

Package ‘immunotation’

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Type Package

Title Tools for working with diverse immune genes

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Description

MHC (major histocompatibility complex) molecules are cell surface complexes that present antigens to T cells. The repertoire of antigens presented in a given genetic background largely depends on the sequence of the encoded MHC molecules, and thus, in humans, on the highly variable HLA (human leukocyte antigen) genes of the hyperpolymorphic HLA locus. More than 28,000 different HLA alleles have been reported, with significant differences in allele frequencies between human populations worldwide. Reproducible and consistent annotation of HLA alleles in large-scale bioinformatics workflows remains challenging, because the available reference databases and software tools often use different HLA naming schemes. The package immunotation provides tools for consistent annotation of HLA genes in typical immunoinformatics workflows such as for example the prediction of MHC-presented peptides in different human donors. Converter functions that provide mappings between different HLA naming schemes are based on the MHC restriction ontology (MRO). The package also provides automated access to HLA alleles frequencies in worldwide human reference populations stored in the Allele Frequency Net Database.

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R topics documented:

assemble_protein_complex	2
build_allele_group	3
decode_MAC	4
encode_MAC	4
get_G_group	5
get_mhcpan_input	6
get_P_group	6
get_serotypes	7
get_valid_organisms	7
human_protein_complex_table	8
plot_allele_frequency	8
query_allele_frequencies	9
query_haplotype_frequencies	10
query_population_detail	11
retrieve_chain_lookup_table	12

Index	13
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assemble_protein_complex
Assemble protein complex

Description

Assemble a table or MHC protein complexes for a given organism.

Usage

assemble_protein_complex(organism)

Arguments

`organism` Organism for which the lookup should be built (e.g. "human", "mouse", ...). The list of valid organisms can be found using the function `get_valid_organisms`

Value

a data frame with the MHC complexes annotated in MRO (only completely annotated complexes are returned)

Examples

```
assemble_protein_complex(organism = "mouse")
```

`build_allele_group` *Building a list of alleles to cover*

Description

`build_allele_group` e.g. A*01:01 -> A*01:01:01, A*01:01:02, A*01:01:03

Usage

```
build_allele_group(allele_selection)
```

Arguments

`allele_selection`
HLA allele for which the allele group should be built.

Value

list of alleles

Examples

```
build_allele_group("A*01:01")
```

decode_MAC

Decode MAC

Description

Decode a multiple allele code (MAC) into a list of HLA alleles. #’ The National Marrow Donor Program (NMDP) uses [MAC](<https://bioinformatics.bethematchclinical.org/hla-resources/allele-codes/allele-code-lists/>) to facilitate the reporting and comparison of HLA alleles. MAC represent groups of HLA alleles and are useful when the HLA typing is ambiguous and does not allow to narrow down one single allele from a list of alleles.

Usage

decode_MAC(MAC)

Arguments

MAC multiple allele code (e.g. "A*01:ATJNV")

Value

list of HLA alleles

Examples

```
MAC <- "A*01:ATJNV"
decode_MAC(MAC)
```

encode_MAC*Encode MAC*

Description

Encode a list of HLA alleles into multiple allele code (MAC). The National Marrow Donor Program (NMDP) uses [MAC](<https://bioinformatics.bethematchclinical.org/hla-resources/allele-codes/allele-code-lists/>) to facilitate the reporting and comparison of HLA alleles. MAC represent groups of HLA alleles and are useful when the HLA typing is ambiguous and does not allow to narrow down one single allele from a list of alleles.

Usage

encode_MAC(allele_list)

Arguments

allele_list list of HLA alleles (e.g. c("A*01:01:01", "A*02:01:01", "A*03:01"))

Value

encoded MAC

Examples

```
allele_list <- c("A*01:01:01", "A*02:01:01", "A*03:01")
encode_MAC(allele_list)
```

get_G_group

G groups

Description

Get the G groups for a list of HLA alleles. [G groups](http://hla.alleles.org/alleles/g_groups.html) are groups of HLA alleles that have identical nucleotide sequences across the exons encoding the peptide binding domains.

Usage

```
get_G_group(allele_list)
```

Arguments

allele_list List of alleles.

Value

Named list of G-groups the input alleles belong to.

Examples

```
allele_list <- c("DQB1*02:02:01", "DQB1*06:09:01")
get_G_group(allele_list)
```

`get_mhcpan_input` *Get format for NetMHCpan tools*

Description

NetMHCpan tools for MHC-peptide binding prediction require HLA complex names in a specific format. `get_mhcpan_input` formats a list of HLA alleles into a list of NetMHC-formated complexes.

Usage

```
get_mhcpan_input(allele_list, mhc_class)
```

Arguments

<code>allele_list</code>	list of HLA alleles (e.g. <code>c("A*01:01:01", "B*27:01")</code>)
<code>mhc_class</code>	<code>["MHC-I" "MHC-II"]</code> indicated which NetMHC you want to use.

Value

protein chain list as formatted for MHCpan input

Examples

```
allele_list <- c("A*01:01:01", "B*27:01")
get_mhcpan_input(allele_list, mhc_class = "MHC-I")
```

`get_P_group` *P groups*

Description

Get the P groups for a list of HLA alleles. [P groups](http://hla.alleles.org/alleles/p_groups.html) are groups of HLA alleles that have identical protein sequences in the peptide binding domains.

Usage

```
get_P_group(allele_list)
```

Arguments

<code>allele_list</code>	list of HLA alleles
--------------------------	---------------------

Value

Named list of P-groups the input alleles belong to.

Examples

```
allele_list <- c("DQB1*02:02:01", "DQB1*06:09:01")
get_P_group(allele_list)
```

get_serotypes	<i>Serotypes</i>
---------------	------------------

Description

Get the serotypes of the MHC complexes encoded by a list of MHC alleles.

Usage

```
get_serotypes(allele_list, organism = "human", mhctype)
```

Arguments

allele_list	List of allele
organism	Organism to be used for MRO lookup. If the organism does not match the given allele, an empty object is returned.
mhctype	["MHC-I" or "MHC-II"] MHC class to use for MRO lookup.

Value

Named list of serotypes, which only contains complexes contained in the MRO. If no serotype is annotated for a given complex, the list element is NA.

Examples

```
allele_list <- c("A*01:01:01", "B*27:01")
get_serotypes(allele_list, mhctype = "MHC-I")
```

get_valid_organisms	<i>get_valid_organisms</i>
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Description

get the list of organisms that are part of the MRO annotation

Usage

```
get_valid_organisms()
```

Value

list of organisms

Examples

```
get_valid_organisms()
```

<code>human_protein_complex_table</code>	<i>human_protein_complex_table</i>
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Description

`human_protein_complex_table`

Usage

```
human_protein_complex_table
```

Format

An object of class `data.frame` with 12385 rows and 8 columns.

Details

`human_protein_complex_table`: `human_protein_complex_table`.

Examples

```
# The human protein complex table is available in the following
# exported variable
human_protein_complex_table
```

<code>plot_allele_frequency</code>	<i>Plotting allele frequencies</i>
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Description

`plot_allele_frequency` Generate a World map displaying the frequency of a given table of HLA alleles. Use the function [query_allele_frequencies](#) to generate a table with allele frequencies.

Usage

```
plot_allele_frequency(allele_frequency)
```

Arguments

allele_frequency
returned by [query_allele_frequencies](#)

Value

ggplot2 object displaying the allele frequencies on a world map.

Examples

```
# select frequency of given allele
sel_allele_freq <- query_allele_frequencies(hla_selection = "A*02:01",
hla_sample_size_pattern = "bigger_than",
hla_sample_size = 10000, standard="g")

plot_allele_frequency(sel_allele_freq)
```

query_allele_frequencies
Query allele frequencies

Description

Query allele frequencies

Usage

```
query_allele_frequencies(
  hla_locus = NA,
  hla_selection = NA,
  hla_population = NA,
  hla_country = NA,
  hla_region = NA,
  hla_ethnic = NA,
  hla_sample_size_pattern = NA,
  hla_sample_size = NA,
  standard = "a"
)
```

Arguments

hla_locus HLA locus that will be used for filtering data. A, B, C, DPA1, DPB1, DQA1, DQB1, DRB1
hla_selection Allele that will be used for filtering data. e.g. A*01:01
hla_population Numeric identifier of the population that will be used for filtering. This identifier is defined by the Allele Frequency Net Database.

<code>hla_country</code>	Country of interest (e.g. Germany, France, ...).
<code>hla_region</code>	Geographic region of interest (e.g. Europe, North Africa, ...)
<code>hla_ethnic</code>	Ethnic origin of interest (e.g. Caucasoid, Siberian, ...)
<code>hla_sample_size_pattern</code>	Keyword used to define the filtering for a specific population size. e.g. "bigger_than", "equal", "less_than", "less_equal_than", "bigger_equal_than"
<code>hla_sample_size</code>	Integer number used to define the filtering for a specific population size, together with the <code>hla_sample_size_pattern</code> argument.
<code>standard</code>	Population standards, as defined in the package vignette. "g" - gold, "s" - silver, "a" - all

Value

`data.frame` object containing the result of the allele frequency query

Examples

```
# select frequencies of the A*02:01 allele,
# for gold standard population with more than 10,000 individuals
sel <- query_allele_frequencies(hla_selection = "A*02:01",
hla_sample_size_pattern = "bigger_than", hla_sample_size = 10000,
standard="g")
```

query_haplotype_frequencies
Query haplotype frequencies

Description

Query haplotype frequencies

Usage

```
query_haplotype_frequencies(
  hla_selection = NA,
  hla_population = NA,
  hla_country = NA,
  hla_region = NA,
  hla_ethnic = NA,
  hla_sample_size_pattern = NA,
  hla_sample_size = NA
)
```

Arguments

- `hla_selection` Alleles that will be used to build the haplotype query. One entry per locus. If no entry for a given locus, the function will search for haplotypes that do not include specifications for this locus. If any allele for a given locus should be considered, the list entry should be "A*" or other locus in same format.
- `hla_population` Numeric identifier of the population that will be used for filtering. This identifier is defined by the Allele Frequency Net Database.
- `hla_country` Country of interest (e.g. Germany, France, ...).
- `hla_region` Geographic region of interest (e.g. Europe, North Africa, ...)
- `hla_ethnic` Ethnic origin of interest (e.g. Caucasoid, Siberian, ...)
- `hla_sample_size_pattern` Keyword used to define the filtering for a specific population size. e.g. "bigger_than", "equal", "less_than", "less_equal_than", "bigger_equal_than"
- `hla_sample_size` Integer number used to define the filtering for a specific population size, together with the `hla_sample_size_pattern` argument.

Value

`data.frame` object containing the result of the allele frequency query

Examples

```
# works only for one haplotype at a time
query_haplotype_frequencies(hla_selection = c("A*02:01", "B*", "C*"),
hla_region = "Europe")
```

query_population_detail

Query population metainformation

Description

Query population metainformation

Usage

```
query_population_detail(population_ids)
```

Arguments

- `population_ids` List of numeric identifiers of the population that will be used for filtering. The identifier is defined by the Allele Frequency Net Database.

Value

data.frame object containing the result of the population detail query

Examples

```
population_detail <- query_population_detail(0001986)
```

```
retrieve_chain_lookup_table
```

Retrieve MHC chain lookup table

Description

Retrieve MHC chain lookup table

Usage

```
retrieve_chain_lookup_table(organism)
```

Arguments

organism name of organism (e.g. "human")

Value

Table containing MHC chain information for the organism. It contains chain names, MHC restriction and protein sequence.

Examples

```
retrieve_chain_lookup_table("mouse")
```

Index

* **datasets**
 human_protein_complex_table, 8

 assemble_protein_complex, 2

 build_allele_group, 3

 decode_MAC, 4

 encode_MAC, 4

 get_G_group, 5
 get_mhcpan_input, 6
 get_P_group, 6
 get_serotypes, 7
 get_valid_organisms, 7

 human_protein_complex_table, 8

 plot_allele_frequency, 8

 query_allele_frequencies, 8, 9, 9
 query_haplotype_frequencies, 10
 query_population_detail, 11

 retrieve_chain_lookup_table, 12