the mutation is close to the start or end of the protein, the region is extended into the other direction to keep the indicated size

Usage

```
calculate_boundary(mut_pos_numeric, length_aa, flanking_region)
```

Arguments

mut_pos_numeric	
length_aa	Amino acid position of mutation
flanking_region	Length of transcript in amino acids
	Vector containing two flanking regions

Value

returns a list with the boundaries for the two regions

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Examples

```
calculate_boundary(250, 500, c(200, 300))
calculate_boundary(250, 500, 300)
```

DominoData *Sample data*

Description

Sample Data

Author(s)

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DominoEffect

*Identification of significant mutation hotspot residues.***Description**

The function identifies individual amino acid residues, which accumulate a high fraction of the overall mutation load within a protein. After detecting mutation hotspots, the function obtains UniProt/Swiss-Prot KB functional and structural annotations for the affected protein regions and checks for the sequence agreement.

Usage

```
DominoEffect(mutation_dataset, gene_data, snp_data,
min_n_muts = 5, MAF_thresh = 0.01,
flanking_region = c(200, 300),
poisson.thr = 0.01, percentage.thr = 0.15,
ratio.thr = 45, approach = "percentage", write_to_file = "NO",
ens_release = "https://feb2023.archive.ensembl.org")
```

Arguments

mutation_dataset	Object containing a table with the mutation data (e.g. TCGA point mutations mapped to protein level).
gene_data	DominoData object containing information about Ensembl gene annotations: gene identifiers and representative transcript cDNA length.
snp_data	Object containing a table with information on population SNPs.
min_n_muts	Numeric vector defining a minimum number of mutations that need to occur at the same residue. Default: 5
MAF_thresh	Numeric vector that defines Minor allele frequency threshold for considering reported mutations as population SNPs.
flanking_region	Numeric vector that defines size of a window around the mutation that will be considered. Up to two window sizes are allowed.
poisson.thr	Numeric vector that defines a threshold for the adjusted p-value. Residues with an associated p-value that is lower than the defined value are reported. Default: 0.01
percentage.thr	Number defining the fraction of mutations within the window that need to fall on a single residue in order for it to be classified as a hotspot. Default: 0.15
ratio.thr	Number defining a requirement that a number of mutations on a single residue should exceed what would be expected by chance given a background mutation rate in the window (i.e. the surrounding region). Default: 45
approach	Option to define selection criteria to use percentage.thr or ratio.thr as criterion for finding single residue mutation clusters. Options: "both", "percentage" or "ratio". Default = "percentage"
write_to_file	Option if the identified and annotated hotspots should be written to a file (YES or NO). Default: NO
ens_release	Release of ensembl to be used. Default: https://feb2023.archive.ensembl.org

Value

Results table

Author(s)

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Examples

```
data("SnpData", package = "DominoEffect")
data("TestData", package = "DominoEffect")
data("DominoData", package = "DominoEffect")

hotspot_mutations <- DominoEffect(mutation_dataset = TestData,
gene_data = DominoData, snp_data = SnpData)
```

GPo_of_hotspots	<i>Converts hotspot mutation table into a GPo object</i>
-----------------	--

Description

This function converts the genomic information on hotspot mutations into a GPo object.

Usage

```
GPo_of_hotspots(hotspot_mutations)
```

Arguments

hotspot_mutations
Data frame with information on hotspot mutations generated by the DominoEffect package.

Value

GPo object that contains the genomic information on hotspot mutations.

Author(s)

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Examples

```
data("SnpData", package = "DominoEffect")
data("TestData", package = "DominoEffect")
data("DominoData", package = "DominoEffect")

hotspot_mutations <- DominoEffect(mutation_dataset = TestData,
gene_data = DominoData, snp_data = SnpData)
GPo_of_hotspots(hotspot_mutations)
```

identify_hotspots *Identify hotspots*

Description

The function identify protein hotspot mutation residues

Usage

```
identify_hotspots(mutation_dataset, gene_data,
  snp_data, min_n_muts = 5, MAF_thresh = 0.01, flanking_region = c(200, 300),
  poisson.thr = 0.01, percentage.thr = 0.15, ratio.thr = 45, approach = "percentage")
```

Arguments

mutation_dataset	Object containing a table with the mutation data (e.g. TCGA point mutations mapped to protein level).
gene_data	Data frame or Txdb object containing information about Ensembl gene annotations: gene identifiers and representative transcript cDNA length.
snp_data	Object containing a table or vcf object with information on population SNPs.
min_n_muts	Numeric vector defining a minimum number of mutations that need to occur at the same residue. Default: 5
MAF_thresh	Numeric vector that defines Minor allele frequency threshold for considering reported mutations as population SNPs.
flanking_region	Numeric vector that defines size of a window around the mutation that will be considered. Up to two window sizes are allowed.
poisson.thr	Numeric vector that defines a threshold for the adjusted p-value. Residues with an associated p-value that is lower than the defined value are reported. Default: 0.01
percentage.thr	Number defining the fraction of mutations within the window that need to fall on a single residue in order for it to be classified as a hotspot. Default: 0.15
ratio.thr	Number defining a requirement that a number of mutations on a single residue should exceed what would be expected by chance given a background mutation rate in the window (i.e. the surrounding region). Default: 45
approach	Option to define selection criteria to use percentage.thr or ratio.thr as criterion for finding single residue mutation clusters. Options: "both", "percentage" or "ratio". Default = "percentage"

Value

An object containing information on the significant hotspots, associated Gene and protein identifiers, number of mutations, percentage of mutations within defined windows that map to the same residue and associated p-values.

Author(s)

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Examples

```
data("SnpData", package = "DominoEffect")
data("TestData", package = "DominoEffect")
data("DominoData", package = "DominoEffect")
hotspot_mutations <- identify_hotspots(mutation_dataset = TestData,
  gene_data = DominoData, snp_data = SnpData)
```

import_txdb	<i>Imports txdb data and converts it into format required for DominoEffect package</i>
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Description

This function imports txdb data and converts into a data frame used in the DominoEffect package only extracting the required information from the txdb object.

Usage

```
import_txdb(txdb_object)
```

Arguments

txdb_object TxDB Object, e.g. from makeTxDbFromEnsembl

Value

Data frame that can be used in DominoEffect package.

Author(s)

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Examples

```
#EnsTxDB <- makeTxDbFromEnsembl(organism="Homo sapiens", release=73,
#                               server="ensemldb.ensembl.org")
#DominoData <- import_txdb(EnsTxDB)
#head(DominoData)
```

map_to_func_elem	<i>Functional annotation of significant hotspot residues.</i>
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Description

This function retrieves Uniprot annotations for the functional elements in the proteins with significant hotspots and overlaps and maps the hotspot residues to these.

Usage

```
map_to_func_elem(hotspot_results, write_to_file = "NO", ens_release = "109")
```

Arguments

hotspot_results	Object containing information about the hotspot residues identified using the function identify_hotspots().
write_to_file	A character vector defining whether the resulting annotated hotspots should be saved in a file (YES or NO).
ens_release	A character vector defining whether the default gene annotations are used, i.e. Ensembl release 109, or if the gene_data correspond to a different Ensembl release. For the current Ensembl version this should be set to: ens_release="www.ensembl.org". For the archive versions to the corresponding archive website.

Value

Updated results file containing an additional column with the information on the annotated functional and structural region within which the mutation is mapped.

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Examples

```
data("TestData", package = "DominoEffect")
data("DominoData", package = "DominoEffect")
data("SnpData", package = "DominoEffect")

hotspot_mutations <- identify_hotspots(TestData, DominoData, SnpData)
hotspot_mutations <- map_to_func_elem(hotspot_mutations,
write_to_file = "NO", ens_release = "109")

head(hotspot_mutations)
```

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