Package 'rfPred'

October 24, 2025
Type Package
Title Assign rfPred functional prediction scores to a missense variants list
Version 1.47.1
Date 2025-07-23
Author Fabienne Jabot-Hanin, Hugo Varet and Jean-Philippe Jais
Depends R ($>= 3.5.0$), methods
Imports utils, Seqinfo, data.table, IRanges, GenomicRanges, parallel, Rsamtools
Suggests BiocStyle
Maintainer Hugo Varet <varethugo@gmail.com></varethugo@gmail.com>
Description Based on external numerous data files where rfPred scores are pre-calculated on all genomic positions of the human exome, the package gives rfPred scores to missense variants identified by the chromosome, the position (hg19 version), the referent and alternative nucleotids and the uniprot identifier of the protein. Note that for using the package, the user has to download the TabixFile and index (approximately 3.3 Go).
License GPL (>=2)
Encoding UTF-8
biocViews Software, Annotation, Classification
RoxygenNote 7.3.2
git_url https://git.bioconductor.org/packages/rfPred
git_branch devel
git_last_commit 330c58d
git_last_commit_date 2025-07-23
Repository Bioconductor 3.22
Date/Publication 2025-10-23
Contents
rfPred-package
1

2 example_GRanges

	rfPred_scores_mo	otor	 6
Index			8
rfPr	ed-package	Assign functional prediction rfPred scores to human missense (random forest method based on SIFT, Polyphen2, PhyloP, Mutation Taster)	

Description

The package provides a function which returns the rfPred score for a list of non-synonymous missense variants. All the rfPred scores are pre-calculated and stored in a TabixFile available on a server and which can be downloaded for using the package while not connected on the Internet. The package does not work without an access to the TabixFile. However, a toy example on the chromosome Y is available within the package to test the rfPred_scores function. curves with numbers of subjects at risk, compare data sets, display spaghetti-plot, build multi-contingency tables...

Author(s)

Fabienne Jabot-Hanin, Hugo Varet and Jean-Philippe Jais

References

dbNSFP database: Liu X, Jian X and Boerwinkle E. 2011. dbNSFP: a lightweight database of human non-synonymous SNPs and their functional predictions. Human Mutation. 32:894-899.

rfPred method: Jabot-Hanin F, Varet H, Tores F and Jais J-P. 2013. rfPred: a new meta-score for functional prediction of missense variants in human exome (submitted).

example_GRanges

Toy example of GRanges object

Description

Toy example of GRanges object

Format

A GRanges object with 11 rows and several columns:

seqnames Chromosome number (only Y in this example)
ranges IRanges object for which start=end: position on the chromosome
reference Referent nucleotid (A, C, G or T)
alteration Alteration nucleotid (A, C, G or T)

rfPred_scores 3

rfPred_scores

Assign functional prediction rfPred scores to human missense variants

Description

rfPred is a statistical method which combines 5 algorithms predictions in a random forest model: SIFT, Polyphen2, LRT, PhyloP and MutationTaster. These scores are available in the dbNFSP database for all the possible missense variants in hg19 version, and the package rfPred gives a composite score more reliable than each of the isolated algorithms.

Usage

```
rfPred_scores(
  variant_list,
  data = system.file("extdata/chrY_rfPred.txtz", package = "rfPred"),
  index = system.file("extdata/chrY_rfPred.txtz.tbi", package = "rfPred"),
  all.col = FALSE,
  file.export = NULL,
  n.cores = 1
)
rfPred_scores(variant_list,
                data=system.file("extdata/chrY_rfPred.txtz", package="rfPred"),
              index=system.file("extdata/chrY_rfPred.txtz.tbi", package="rfPred"),
                     all.col=FALSE, file.export=NULL, n.cores=1)
## S4 method for signature 'character'
rfPred_scores(
  variant_list,
  data = system.file("extdata/chrY_rfPred.txtz", package = "rfPred"),
  index = system.file("extdata/chrY_rfPred.txtz.tbi", package = "rfPred"),
  all.col = FALSE,
  file.export = NULL,
  n.cores = 1
)
## S4 method for signature 'GRanges'
rfPred_scores(
  variant_list,
  data = system.file("extdata/chrY_rfPred.txtz", package = "rfPred"),
  index = system.file("extdata/chrY_rfPred.txtz.tbi", package = "rfPred"),
  all.col = FALSE,
  file.export = NULL,
  n.cores = 1
)
```

Arguments

variant_list A variants list in a data.frame containing 4 or 5 columns: chromosome number, hg19 genomic position on the chromosome, reference nucleotid, variant nucleotid and uniprot protein identifier (optional); or a character string of the

4 rfPred_scores

path to a VCF (Variant Call Format) file; or a GRanges object with metadata containing textually reference, alteration and proteine (optional) columns

names for reference and alteration

data Path to the compressed TabixFile, either on the server (default) or on the user's

computer

index Path to the index of the TabixFile, either on the server (default) or on the user's

computer

all.col TRUE to return all available information, FALSE to return a more compact result

(the most informative columns, see Value)

file.export Optional, name of the CSV file in which export the results (default is NULL)

n. cores number of cores to use when scaning the TabixFile, can be efficient for large

request (default is 1)

Value

The variants list with the assigned rfPred scores, as well as the scores used to build rfPred meta-score: SIFT, phyloP, MutationTaster, LRT (transformed) and Polyphen2 (corresponding to Polyphen2_HVAR_score). The data frame returned contains these columns:

chromosome chromosome number

position_hg19 physical position on the chromosome as to hg19 (1-based coordinate)

reference reference nucleotide allele (as on the + strand)
alternation alternative nucleotide allele (as on the + strand)

proteine Uniprot accession number aaref reference amino acid alternative amino acid

aapos amino acid position as to the protein

rfPred_score rfPred score betwen 0 and 1 (higher it is, higher is the probability of pathogenic-

ity)

SIFT_score SIFT score between 0 and 1 (higher it is, higher is the probability of pathogenic-

ity contrary to the original SIFT score) = 1-original SIFT score

Polyphen2_score

Polyphen2 (HVAR one) score between 0 and 1, used to calculate rfPred (higher

it is, higher is the probability of pathogenicity)

MutationTaster_score

MutationTaster score between 0 and 1 (higher it is, higher is the probability of

pathogenicity)

PhyloP_score PhyloP score between 0 and 1 (higher it is, higher is the probability of pathogenic-

ity): PhyloP_score=1-0.5x10^phyloP if phyloP>0 or PhyloP_score=0.5x10^-

phyloP if phyloP<0

LRT score between 0 and 1 (higher it is, higher is the probability of pathogenic-

ity): LRT_score=1-LRToriginalx0.5 if LRT_Omega<1 or LRT_score=LRToriginalx0.5

if LRT_Omega>=1

The following columns are also returned if all.col is TRUE:

Uniprot_id Uniprot ID number

genename gene name

rfPred_scores 5

position_hg18 physical position on the chromosome as to hg18 (1-based coordinate) Polyphen2_HDIV_score

Polyphen2 score based on HumDiv, i.e. hdiv_prob. The score ranges from 0 to 1: the corresponding prediction is "probably damaging" if it is in [0.957,1]; "possibly damaging" if it is in [0.453,0.956]; "benign" if it is in [0,0.452]. Score cut-off for binary classification is 0.5, i.e. the prediction is "neutral" if the score is lower than 0.5 and "deleterious" if the score is higher than 0.5. Multiple entries separated by ";"

Polyphen2_HDIV_pred

Polyphen2 prediction based on HumDiv: D (probably damaging), P (possibly damaging) and B (benign). Multiple entries separated by ";"

Polyphen2_HVAR_score

Polyphen2 score based on HumVar, i.e. hvar_prob. The score ranges from 0 to 1, and the corresponding prediction is "probably damaging" if it is in [0.909,1]; "possibly damaging" if it is in [0.447,0.908]; "benign" if it is in [0,0.446]. Score cut-off for binary classification is 0.5, i.e. the prediction is "neutral" if the score is lower than 0.5 and "deleterious" if the score is higher than 0.5. Multiple entries separated by ";"

Polyphen2_HVAR_pred

Polyphen2 prediction based on HumVar: D (probably damaging), P (possibly damaging) and B (benign). Multiple entries separated by ";"

MutationTaster_pred

MutationTaster prediction: A (disease_causing_automatic), D (disease_causing), N (polymorphism) or P (polymorphism_automatic)

phyloP original phyloP score

LRT_Omega estimated nonsynonymous-to-synonymous-rate ratio
LRT_pred LRT prediction, D(eleterious), N(eutral) or U(nknown)

Author(s)

Fabienne Jabot-Hanin, Hugo Varet and Jean-Philippe Jais

References

Jabot-Hanin F, Varet H, Tores F and Jais J-P. 2013. rfPred: a new meta-score for functional prediction of missense variants in human exome (submitted).

Examples

6 rfPred_scores_motor

rfPred_scores_motor

Motor of rfPred_scores

Description

Motor of rfPred_scores

Usage

```
rfPred_scores_motor(variant_list, data, index, all.col, file.export, n.cores)
```

Arguments

variant_list	Variants list in a data.frame containing 4 or 5 columns: chromosome number, hg19 genomic position on the chromosome, reference nucleotid, variant nucleotid and uniprot protein identifier (optional)
data	Path to the compressed TabixFile, either on the server (default) or on the user's computer
index	Path to the index of the TabixFile, either on the server (default) or on the user's computer
all.col	TRUE to return all available information, FALSE to return a more compact result (the most informative columns, see Value)
file.export	Optional, name of the CSV file in which export the results (default is NULL)
n.cores	number of cores to use when scaning the TabixFile, can be efficient for large request (default is 1)

Value

see the rfPred_scores function

Note

This function is called by the rfPred_scores S4 method

variant_list_Y 7

variant_list_Y

Toy example of data.frame

Description

Toy example of data.frame

Format

A data frame with 5 observations on the following 5 variables:

chr Chromosome number (only Y in this example)

pos Position on the chromosome (numeric)

ref Referent nucleotid (A, C, G or T)

alt Alteration nucleotid (A, C, G or T)

uniprot Uniprot protein identifier (factor)

Index