

# Package ‘rBLAST’

April 1, 2025

**Title** R Interface for the Basic Local Alignment Search Tool

**Description** Seamlessly interfaces the Basic Local Alignment Search Tool (BLAST) to search genetic sequence data bases. This work was partially supported by grant no. R21HG005912 from the National Human Genome Research Institute.

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**Depends** Biostrings (>= 2.26.2)

**Imports** methods, utils, BiocFileCache

**Suggests** knitr, rmarkdown, testthat

**URL** <https://github.com/mhahsler/rBLAST>

**BugReports** <https://github.com/mhahsler/rBLAST/issues>

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**License** GPL-3

**VignetteBuilder** knitr

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**Author** Michael Hahsler [aut, cre] (<<https://orcid.org/0000-0003-2716-1405>>),  
Nagar Anurag [aut]

**Maintainer** Michael Hahsler <[mhahsler@lyle.smu.edu](mailto:mhahsler@lyle.smu.edu)>

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rBLAST-package	<i>rBLAST: R Interface for the Basic Local Alignment Search Tool</i>
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### Description

Seamlessly interfaces the Basic Local Alignment Search Tool (BLAST) to search genetic sequence data bases. This work was partially supported by grant no. R21HG005912 from the National Human Genome Research Institute.

### Citation Request

Cite the use of this package as:

Hahsler M, Nagar A (2019). rBLAST: R Interface for the Basic Local Alignment Search Tool. R package version 0.99.2, URL: <https://github.com/mhahsler/rBLAST>.

### Author(s)

**Maintainer:** Michael Hahsler <[mhahsler@lyle.smu.edu](mailto:mhahsler@lyle.smu.edu)> ([ORCID](#))

Authors:

- Nagar Anurag

### See Also

Useful links:

- <https://github.com/mhahsler/rBLAST>
- Report bugs at <https://github.com/mhahsler/rBLAST/issues>

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blast	<i>Basic Local Alignment Search Tool (BLAST)</i>
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### Description

Open a BLAST database and execute `blastn` (`blastp` or `blastx`) from `blast+` to find sequences matches.

**Usage**

```
blast(db = NULL, remote = FALSE, type = "blastn")

blast_help(type = "blastn")

## S3 method for class 'BLAST'
print(x, info = TRUE, ...)

## S3 method for class 'BLAST'
predict(
  object,
  newdata,
  BLAST_args = "",
  custom_format = "",
  verbose = FALSE,
  keep_tmp = FALSE,
  ...
)

has_blast()
```

**Arguments**

db	the database file to be searched (without file extension).
remote	logical execute the query remotely on the NCBI server. db needs to be the name of a database available in the server.
type	BLAST program to use (e.g., blastn, blastp, blastx).
info	show additional data base information.
...	additional arguments are ignored.
object, x	An open BLAST database as a BLAST object created with <code>blast()</code> .
newdata	the query as an object of class <code>Biostrings::XStringSet</code> .
BLAST_args	additional arguments in command-line style.
custom_format	custom format specified by space delimited format specifiers.
verbose	logical; print progress and debugging information.
keep_tmp	logical; keep temporary files for debugging.

**Value**

- `blast()` returns a BLAST database object which can be used for queries (via `predict`).
- `predict` returns a data.frame containing the BLAST results.
- `has_blast()` returns TRUE if the blast software installation can be found and FALSE otherwise.

**Installing BLAST+**

The BLAST+ software needs to be installed on your system. Installation instructions are available in this package's `INSTALL` file and at <https://www.ncbi.nlm.nih.gov/books/NBK569861/>.

R needs to be able to find the executable. After installing the software, try in R

```
Sys.which("blastn")
```

If the command returns "" instead of the path to the executable, then you need to set the environment variable called PATH. In R

```
Sys.setenv(PATH = paste(Sys.getenv("PATH"),
  "path_to_your_BLAST_installation", sep=.Platform$path.sep))
```

## BLAST Databases

You will also need a database. NCBI BLAST databases are updated daily and may be downloaded via FTP from <https://ftp.ncbi.nlm.nih.gov/blast/db/>. See `blast_db_cache()` on how to manage a local cache of database files.

BLAST databases are a set of database files with different extensions. All files start with the same database name. For example, `16S_ribosomal_RNA.tar.gz` contains files starting with `16S_ribosomal_RNA` which is the database name used for calling `blast()`.

Large databases are separated into several archives numbered `00`, `01`, etc. Download all archives and extract the files in the same directory. All files will have a common name which is the database name used for calling `blast()`.

## Author(s)

Michael Hahsler

## References

BLAST Help - BLAST+ Executable: <https://blast.ncbi.nlm.nih.gov/doc/blast-help/downloadblastdata.html>

BLAST Command Line Applications User Manual, <https://www.ncbi.nlm.nih.gov/books/NBK279690/>

## See Also

Other blast: `blast_db_cache()`, `makeblastdb()`

## Examples

```
## check if blastn is correctly installed. Should return the path to the
## executable
Sys.which("blastn")

## only run if blast is installed
if (has_blast()) {
  ## check version you should have version 1.8.1+
  system2("blastn", "-version")

  ## download the latest version of the 16S Microbial
  ## rRNA data base from NCBI using the local cache
  tgz_file <- blast_db_get("16S_ribosomal_RNA.tar.gz")

  ## extract the database files
  untar(tgz_file, exdir = ".16S_rRNA_DB")

  ## Note the database file can also downloaded without using a
  ## cache using download.file
  # download.file(paste("https://ftp.ncbi.nlm.nih.gov/blast/db",
  # "16S_ribosomal_RNA.tar.gz", sep = "/"),
  # "16S_ribosomal_RNA.tar.gz", mode = "wb")
```

```

# untar("16S_ribosomal_RNA.tar.gz", exdir = "./16S_rRNA_DB")

## A BLAST database is just a set of files. It is a good idea to
## organize the files in a directory.
list.files("./16S_rRNA_DB")

## load a BLAST database (replace db with the location + name of
## the BLAST DB without the extension)
bl <- blast(db = "./16S_rRNA_DB/16S_ribosomal_RNA")
bl

## read a single example sequence to BLAST
seq <- readRNAStringSet(system.file("examples/RNA_example.fasta",
  package = "rBLAST"
))[1]
seq

## query a sequence using BLAST
cl <- predict(bl, seq)
cl[1:5, ]

## Pass on BLAST arguments (99% identity) and use a custom format
## (see BLAST documentation)
fmt <- paste(
  "qaccver saccver pident length mismatch gapopen qstart qend",
  "sstart send eval evalue bitscore qseq sseq"
)
cl <- predict(bl, seq,
  BLAST_args = "-perc_identity 99",
  custom_format = fmt
)
cl

## cleanup the example: delete the database files
unlink("./16S_rRNA_DB", recursive = TRUE)
}

```

blast\_db\_cache

*Manage BLAST Database Downloads using BioCFileCache***Description**

Use [BioCFileCache::BioCFileCache](#) to manage local copies of BLAST database downloads. NCBI BLAST databases are updated daily and may be downloaded via FTP from <https://ftp.ncbi.nlm.nih.gov/blast/db/>.

**Usage**

```
blast_db_cache()
```

```
blast_db_get(
  file = "16S_ribosomal_RNA.tar.gz",
  baseURL = "https://ftp.ncbi.nlm.nih.gov/blast/db/",
  check_update = TRUE,

```

```
    verbose = TRUE
  )
```

### Arguments

file	the filename of the database.
baseURL	URL to download blast databases from. The default is NCBI's ftp server.
check_update	logical; update the local cache if there is a newer version of the file available on the server. This may take some time.
verbose	logical; display download information.

### Details

The package maintains its own local cache which can be accessed using `blast_db_cache()`.

### Value

- `blast_db_cache()` returns the path to the local `BiocFileCache::BiocFileCache` cache.
- `blast_db_get()` returns the file path to a downloaded BLAST database file.

### Author(s)

Michael Hahsler

### See Also

Other blast: `blast()`, `makeblastdb()`

### Examples

```
## get a database file (will be downloaded if the
##   local copy is not up-to-date)
db_16S <- blast_db_get("16S_ribosomal_RNA.tar.gz")
db_16S

## directly interacting with the local cache
library(BiocFileCache)

## show the package's cache directory
local_cache <- blast_db_cache()
local_cache

## bfc functions can be used to manage the local cache
bfcinfo(local_cache)
```

---

`makeblastdb`*Create BLAST Databases*

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## Description

Call the `makeblastdb` utility to create a BLAST database from a FASTA file.

## Usage

```
makeblastdb(  
    file,  
    db_name = NULL,  
    dbtype = "nucl",  
    hash_index = TRUE,  
    args = "",  
    verbose = TRUE  
)
```

## Arguments

<code>file</code>	input file/database name. <b>Note</b> that the filename and path cannot contain whitespaces.
<code>db_name</code>	name of the database (files).
<code>dbtype</code>	molecule type of target db ("nucl" or "prot").
<code>hash_index</code>	logical; create index of sequence hash values.
<code>args</code>	string including additional arguments passed on to <code>makeblastdb</code> .
<code>verbose</code>	logical; show the progress report produced by <code>makeblastdb</code> ?

## Details

R needs to be able to find the executable (mostly an issue with Windows). Try `Sys.which("makeblastdb")` to see if the program is properly installed.

Use `blast_help("makeblastdb")` to see all possible extra arguments. Arguments need to be formatted in exactly the way as they would be used for the command line tool.

## Value

Nothing but creates a BLAST database directory.

## Author(s)

Michael Hahsler

## See Also

Other blast: [blast\(\)](#), [blast\\_db\\_cache\(\)](#)

**Examples**

```
## check if makeblastdb is correctly installed
Sys.which("makeblastdb")

## only run if blast is installed
if (has_blast()) {
  ## see possible arguments
  blast_help("makeblastdb")

  ## read some example sequences
  seq <- readRNAStringSet(system.file("examples/RNA_example.fasta",
    package = "rBLAST"
  ))

  ## 1. write the FASTA file
  writeXStringSet(seq, filepath = "seqs.fasta")

  ## 2. make database
  makeblastdb(file = "seqs.fasta", db_name = "db/small", dbtype = "nucl")

  ## 3. open database
  db <- blast("db/small")
  db

  ## 4. perform search (first sequence in the db should be a perfect match)
  predict(db, seq[1])

  ## clean up
  unlink("seqs.fasta")
  unlink("db", recursive = TRUE)
}
```

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