# Package 'igblastr'

June 29, 2025

```
Title User-friendly R Wrapper to IgBLAST
Description The igblastr package provides functions to conveniently install
      and use a local IgBLAST installation from within R.
      IgBLAST is described at <a href="https://pubmed.ncbi.nlm.nih.gov/23671333/">https://pubmed.ncbi.nlm.nih.gov/23671333/</a>.
      Online IgBLAST: <a href="mailto:right-num.nih.gov/igblast/">https://www.ncbi.nlm.nih.gov/igblast/</a>.
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URL https://bioconductor.org/packages/igblastr
BugReports https://github.com/HyrienLab/igblastr/issues
Version 0.99.11
License Artistic-2.0
Encoding UTF-8
Depends R (>= 4.2.0), tibble, Biostrings
Imports methods, utils, stats, tools, R.utils, curl, httr, xml2,
      rvest, xtable, jsonlite, S4Vectors, IRanges, GenomeInfoDb
Suggests parallel, testthat, knitr, rmarkdown, BiocStyle, ggplot2,
      dplyr, scales, ggseqlogo
SystemRequirements Perl (for install_IMGT_germline_db() only)
VignetteBuilder knitr
Collate utils.R long_to_wide_airr.R file-utils.R db-utils.R
      IMGT-utils.R AIRR-utils.R precompiled-igblast-utils.R
      cache-utils.R get_igblast_root.R edit_imgt_file.R
      igblast_info.R auxiliary-data-utils.R install_igblast.R
      make_blastdbs.R create_region_db.R create_germline_db.R
      create_c_region_db.R list_germline_dbs.R list_c_region_dbs.R
      install_IMGT_germline_db.R install_AIRR_germline_db.R
      augment_germline_db.R igblastn-args-utils.R outfmt7-utils.R
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```
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augment\_germline\_db
Add novel gene alleles to a germline db

# Description

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Three functions to add novel V, D, or J gene alleles to a germline db.

Note that these functions can also be used to combine germline databases from two different organisms. See "COMBINE GERMLINE DATABASES FROM TWO ORGANISMS" in the Examples section below for how to do this.

# Usage

```
augment_germline_db_V(db_name, novel_alleles, destdir=".", overwrite=FALSE)
augment_germline_db_D(db_name, novel_alleles, destdir=".", overwrite=FALSE)
augment_germline_db_J(db_name, novel_alleles, destdir=".", overwrite=FALSE)
```

# **Arguments**

db\_name

A single string that is the name of the cached germline db that contains the set of gene alleles to augment. Use <code>list\_germline\_dbs()</code> to list the cached germline dbs.

The exact function used (i.e. augment\_germline\_db\_V(), augment\_germline\_db\_D(), or augment\_germline\_db\_J()) determines the set of alleles to augment (i.e. alleles from the V, D, or J region).

novel\_alleles A single string that is the path to a FASTA file (possibly gz-compressed) where

the novel alleles are stored.

Alternatively, the novel alleles can be supplied as a named DNAStringSet object.

destdir A single string that is the path to the "destination directory", that is, the directory

where the augmented V-, D-, or J-region db is to be created. This directory will be created if it doesn't exist already. Note that, by default, the augmented region

db will be created in the current directory.

overwrite If the "destination directory" already contains a V-, D-, or J-region db, should it

be overwritten?

#### Value

An invisible NULL.

#### See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- list\_germline\_dbs to list the cached germline dbs.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

```
if (!has_igblast()) install_igblast()
query <- system.file(package="igblastr", "extdata",</pre>
                  "heavy_sequences.fasta")
use_c_region_db("_IMGT.human.IGH+IGK+IGL.202412")
## USE HUMAN GERMLINE DATABASE FROM AIRR
## -----
use_germline_db("_AIRR.human.IGH+IGK+IGL.202501")
AIRR_df <- igblastn(query)
## ADD NOVEL V ALLELES
## -----
## 'fake_human_V_alleles.fasta' contains made-up novel V alleles:
## - 2 novel alleles for gene IGHV1-8: IGHV1-8*fake1, IGHV1-8*fake2
## - 1 novel allele for gene IGHV4-61: IGHV4-61*fake
my_novel_V_alleles <- system.file(package="igblastr", "extdata",</pre>
                              "novel_germline_alleles",
                              "fake_human_V_alleles.fasta")
## Take a quick look at these novel V alleles:
readDNAStringSet(my_novel_V_alleles)
## Create a new V germline database that combines the V alleles
## from _AIRR.human.IGH+IGK+IGL.202501 with our novel V alleles:
```

```
myVdb_path <- file.path(tempdir(), "myVdb")</pre>
augment_germline_db_V("_AIRR.human.IGH+IGK+IGL.202501",
                      my_novel_V_alleles,
                      destdir=myVdb_path)
## To use this new augmented V germline database with igblastn(),
## supply its path via the 'germline_db_V' argument:
AIRR_df2 <- igblastn(query, germline_db_V=myVdb_path)
## A QUICK COMPARISON BETWEEN 'AIRR_df' AND 'AIRR_df2'
## Index of rows where "v_call" has changed between 'AIRR_df'
## and 'AIRR_df2':
idx <- which(AIRR_df$v_call != AIRR_df2$v_call)</pre>
idx # 2 rows
AIRR_df[idx, c("v_call", "v_cigar", "v_identity")]
AIRR_df2[idx, c("v_call", "v_cigar", "v_identity")]
## Besides these 2 rows, all the other rows are the same:
stopifnot(all.equal(AIRR_df[-idx, ], AIRR_df2[-idx, ]))
## COMBINE GERMLINE DATABASES FROM TWO ORGANISMS
## The augment_germline_db_[VDJ]() functions can be used to combine
## germline databases from two different organisms. This can be useful
## for example when working with BCR sequences from mice that have been
## engineered to have both mouse and some human immunoglobulin genes.
## To create a hybrid human/mouse V germline database, we can either:
##
## (1) Add all (or a subset of) mouse V alleles to all human V alleles.
##
       This is done by extracting mouse V germline allele sequences from
##
       a cached germline database and using them to augment a cached
##
       germline database for human.
##
## (2) Add all (or a subset of) human V alleles to all mouse V alleles.
       This is done by extracting human V germline allele sequences from
##
##
       a cached germline database and using them to augment a cached
##
       germline database for mouse.
##
## Note that:
## - We can choose to subset or not the V germline allele sequences
    extracted from one \ensuremath{\text{V}} germline database before adding them to the
##
    other V germline database.
## - The two approaches above are equivalent if we don't subset, that
## is, if we combine **all** human V alleles with **all** mouse V
    alleles.
## - However if our engineered mice only have a small known subset of
## human immunoglobulin genes (e.g. IGHV1-2), then we might want to
    create a hybrid human/mouse germline database that only adds the
##
    human alleles for genes IGHV1-2 to the mouse V alleles. In this
```

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```
case we need to use (2).
## Let's do (2):
db_name1 <- "_AIRR.mouse.PWD_PhJ.IGH+IGK+IGL.202501"</pre>
db_name2 <- "_AIRR.human.IGH+IGK+IGL.202501"
## Extract human V germline alleles:
human_V_alleles <- load_germline_db(db_name2, "V")</pre>
## Subset to keep only alleles for genes IGHV1-2:
idx <- grep("^IGHV[12]", names(human_V_alleles))</pre>
human_V12_alleles <- human_V_alleles[idx]</pre>
## Create a new V germline database that combines the mouse V
## alleles from 'db_name1' with the alleles in 'human_V12_alleles':
engmouseVdb_path <- file.path(tempdir(), "engmouseVdb")</pre>
augment_germline_db_V(db_name1, human_V12_alleles,
                      destdir=engmouseVdb_path)
## Then, assuming that 'query' contains BCR sequences from the
## engineered mice:
## Not run:
  use_germline_db(db_name1)
  use_c_region_db("_IMGT.mouse.IGH.202412")
  igblastn(query, germline_db_V=engmouseVdb_path, ...)
## End(Not run)
## Note that, by default, the mouse-only D and J databases that
## we selected with 'use_germline_db(db_name1)' are being used.
## If we also want to create hybrid D and J databases, we need
## to repeat the above steps for each of them. Then we need to
## specify the paths to the 3 hybrid databases when we call igblastn():
## Not run:
  igblastn(query, germline_db_V=engmouseVdb_path,
                  germline_db_D=engmouseDdb_path,
                  germline_db_J=engmouseJdb_path,
                  . . . )
## End(Not run)
```

auxiliary-data-utils Manipulation of IgBLAST auxiliary data

# **Description**

A standard IgBLAST installation – like the one used by the **igblastr** package – typically includes *auxiliary data* for various organisms, in the form of one tabulated file per organism. Each file indicates the germline J gene coding frame start position, the J gene type, and the CDR3 end position for each sequence in the germline J sequence database. See <a href="https://ncbi.github.io/igblast/cook/How-to-set-up.html">https://ncbi.github.io/igblast/cook/How-to-set-up.html</a> for the details.

You can use get\_igblast\_auxiliary\_data() to obtain the path to the file containing the auxiliary data for a given organism.

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#### **Usage**

```
get_igblast_auxiliary_data(organism)
load_igblast_auxiliary_data(organism)
```

# **Arguments**

organism

A single string containing the name of an organism as returned by list\_igblast\_organisms().

#### Value

get\_igblast\_auxiliary\_data(): Returns a single string containing the path to the auxiliary data included in the IgBLAST installation used by **igblastr**, for the specified organism. Not necessarily suitable to use with igblastn() (see WARNING below).

load\_igblast\_auxiliary\_data(): Returns the auxiliary data in a data.frame with 1 row per germline J sequence and the following columns:

- 1. sseqid: gene/allele name a.k.a. subject sequence id;
- 2. coding\_frame\_start: first coding frame start position (position is 0-based);
- 3. chaintype: chain type;
- 4. CDR3\_stop: CDR3 stop;
- 5. extra\_bps: extra base pairs beyond J coding end.

# WARNING

According to https://ncbi.github.io/igblast/cook/How-to-set-up.html the auxiliary data included in IgBLAST is specific to a particular NCBI or IMGT germline db. Unfortunately this means that this data is NOT guaranteed to be compatible with the germline db that you will use with igblastn(). See documentation of the auxiliary\_data argument in ?igblastn for more information about this.

# See Also

- https://ncbi.github.io/igblast/cook/How-to-set-up.html for important information about the IgBLAST auxiliary data.
- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- install\_igblast to perform an internal IgBLAST installation.
- get\_igblast\_root to get (or set) the IgBLAST installation used (or to be used) by the **ig-blastr** package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

```
if (!has_igblast()) install_igblast()
igblast_info()
list_igblast_organisms()
## Make sure to read the WARNING above before using the auxiliary
## data below with igblastn()!
```

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```
get_igblast_auxiliary_data("human")
load_igblast_auxiliary_data("human")
get_igblast_auxiliary_data("rhesus_monkey")
load_igblast_auxiliary_data("rhesus_monkey")
```

bcr\_browser

Display annotated BCR sequences in the browser

# **Description**

Use bcr\_browser() to display the annotated BCR sequences returned by igblastn() in the browser. For each sequence, the V, D, and J segments are shown as well as the FWR1-4 and CDR1-3 regions. Additionally, the C segments are shown if the C-region information is available.

# Usage

# **Arguments**

AIRR\_df

The AIRR-formatted data.frame or tibble returned by igblastn(). Note that calling bcr\_browser() on a data.frame with thousands of rows is quite resource-intensive (it can even crash your browser!), so in this case we recommend subsetting the data.frame before passing it to bcr\_browser() to keep the number of rows under 2000.

 $\verb|show.full.sequence||$ 

By default, the part of the BCR sequences upstream of the V region is not shown. Set show. full.sequence to TRUE to show it.

dna.coloring

Whether the nucleotides in the BCR sequences (sequence column in AIRR\_df) should be colored or not.

Vcolor, Dcolor, Jcolor, Ccolor

The background colors of the V, D, J, and C segments of the BCR sequences. Note that the C segments are shown only if AIRR\_df contains C-region information.

FWRcolor, CDRcolor

The background colors of the Framework Regions (FWR1-4) and Complementarity-Determining Regions (CDR1-3), respectively.

# Value

0 or the error code returned by the internal call to browseURL(), invisibly.

#### See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.
- tibble objects implemented in the tibble package.

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#### **Examples**

```
if (!has_igblast()) install_igblast()
query <- system.file(package="igblastr", "extdata",</pre>
               "heavy_sequences.fasta")
use_germline_db("_AIRR.human.IGH+IGK+IGL.202501")
## -----
## With C regions
## -----
use_c_region_db("_IMGT.human.IGH+IGK+IGL.202412")
AIRR_df <- igblastn(query)
bcr_browser(AIRR_df)
## By default, the part of the sequences upstream of the V region is
## not shown. Use 'show.full.sequence=TRUE' to show the full sequences:
bcr_browser(AIRR_df, show.full.sequence=TRUE)
## -----
## No C regions
## -----
use_c_region_db("")
AIRR_df2 <- igblastn(query)
bcr_browser(AIRR_df2)
```

get\_igblast\_root

Control IgBLAST installation to use

#### **Description**

Get (or set) the IgBLAST installation used (or to be used) by the igblastr package.

# Usage

```
get_igblast_root()
set_igblast_root(version_or_path)
```

# **Arguments**

```
version_or_path
```

A single string that is either a version number (e.g. "1.22.0") or the path to an IgBLAST installation.

# **Details**

set\_igblast\_root can be used to set or change the path to the IgBLAST installation to use. This can be an *internal* or *external* installation.

In the former case, version\_or\_path should be the version of an existing *internal* installation. The setting will be persistent.

In the latter case, it should be the full path (absolute or relative) to the *root directory* of a valid *external* installation. Note that the setting won't be persistent i.e. it won't be remembered across R

sessions. See ?IGBLAST\_ROOT for how to set the *external* IgBLAST installation to use in **igblastr** in a persistent manner.

# Value

get\_igblast\_root() returns a single string containing the path to the *root directory* of the Ig-BLAST installation used by **igblastr**.

set\_igblast\_root() returns a single string containing the path to the *root directory* of the newly selected IgBLAST installation. The string is returned invisibly.

#### See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- install\_igblast to perform an internal IgBLAST installation.
- igblast\_info to collect basic information about the IgBLAST installation used by the igblastr package.
- IGBLAST\_ROOT to set the *external* IgBLAST installation to be used by the **igblastr** package in a persistent manner.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

# **Examples**

```
if (!has_igblast()) install_igblast()
get_igblast_root()
```

igblastn

BLAST for Ig and TCR sequences

# **Description**

The igblastn() function is a wrapper to the igblastn *standalone executable* included in Ig-BLAST. This is the main function in the **igblastr** package.

# Usage

# **Arguments**

query

A character vector containing the paths to the input files (FASTA), or a named DNAStringSet object.

If a character vector, then query must be of length >= 1 and each vector element must be the path to a FASTA file (possibly gz-compressed). In the context of IgBLAST, the DNA sequences in the FASTA files are referred to as the query sequences, and the sequence names found in the description lines of the FASTA records are referred to as the query sequence ids.

Note that the query sequences are typically (but not always) stored in a single file, in which case query will be a single string. If more than one FASTA file is specified via query, then igblastn() will concatenate all the files together and pass the resulting file to the igblastn standalone executable.

If query is a DNAStringSet object, then it must have names on it. These will be considered the query sequence ids.

One of "AIRR", 3, 4, 7, or 19. "AIRR" is the default and is an alias for 19. outfmt can also be a string describing a customized format 7 e.g. "7 qseqid sseqid pident nident length score". See ?list\_outfmt7\_specifiers for more information about customizing format 7.

germline\_db\_V "auto" (the default), or the path to a V-region db.

> Note that, by default (i.e. when germline\_db\_V is omitted or set to "auto"), igblastn() uses the V-region db that belongs to the cached germline db currently selected. See ?use\_germline\_db for how to select the cached germline db to use with igblastn().

Same as germline\_db\_V but for the D-region db. germline\_db\_D

germline\_db\_J Same as germline\_db\_V but for the J-region db.

germline\_db\_V\_seqidlist, germline\_db\_D\_seqidlist, germline\_db\_J\_seqidlist

> Restrict search of germline database to list of gene alleles. A list of gene alleles can be specified either as a character vector of gene allele identifiers (e.g. IGHV3-23\*01, IGHV3-23\*04, etc...) or as the path to a file containing the identifiers (one identifier per line). In the latter case, a file object must be passed to the germline\_db\_V\_seqidlist, germline\_db\_D\_seqidlist, or germline\_db\_J\_seqidlist argument. The file object will typically be constructed with something like file("path/to/some/file").

organism

"auto" (the default), or the organism associated with the query sequences. Supported organisms include human, mouse, rat, rabbit and rhesus\_monkey. Use list\_igblast\_organisms() to obtain this list programmatically.

Note that, by default (i.e. when organism is omitted or set to "auto"), igblastn() infers the organism from the name of the cached germline db currently selected. See ?use\_germline\_db for how to select the cached germline db to use with igblastn().

c\_region\_db

"auto" (the default), NULL, or the path to a C-region db.

Note that, by default (i.e. when c\_region\_db is omitted or set to "auto"), igblastn() uses the cached C-region db currently selected. See ?use\_c\_region\_db for how to select the cached C-region db to use with igblastn().

auxiliary\_data

"auto" (the default), or the path to a file containing the coding frame start positions for the sequences in the J-region db, or NULL.

outfmt

Note that, by default (i.e. when auxiliary\_data is omitted or set to "auto"), igblastn() uses one of the auxiliary data files included in the IgBLAST installation used by **igblastr**. More precisely, igblastn() uses get\_igblast\_auxiliary\_data() internally to obtain the path to the organism-specific auxiliary data file.

# IMPORTANT NOTES:

- Supplying auxiliary data that is not compatible with the V gene sequences
  of the selected germline db can cause igblastn() to return improper frame
  status or CDR3 information (other returned information will still be correct). See ?get\_igblast\_auxiliary\_data for more information.
- When auxiliary\_data is set to NULL, then no auxiliary data is used. In this case, igblastn() can emit a significant number of the following warning:

Warning: Auxilary data file could not be found and various columns of the returned AIRR-formatted tibble (e.g. columns vj\_in\_frame, productive, cdr3, fwr4, and others) will be filled with NAs.

Extra arguments to be passed to the igblastn *standalone executable*. The list of valid arguments can be displayed with igblastn\_help().

Note that the argument/value pairs must be passed to the igblastn() function in the usual R fashion. For example, what would be passed as -num\_alignments 1 -num\_threads 8 when invoking the igblastn *standalone executable* in a terminal should be passed as num\_alignments\_V=1, num\_threads=8 when calling the igblastn() function:

igblastn(query, num\_alignments\_V=1, num\_threads=8)

NULL (the default), or the path to the file where the igblastn *standalone executable* should write its output.

Note that, by default (i.e. when out is omitted or set to NULL), igblastn() instructs the igblastn *standalone executable* to write its output to a temporary file.

Whether igblastn() should parse the plain-text output produced by the igblastn *standalone executable* or not, before returning it to the user. TRUE by default.

If set to FALSE, then igblastn() returns the output as-is in a character vector, with one line per element in the vector. Note that igblastn() sets the "igblastn\_raw\_output" class attribute on this character vector, which allows compact display of the vector (this is achieved via a dedicated print() method defined in the **igblastr** package). The class attribute can be dropped with unclass().

show.in.browser

For igblastn(): Whether the output of the igblastn *standalone executable* should also be displayed in the browser or not (in addition to being returned by the igblastn() function call). FALSE by default.

For igblastn\_help(): Whether the help printed by the igblastn *standalone executable* (when invoked with the -h or -help argument) should be displayed in the browser or not. FALSE by default.

show.command.only

TRUE or FALSE. If set to TRUE, igblastn() won't invoke the igblastn *standalone executable* and instead will display the full command that shows how it would have invoked it. Note that the command is also returned in an invisible character vector. FALSE by default.

TRUE or FALSE. If set to FALSE (the default), the igblastn *standalone executable* is invoked with the -h argument. Otherwise, it's invoked with the -help argument.

. .

out

parse.out

long.help

#### Value

igblastn() captures the output produced by the igblastn standalone executable and returns it as:

- A tibble with 1 row per query sequence if outfmt is "AIRR" or 19 and parse.out is TRUE.
- A nested list with two top-level components (records and footer) if outfmt is 7 (or a customized format 7) and parse.out is TRUE. See ?parse\_outfmt7 for more information.
- A character vector with class attribute "igblastn\_raw\_output" on it in all other cases, that is, if parse.out is FALSE or outfmt is 3 or 4. See the parse.out argument above for more information.

#### Note

By default, the NCBI BLAST+ and IgBLAST programs will "call home" to report usage when they run on a computer with internet access. See <a href="https://www.ncbi.nlm.nih.gov/books/NBK569851/">https://www.ncbi.nlm.nih.gov/books/NBK569851/</a> for the details. This can induce a significant slowdown in some situations e.g. when the igblastn standalone executable is called in a loop on a small set of query sequences at each iteration.

For this reason, the "call home" feature is disabled in **igblastr** by default, unless environment variable BLAST\_USAGE\_REPORT is set to true. See ?igblastr\_usage\_report for more information.

#### See Also

- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.
- install\_igblast to perform an internal IgBLAST installation.
- igblast\_info to collect basic information about the IgBLAST installation used by the igblastr package.
- install\_IMGT\_germline\_db to install a germline db from IMGT.
- use\_germline\_db to select the cached germline db to use with igblastn().
- use\_c\_region\_db to select the cached C-region db to use with igblastn().
- bcr\_browser to display the annotated sequences returned by igblastn() in the browser.
- list\_outfmt7\_specifiers for how to customize output format 7.
- list\_igblast\_organisms to list the organisms supported by IgBLAST.
- augment\_germline\_db to add novel gene alleles to a germline db.
- igblastr\_usage\_report to turn "Usage Reporting" on or off.
- DNAStringSet objects implemented in the **Biostrings** package.
- tibble objects implemented in the tibble package.

```
if (!has_igblast()) install_igblast()

## ------
## Preliminary steps
## -----
igblast_info()

## Files 'heavy_sequences.fasta' and 'light_sequences.fasta' included
## in igblastr contain 250 paired heavy- and light- chain sequences (125)
```

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```
## sequences in each file) downloaded from OAS (the Observed Antibody
## Space database):
filenames <- paste0(c("heavy", "light"), "_sequences.fasta")</pre>
query <- system.file(package="igblastr", "extdata", filenames)</pre>
## Install Human germline db from IMGT (Perl required!):
db_name <- install_IMGT_germline_db("202506-1", "Homo_sapiens", force=TRUE)</pre>
## Select germline db to use with igblastn():
use_germline_db(db_name)
## Select C-region db to use with igblastn():
use_c_region_db("_IMGT.human.IGH+IGK+IGL.202412")
## -----
## Call igblastn()
## We don't specify the 'outfmt' argument so output will be in AIRR format.
AIRR_df <- igblastn(query)
AIRR df
## The result is a tibble with one row per query sequence:
class(AIRR_df)
dim(AIRR_df)
## You can call bcr_browser() on 'AIRR_df' to vizualize the annotated
\#\# sequences in the browser. See '?bcr_browser'.
## Note that this tibble can easily be converted to an ordinary data.frame
## with 'as.data.frame()', or to a DataFrame with 'as(., "DataFrame")':
as(AIRR_df, "DataFrame")
## -----
## More examples
## -----
## See '?parse_outfmt7' for more examples.
```

igblastr\_usage\_report Turn "Usage Reporting" on or off

# **Description**

By default, the NCBI BLAST+ and IgBLAST programs will "call home" to report usage when they run on a computer with internet access. See <a href="https://www.ncbi.nlm.nih.gov/books/NBK569851/">https://www.ncbi.nlm.nih.gov/books/NBK569851/</a> for the details. This can induce a significant slowdown in some situations e.g. when the igblastn standalone executable is called in a loop on a small set of query sequences at each iteration.

For this reason, the "call home" feature is disabled in **igblastr** by default, unless environment variable BLAST\_USAGE\_REPORT is set to true.

More precisely, the "call home" feature is controlled by global option igblastr\_usage\_report in **igblastr**. On package startup, this option is set to TRUE if environment variable BLAST\_USAGE\_REPORT is set to true. Otherwise (i.e. if BLAST\_USAGE\_REPORT is not set, or is set to false or gibberish) it is set to FALSE.

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#### **Details**

The user can change the value of global option igblastr\_usage\_report any time with:

```
options(igblastr_usage_report=TRUE)
or with:
    options(igblastr_usage_report=FALSE)
To get the value of this option, use:
```

getOption("igblastr\_usage\_report")

Note that changing the value of a global option interactively with options (...) won't be remembered across R sessions. For a persistent change, you can either:

- Put the options (...) command in your .Rprofile file. See ?Rprofile for more information. Note that this is the standard way of setting global options persistently.
- In the particular case of global option igblastr\_usage\_report an alternative is to define environment variable BLAST\_USAGE\_REPORT outside R. The exact way to do this is OS-dependent e.g. on Linux and Mac you can define it in your user's .profile by adding the following line to it:

```
export BLAST_USAGE_REPORT=true
```

# See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

# **Examples**

```
## Check current status of usage reporting:
getOption("igblastr_usage_report")

## Turn on usage reporting:
options(igblastr_usage_report=TRUE)

## Turn off usage reporting:
options(igblastr_usage_report=FALSE)
```

 $igblast\_info$ 

Check IgBLAST used by igblastr

# **Description**

Collect basic information about the IgBLAST installation used by the **igblastr** package, or about any IgBLAST installation on the user machine.

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#### **Usage**

```
igblast_info(igblast_root=get_igblast_root())
igblast_build(igblast_root=get_igblast_root())
igblastn_version(igblast_root=get_igblast_root(), raw.version=FALSE)
makeblastdb_version(igblast_root=get_igblast_root(), raw.version=FALSE)
list_igblast_organisms(igblast_root=get_igblast_root())
has_igblast()
```

# **Arguments**

igblast\_root

A single string that is the path to an IgBLAST installation. By default igblast\_root is set to get\_igblast\_root(), which is the path to the IgBLAST installation used by the **igblastr** package. See ?get\_igblast\_root for more information.

Note that the supplied string must contain the path to the *root directory* of an IgBLAST installation, that is, to a directory with a bin subdirectory in it that has the igblastn, igblastp, and makeblastdb *standalone executables* (on Windows these executables are files named igblastn.exe, igblastp.exe, and makeblastdb.exe, respectively).

raw.version

By default (i.e. when raw.version is omitted or set to FALSE), igblastn\_version() and makeblastdb\_version() return the version string of the igblastn and makeblastdb *standalone executables* included in IgBLAST. This string is extracted from the output produced by system commands:

```
\begin{array}{c} \text{igblastn -version} \\ \\ \text{and} \\ \\ \text{makeblastdb -version} \end{array}
```

When raw.version is set to TRUE, igblastn\_version() and makeblastdb\_version() return the *full ouput* produced by the above commands.

#### Value

igblast\_info() returns a named list containing basic information about the IgBLAST installation. igblast\_build() returns a single string containing IgBLAST build information.

By default, igblastn\_version() returns a single string containing the version of the igblastn *standalone executable* included in IgBLAST.

By default, makeblastdb\_version() returns a single string containing the version of the makeblastdb standalone executable included in IgBLAST.

list\_igblast\_organisms() returns a character vector that lists the organisms for which IgBLAST provides internal data. Note that this is obtained by simply listing the content of the internal\_data directory located in the IgBLAST installation.

has\_igblast() returns TRUE or FALSE.

# See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- install\_igblast to perform an internal IgBLAST installation.

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• get\_igblast\_root to get (or set) the IgBLAST installation used (or to be used) by the **ig-blastr** package.

- IGBLAST\_ROOT to set the *external* IgBLAST installation to be used by the **igblastr** package in a persistent manner.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

# **Examples**

```
if (!has_igblast()) install_igblast()
igblast_info()
list_igblast_organisms()
```

IGBLAST\_ROOT

Use an external IgBLAST installation

# **Description**

Select the *external* IgBLAST installation to use in **igblastr** in a persistent manner.

#### **Details**

The **igblastr** package can use 2 types of IgBLAST installation:

- 1. Internal (a.k.a. igblastr-managed): refers to an installation obtained with install\_igblast().
- 2. External: refers to an installation that is not managed by the **igblastr** package. This is usually an installation that was manually performed by you or a system administrator on your machine. It can be a system-wide installation or a per-user installation.

To use an *external* installation of IgBLAST in **igblastr**, set environment variable IGBLAST\_ROOT to the path of the installation. Note that this must be the path to the *root directory* of the IgBLAST installation, that is, to a directory with a bin subdirectory in it that has the igblastn, igblastp, and makeblastdb *standalone executables* (on Windows these executables are files named igblastn.exe, igblastp.exe, and makeblastdb.exe, respectively).

This can be done within your current R session with Sys. setenv(IGBLAST\_ROOT="path/to/igblast\_root") for testing. However, this won't be remembered across R sessions.

To set IGBLAST\_ROOT in a persistent manner, define it outside R. The exact way to do this is OS-dependent e.g. on Linux and Mac you can define it in your user's .profile by adding the following line to it:

```
export IGBLAST_ROOT="path/to/igblast_root"
```

# See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- install\_igblast to perform an *internal* IgBLAST installation.
- igblast\_info to collect basic information about the IgBLAST installation used by the igblastr package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

install\_igblast 17

# Description

Download and install a pre-compiled IgBLAST from NCBI FTP site for use with igblastr.

# Usage

```
install_igblast(release="LATEST", force=FALSE, ...)
```

# Arguments

release	A single string specifying the IgBLAST release version to install. For example "LATEST" (recommended), or one of the IgBLAST release versions listed at <a href="https://ftp.ncbi.nih.gov/blast/executables/igblast/release/">https://ftp.ncbi.nih.gov/blast/executables/igblast/release/</a> (e.g. "1.21.0"). Note that old versions have not been tested and are not guaranteed to be compatible with the <b>igblastr</b> package.
force	Set to TRUE to reinstall if the specified IgBLAST release version is already installed.
	Extra arguments to be passed to the internal call to download.file(). See ?download.file in the utils package for more information.

# Value

The path to the root directory of the IgBLAST installation, as an invisible string.

#### See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- IGBLAST\_ROOT to use an external IgBLAST installation.
- igblast\_info to collect basic information about the IgBLAST installation used by the igblastr package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

```
if (!has_igblast()) install_igblast()
igblast_info()
```

# **Description**

The install\_IMGT\_germline\_db() function downloads IMGT/V-QUEST germline sequences from the IMGT website for a given organism, and stores them in a local germline database. This local database gets installed in **igblastr**'s persistent cache. It can then be used later with igblastn().

# Usage

# **Arguments**

release	A single string specifying the IMGT/V-QUEST release to get the germline sequences from (or to list the organisms from for list_IMGT_organisms()). Use list_IMGT_releases() to list all releases.
organism	A single string specifying the organism for which to get the germline sequences.
force	Set to TRUE to reinstall if the requested database is already installed.
	Extra arguments to be passed to the internal call to download.file(). See ?download.file in the <b>utils</b> package for more information.
recache	list_IMGT_releases() uses a caching mechanism so that the list of IMGT/V-QUEST releases gets downloaded only once from the IMGT website during an R session (note that this caching is done in memory so it does not persist across sessions). Set recache to TRUE to force a new download (and recaching) of the list of IMGT/V-QUEST releases.

# Value

install\_IMGT\_germline\_db() returns the name to the newly installed germline db as an invisible string.

list\_IMGT\_releases() returns the list of IMGT/V-QUEST releases in a character vector. The releases are sorted from newest to oldest (latest release is first).

list\_IMGT\_organisms() returns the list of organisms included in the specified IMGT/V-QUEST release in a character vector.

IMGT\_is\_up() returns TRUE or FALSE, indicating whether the IMGT website at <a href="https://www.imgt.org">https://www.imgt.org</a> is up and running or down.

list\_c\_region\_dbs

#### Note

install\_IMGT\_germline\_db() generates the local database by performing the instructions provided at https://ncbi.github.io/igblast/cook/How-to-set-up.html.

In particular, utility tool *edit\_imgt\_file.pl* (Perl script included in the IgBLAST installation) is invoked to process the sequences. This means that Perl must be available on your machine.

#### See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- use\_germline\_db to select the cached germline db to use with igblastn().
- The IMGT website: https://www.imgt.org/.
- The IMGT/V-QUEST download site: https://www.imgt.org/download/V-QUEST/.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

# **Examples**

```
if (!has_igblast()) install_igblast()

if (IMGT_is_up()) {
    ## As of March 26, 2025, the latest IMGT/V-QUEST release is 202506-1:
    list_IMGT_releases()

    list_IMGT_organisms("202506-1")

## Download Mouse germline sequences from IMGT/V-QUEST 202506-1, and
    ## store them in a cached germline database:
    install_IMGT_germline_db("202506-1", organism="Mus musculus", force=TRUE)

## List the cached germline databases:
    list_germline_dbs()

## Tell igblastn() to use the newly installed germline db:
    use_germline_db("IMGT-202506-1.Mus_musculus.IGH+IGK+IGL")
}
```

list\_c\_region\_dbs

List cached C-region dbs and select one to use with igblastn()

# **Description**

Use list\_c\_region\_dbs() to list all the *cached C-region dbs*, that is, all the C-region databases currently installed in **igblastr**'s persistent cache.

Use use\_c\_region\_db() to select the cached C-region db to use with igblastn(). This choice will be remembered for the duration of the current R session but can be changed anytime.

Use load\_c\_region\_db() to load the nucleotide sequences of the gene regions stored in a cached C-region db.

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#### **Usage**

```
list_c_region_dbs(builtin.only=FALSE, names.only=FALSE, long.listing=FALSE)
use_c_region_db(db_name=NULL, verbose=FALSE)
load_c_region_db(db_name)
```

# **Arguments**

builtin.only By default list\_c\_region\_dbs() returns the list of all cached C-region dbs,

including built-in C-region dbs. Set builtin.only to TRUE to return only the list of built-in C-region dbs. Note that built-in dbs are prefixed with an underscore

(\_).

names.only By default list\_c\_region\_dbs() returns the list of cached C-region dbs in a

data.frame with one db per row. Set names  $.\,\mbox{only}$  to TRUE to return only the db

names in a character vector.

long.listing TRUE or FALSE. If set to TRUE, then list\_c\_region\_dbs() returns a named list

with one list element per C-region db. Each list element is a named integer

vector that indicates the number of C-region sequences per locus.

Ignored if names.only is set to TRUE.

db\_name For use\_c\_region\_db():

NULL or a single string specifying the name of the cached C-region db to use.

Use list\_c\_region\_dbs() to list all the cached C-region dbs.

If set to NULL (the default), then use\_c\_region\_db() returns the name of the cached C-region db that is currently in use, if any. Otherwise it returns the empty

string ("").

Note that the current selection can be cancelled with use\_c\_region\_db("").

For load\_c\_region\_db():

A single string specifying the name of the cached C-region db from which to load the gene regions. Use list\_c\_region\_dbs() to list all the cached C-

region dbs.

verbose If set to TRUE, then use\_c\_region\_db() will display some information about

its internal operations.

# **Details**

The **igblastr** package provides a small set of utilities to manage the cached germline and C-region databases to use with igblastn().

Terminology:

- A *cached germline db* contains the nucleotide sequences of the V, D, and J gene regions for a given organism.
- A *cached C-region db* contains the nucleotide sequences of the C regions (i.e. constant gene regions) for a given organism.

This man page documents the basic utilities to operate on cached C-region dbs: list\_c\_region\_dbs(), use\_c\_region\_db(), and load\_c\_region\_db().

The basic utilities to operate on cached germline dbs are documented in the man page for list\_germline\_dbs.

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#### Value

list\_c\_region\_dbs() returns the list of all cached C-region dbs in a data.frame with one db per row (if names.only is FALSE, which is the default), or in a character vector (if names.only is TRUE). Column C in the data.frame indicates the number of C-region sequences in each db.

Built-in dbs are prefixed with an underscore (\_). Note that the C-region built-in dbs from IMGT were downloaded from https://www.imgt.org/vquest/refseqh.html#constant-sets and included in the igblastr package on the date indicated by the suffix of the db name.

When called with no argument, use\_c\_region\_db() returns a single string containing the name of the cached C-region db currently used by igblastn() if any, or the empty string ("") if igblastn() is not using any C-region db.

When called with the db\_name argument, use\_c\_region\_db(db\_name) returns db\_name invisibly.

load\_c\_region\_db() returns the nucleotide sequences from the specified C-region db in a named DNAStringSet object.

# See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- use\_germline\_db to select the cached germline db to use with igblastn().
- DNAStringSet objects in the **Biostrings** package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

```
if (!has_igblast()) install_igblast()

## 3 built-in C-region dbs prefixed with an underscore:
list_c_region_dbs()

list_c_region_dbs(names.only=TRUE) # db names only

list_c_region_dbs(long.listing=TRUE) # long listing

## Select C-region db to use with igblastn():
use_c_region_db("_IMGT.human.IGH+IGK+IGL.202412")

use_c_region_db() # get current selection

use_c_region_db("") # cancel current selection

use_c_region_db()

## Load C-region sequences:
load_c_region_db("_IMGT.human.IGH+IGK+IGL.202412")
load_c_region_db("_IMGT.mouse.IGH.202412")
```

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list\_germline\_dbs

List cached germline dbs and select one to use with igblastn()

#### **Description**

Use list\_germline\_dbs() to list all the *cached germline dbs*, that is, all the germline databases currently installed in **igblastr**'s persistent cache.

Use use\_germline\_db() to select the cached germline db to use with igblastn(). This choice will be remembered for the duration of the current R session but can be changed anytime.

Use load\_germline\_db() to load the nucleotide sequences of the gene regions stored in a cached germline db.

# Usage

```
list_germline_dbs(builtin.only=FALSE, names.only=FALSE, long.listing=FALSE)
use_germline_db(db_name=NULL, verbose=FALSE)
load_germline_db(db_name, region_types=NULL)
```

#### **Arguments**

builtin.only By default list\_germline\_dbs() returns the list of all cached germline dbs,

including built-in germline dbs. Set builtin.only to TRUE to return only the list of built-in germline dbs. Note that built-in dbs are prefixed with an underscore

(\_).

names.only By default list\_germline\_dbs() returns the list of cached germline dbs in a

data.frame with one db per row. Set names.only to TRUE to return only the db

names in a character vector.

long.listing TRUE or FALSE. If set to TRUE, then list\_germline\_dbs() returns a named list

with one list element per germline db. Each list element is an integer matrix that

indicates the number of germline sequences per locus and region type.

Ignored if names.only is set to TRUE.

db\_name For use\_germline\_db():

NULL or a single string specifying the name of the cached germline db to use.

Use list\_germline\_dbs() to list all the cached germline dbs.

If set to NULL (the default), then use\_germline\_db() returns the name of the cached germline db that is currently in use, if any. Otherwise it raises an error.

For load\_germline\_db():

A single string specifying the name of the cached germline db from which to load the V, D, and/or J regions. Use list\_germline\_dbs() to list all the cached

germline dbs.

verbose If set to TRUE, then use\_germline\_db() will display some information about

its internal operations.

region\_types The types of regions (V, D, and/or J) to load from the database. Specified as a

single string (e.g. "DJ") or as a character vector of single-letter elements (e.g. c("D", "J")). By default (i.e. when region\_types is NULL), all the regions are

returned.

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#### **Details**

The **igblastr** package provides a small set of utilities to manage the cached germline and C-region databases to use with igblastn().

Terminology:

- A *cached germline db* contains the nucleotide sequences of the V, D, and J gene regions for a given organism.
- A *cached C-region db* contains the nucleotide sequences of the C regions (i.e. constant gene regions) for a given organism.

This man page documents the basic utilities to operate on cached germline dbs: list\_germline\_dbs(), use\_germline\_db(), and load\_germline\_db().

The basic utilities to operate on cached C-region dbs are documented in the man page for list\_c\_region\_dbs.

#### Value

list\_germline\_dbs() returns the list of all cached germline dbs in a data.frame with one db per row (if names.only is FALSE, which is the default), or in a character vector (if names.only is TRUE). Columns V, D, J in the data.frame indicate the number of germline sequences for each region in each db.

Built-in dbs are prefixed with an underscore (\_). Note that the germline built-in dbs from AIRR were obtained from https://ogrdb.airr-community.org/germline\_sets/Homo%20sapiens and https://ogrdb.airr-community.org/germline\_sets/Mus%20musculus and included in the igblastr package on the date indicated by the suffix of the db name.

When called with no argument, use\_germline\_db() returns a single string containing the name of the cached germline db currently used by igblastn() if any, or it raises an error if no germline db has been selected yet.

When called with the db\_name argument, use\_germline\_db(db\_name) returns db\_name invisibly. load\_germline\_db() returns the nucleotide sequences from the specified germline db in a named DNAStringSet object.

# See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- install\_IMGT\_germline\_db to install a germline db from IMGT.
- use\_c\_region\_db to select the cached C-region db to use with igblastn().
- DNAStringSet objects in the **Biostrings** package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

```
if (!has_igblast()) install_igblast()

## Get list of built-in germline dbs only.
list_germline_dbs(builtin.only=TRUE)
list_germline_dbs(builtin.only=TRUE, names.only=TRUE) # db names only

## Long listing:
list_germline_dbs(long.listing=TRUE)
```

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```
if (IMGT_is_up()) {
  ## Install Mouse germline db from IMGT (Perl required!):
  install_IMGT_germline_db("202506-1", "Mus_musculus", force=TRUE)
  list_germline_dbs() # all germline dbs
  ## Select germline db to use with igblastn():
  db_name <- "IMGT-202506-1.Mus_musculus.IGH+IGK+IGL"</pre>
  use_germline_db(db_name) # select germline db to use
  use_germline_db() # get current selection
  ## Load germline sequences:
  load_germline_db(db_name)
  load_germline_db(db_name, region_types="D")
  load_germline_db(db_name, region_types="DJ")
}
```

OAS-utils

Download and manipulate OAS data

# **Description**

Some utility functions to query the Observed Antibody Space database, a.k.a. OAS, and to download and manipulate data from OAS.

```
OAS's homepage: https://opig.stats.ox.ac.uk/webapps/oas/
```

Note that OAS has two databases: the "Unpaired Sequences" database and the "Paired Sequences" database. Some of the utilities documented in this man page only work on data coming from the latter.

# Usage

```
## Read metadata/data from a single OAS unit file:
read_OAS_csv_metadata(file)
read_OAS_csv(file, skip=1, ...)
extract_sequences_from_paired_OAS_df(df, add.prefix=FALSE)
## Basic query of OAS website:
list_paired_OAS_studies(as.df=FALSE, recache=FALSE)
list_paired_OAS_units(study, as.df=FALSE, recache=FALSE)
download_paired_OAS_units(study, units=NULL, destdir=".", ...)
## Read metadata/data from a batch of downloaded OAS unit files:
extract_metadata_from_OAS_units(dir=".", pattern="\\.csv\\.gz$")
extract_sequences_from_paired_OAS_units(dir=".", pattern="\\.csv\\.gz$")
```

#### **Arguments**

file A single string that is the path to an OAS unit file.

The number of lines of the data file to skip before beginning to read data. The skip first line in an OAS unit file contains metadata in JSON format, so must always

be skipped.

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For read\_OAS\_csv(): Extra arguments to be passed to the internal call to read.table(). . . .

See ?read. table in the **utils** package for more information.

For download\_paired\_OAS\_units(): Extra arguments to be passed to the internal call to download.file(). See ?download.file in the utils package for

more information.

df The data.frame or tibble returned by read\_OAS\_csv().

TRUE or FALSE. Should the names on the returned DNAStringSet object be the add.prefix

original sequence ids as-is (this is the default), or should the heavy\_chain\_ and

light\_chain\_ prefixes be added to them?

extract\_sequences\_from\_paired\_OAS\_df() returns a DNAStringSet object

with the sequence ids as names. The sequence ids are obtained from the sequence\_id\_heavy

and sequence\_id\_light columns of the supplied data.frame or tibble. By default, they are propagated as-is to the DNAStringSet object, which makes it difficult to recognize which chain (heavy or light) the antibody sequences are coming from. Setting add.prefix to TRUE will add the heavy\_chain\_ or light\_chain\_ prefix to the names on the DNAStringSet object, hence making

it easy to identify which chain a given antibody sequence is coming from.

TRUE or FALSE. By default, i.e. when as.df is FALSE, list\_paired\_OAS\_studies()

and list\_paired\_OAS\_units() return the list of studies or units in a character vector. Alternatively you can set as.df to TRUE to get the list in a 3column data.frame that contains a directory index as displayed at https:// opig.stats.ox.ac.uk/webapps/ngsdb/paired/orathttps://opig.stats.

ox.ac.uk/webapps/ngsdb/paired/Jaffe\_2022/csv/.

TRUE or FALSE. list\_paired\_OAS\_studies() and list\_paired\_OAS\_units() recache

> both cache the information retrieved from OAS website for the duration of the R session (note that this caching is done in memory so it does not persist across sessions). Set recache to TRUE to force a new retrieval (and recaching) of the

results.

study A single string containing the name of a study as returned by list\_paired\_OAS\_studies().

units NULL, or a character vector that must be a subset of list\_paired\_OAS\_units(study)

in which case the download will be restricted to these units only.

destdir A single string that is the path to the directory where the OAS unit files are to be

downloaded.

dir A single string that is the path to a directory containing OAS unit files. This will

typically be the same as destdir above if the unit files were downloaded with

download\_paired\_OAS\_units().

pattern Regular expression passed to the internal call to list.files() to obtain the list

of OAS unit files located in dir. No reason to change this unless you know what

you are doing.

# **Details**

as.df

OAS delivers data in the form of OAS unit files. These files are typically obtained by running the bulk\_download. sh script that OAS generates based on one's search criteria. They are compressed CSV (comma-separated values) files with the .csv.gz extension.

OAS unit files can vary a lot in size: from only a few KB to 25 MB or more.

The first line in an OAS unit file contains metadata in JSON format (which means that these files cannot strictly be considered CSV files).

The CSV data is MiAIRR-compliant (see The "MiAIRR format" paper in the References section below).

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#### Value

read\_OAS\_csv\_metadata() extracts the metadata from the specified OAS unit file and returns it in a named list.

read\_OAS\_csv() extracts the data from the specified OAS unit file and returns it in a tibble. The tibble has 1 row per antibody sequence if the data is unpaired (i.e. comes from the "Unpaired Sequences" database), or 1 row per sequence pair if the data is paired (i.e. comes from the "Paired Sequences" database).

extract\_sequences\_from\_paired\_OAS\_df() returns the sequence pairs in a named DNAStringSet object where the names are the sequence ids. See add.prefix above for how the sequence ids are obtained.

list\_paired\_OAS\_studies() returns the list of studies that populate the "Paired Sequences" database in a character vector. This list can be seen here: https://opig.stats.ox.ac.uk/webapps/ngsdb/paired/.

list\_paired\_OAS\_units() returns the list of all the OAS unit files that belong to a given study from the "Paired Sequences" database.

download\_paired\_OAS\_units() returns an invisible NULL.

extract\_metadata\_from\_OAS\_units() returns the metadata of all the OAS unit files found in the specified directory in a data.frame with 1 row per file.

extract\_sequences\_from\_paired\_OAS\_units() extracts the sequence pairs from all the OAS unit files found in the specified directory and returns them in a named DNAStringSet object where the names are the sequence ids. The sequence ids are obtained by prefixing the original sequence ids found in the files with the name of the unit followed by \_heavy\_chain\_ or \_light\_chain\_.

#### References

• The OAS paper:

Tobias H. Olsen, Fergus Boyles, Charlotte M. Deane. Observed Antibody Space: A diverse database of cleaned, annotated, and translated unpaired and paired antibody sequences. Protein Science (2021). https://doi.org/10.1002/pro.4205

• The "MiAIRR format" paper:

Rubelt, F., Busse, C., Bukhari, S. et al. Adaptive Immune Receptor Repertoire Community recommendations for sharing immune-repertoire sequencing data. Nat Immunol 18, 1274–1278 (2017). https://doi.org/10.1038/ni.3873

# See Also

- OAS's homepage at: https://opig.stats.ox.ac.uk/webapps/oas/
- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- tibble objects implemented in the tibble package.
- DNAStringSet objects implemented in the **Biostrings** package.

```
list_paired_OAS_studies()
list_paired_OAS_units("Eccles_2020")
## Import all the pairs of antibody sequences from the Eccles_2020 study:
```

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```
download_dir <- tempdir()
download_paired_OAS_units("Eccles_2020", destdir=download_dir)

metadata <- extract_metadata_from_OAS_units(download_dir)
metadata # data.frame with 1 row per unit file

sequences <- extract_sequences_from_paired_OAS_units(download_dir)
sequences # DNAStringSet object

## Odd indices correspond to heavy chain sequences and even indices
## to light chain sequences:
head(names(sequences))

sequences[1:2] # 1st pair
sequences[3:4] # 2nd pair
sequences[5:6] # 3rd pair
# etc...</pre>
```

outfmt7-utils

Handle igblastn output format 7

# **Description**

Some utilities to handle igblastn output format 7.

# Usage

```
list_outfmt7_specifiers()
parse_outfmt7(out_lines)
```

# **Arguments**

```
out_lines The character vector returned by igblatsn(query, outfmt=7, parse.out=FALSE, ...).
```

# Value

list\_outfmt7\_specifiers() returns the list of format specifiers supported by igblastn() formatting option 7.

parse\_outfmt7(out\_lines) returns the parsed form of out\_lines in a list.

# See Also

- The igblastn function to run the igblastn *standalone executable* included in IgBLAST from R. This is the main function in the **igblastr** package.
- IgBLAST is described at https://pubmed.ncbi.nlm.nih.gov/23671333/.

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```
if (!has_igblast()) install_igblast()
## Files 'heavy_sequences.fasta' and 'light_sequences.fasta' included
## in igblastr contain 250 paired heavy- and light- chain sequences (125
## sequences in each file) downloaded from OAS (the Observed Antibody
## Space database):
filenames <- paste0(c("heavy", "light"), "_sequences.fasta")</pre>
query <- system.file(package="igblastr", "extdata", filenames)</pre>
## Keep only the first 10 sequences from each file:
query <- c(head(readDNAStringSet(query[[1L]]), n=10),</pre>
          head(readDNAStringSet(query[[2L]]), n=10))
## Select the germline and C-region dbs to use with igblastn():
use_germline_db("_AIRR.human.IGH+IGK+IGL.202501")
use_c_region_db("_IMGT.human.IGH+IGK+IGL.202412")
## -----
## FIRST igblastn RUN: GET OUTPUT IN FORMAT 7
## For this first run we specify 'outfmt=7' and 'parse.out=FALSE':
out_lines <- igblastn(query, outfmt=7, parse.out=FALSE)</pre>
out_lines # raw output
out <- parse_outfmt7(out_lines) # parse the output</pre>
## Output contains one record per query sequence:
length(out$records) # 20
## Each record can have 5 or 6 sections:
## 1. query_details
## 2. VDJ_rearrangement_summary
## 3. VDJ_junction_details
## 4. subregion_sequence_details (can be missing)
## 5. alignment_summary
## 6. hit_table
## Taking a close look at the first record:
rec1 <- out$records[[1]]</pre>
qseqid(rec1)
              # query sequence id associated with this record
rec1$hit_table # data.frame with the standard columns
## -----
## SECOND igblastn RUN: GET OUTPUT IN CUSTOMIZED FORMAT 7
## -----
## For this second run we request a customized format 7 by supplying
## space delimited format specifiers:
outfmt <- "7 qseqid sseqid pident nident length score"
out <- igblastn(query, outfmt=outfmt)</pre>
```

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