Package 'epitopR'

August 19, 2022

Type Package

Title Predict Peptide-MHC Binding

Version 0.1.1

Description A suite of tools to predict peptide MHC (major histocompatibility complex) presentation in the context of both human and mouse. Polymorphic peptides between self and foreign proteins are identified. The ability of peptides to bind self MHC is assessed and scored. Based on half maximal inhibitory concentration as queried through the immune epitope database API <<u>http://tools.iedb.org/mhcii/></u> using user defined methods, the foreign peptides most likely to be presented are output along with their predicted binding strength, amino acid position, the protein from which each peptide was derived, and the presenting allele. ``References:'' Vita R, Mahajan S, Overton JA, Dhanda SK, Martini S, Cantrell JR, Wheeler DK, Sette A, Peters B. <<u>doi:10.1093/nar/gky1006></u>.

Depends R (>= 4.1.0)

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Encoding UTF-8

RoxygenNote 7.2.1

Suggests knitr, rmarkdown, testthat (>= 3.0.0)

VignetteBuilder knitr

Imports devtools, dplyr, usethis, rstudioapi, fs, here, httr, janitor, purrr, stringr, tibble, tidyverse, utils, readr, ggseqlogo, Biostrings, seqinr

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core_mut

Determine presence of mutation in core binding sequence

Description

The core_mut() function appends a new column to the peptide dataframe, identifying those that have a mutation in the core binding sequence. Since MHCII has an open binding pocket, it presents peptides that may be several amino acids longer than the core sequence pattern required to bind a particular MHC. In some cases, the user may want to filter their results, in order to keep only peptides with a mutation in the core binding sequence (when compared to the equivalent self peptide). In order to achieve this, the stimulating and self antigens are aligned using a multiple sequence alignment tool, from the bioconductor package msa. The sequence positions of the core in the stimulating peptide are determined, and sequences are kept only if there is a sequence difference between stimulating and self peptides at that position.

Usage

core_mut(dat_in, ag_stim, ag_self)

Arguments

dat_in	dataframe, output of mhcII_hu(). The stimulating peptide, core pattern, start, and end positions will be pulled from this dataframe.
ag_stim	string, amino acid sequence of the stimulating antigen, aligned with self
ag_self	string, amino acid sequence of the self antigen, aligned with stimulating

Value

data frame, peptide with flag of whether or not a mutation is in the core binding sequence

mhcII_hu

Description

It determines which non-self peptides can be presented by a given HLA class II allele. This function takes a sequence for a stimulating antigen and the corresponding self antigen, and given a defined sequence length, queries the IEDB API with the user's choice of peptide binding prediction method. The set of peptides present in the results for the stimulating antigen but not the self antigen are then carried forward as non-self peptides. If desired, the user can adjust the default thresholds (by IC50 binding affinity or percentile rank) used to define "strong" and "weak" binders. The output is a dataframe of non-self peptides that are predicted to bind to the presenting allele.

Usage

```
mhcII_hu(
    ag_present,
    ag_stim,
    ag_self,
    seq_len = "15",
    fd_out = as.character(paste0(tempdir(), "/", "outputs", "/")),
    method = "netmhciipan",
    cutoff_score = list(cutoff_netpan = c(50, 500), cutoff_comblib = c(50, 500),
        cutoff_nn_align = c(50, 500), cutoff_sturniolo = c(2), cutoff_el = c(2, 10)),
    cutoff_rank = c(2, 10),
    url_iedb = "http://tools-cluster-interface.iedb.org/tools_api/mhcii/"
)
```

Arguments

ag_present	character vector, presenting allele, formatted with either "_", "*", or":" separat- ing loci, antigen, and allele. For example, "DRB1_08_01".
ag_stim	character vector, stimulating antigen, can either be an HLA class II allele en- tered in the same format as ag_present, or a character vector of the amino acid sequence of the protein
ag_self	character, self antigen, can either be an HLA class II allele entered in the same format as ag_present, or a character vector of the amino acid sequence of the protein
seq_len	string, length of peptides to consider
fd_out	string, output folder name; default output is current working directory
method	string, IEDB prediction method to be used. Options are "netmhciipan', "netmhci- ipan_el" or "recommended." Default is netmhciipan.
cutoff_score	list of vectors. Defines the thresholds required to be included in results, and to be labeled, "strong" or "weak" binder. Multiple prediction methods are used, each of which provide different raw outputs (i.e. IC50, "strength", "score"). Our

	justification for the default thresholds is listed in the mhcII_hu vignette, however the user may choose to specify alternate cutoffs if desired.
cutoff_rank	vector, IEDB adjusts all outputs in comparison to a set of random natural pep- tides in order to determine an normalized adjusted percentile rank. With normal- ized ranks, the same thresholds can be used across different methods. Default thresholds are <2% for strong binders and <10% for weak binders.
url_iedb	string, iedb api url

Value

data frame, MHC II binding prediction result table

Examples

mhcII_hu(ag_present=c("DRB1_08_01"),ag_stim=c("DQA1_01_01","DQA1_04_01"),ag_self=c("DQA1_02_01"))

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prep human reference tables based on IMGT database

Description

prep human reference tables based on IMGT database

Usage

```
prep_ref_hu(
    url_imgt = "https://raw.githubusercontent.com/ANHIG/IMGTHLA/Latest/fasta/"
)
```

Arguments

url_imgt, string, github link of IMGT database

Value

data frame, MHC II sequence of officially named alleles

Examples

out <- prep_ref_hu()</pre>

utils

Description

preproc_huII() format and validate allele names

Usage

```
preproc_huII(allele_in)
```

```
pull_seq_huII(alleles_in, tbl_ref_in)
```

prep_lbl_huII(alleles_in)

comb_pred_tbl(nm_method, nm_sht, nm_fd, thold_score, thold_rank)

find_nonself_huII(dat_in)

pull_ag_self(dat_in)

find_core_mut(dat_in)

Arguments

allele_in	a vector contains allele name(s)
	pull_seq_huII() pull out sequence of each allele based on ref table
alleles_in	vector, allele names
	comb_pred_tbl() combine individual prediction tables by method, exclude none- binders and keep strong and weak binders only
tbl_ref_in	dataframe, reference table, default is human_all.csv from github
	prep_lbl_huII() concatenate allele strings for prediction table names
nm_method	string, prediction method used for IEDB prediction
nm_sht	string, short name of alleles
nm_fd	string, folder name which contains predict tables from IEDB
thold_score	list of vectors, binder thresholds by score
thold_rank	vector, binder thresholds by rank
	find_nonself_huII() find nonself binding peptides
dat_in	dataframe with pep_stim, core, pep_self selected from pull_ag_self

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