

# Package: hoardeR (via r-universe)

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**Type** Package

**Title** Collect and Retrieve Annotation Data for Various Genomic Data  
Using Different Webservices

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**Description** Cross-species identification of novel gene candidates using the NCBI web service is provided. Further, sets of miRNA target genes can be identified by using the targetscan.org API.

**VignetteBuilder** knitr

**License** GPL (>= 2)

**Repository** <https://fischuu.r-universe.dev>

**RemoteUrl** <https://github.com/fischuu/hoarder>

**RemoteRef** HEAD

**RemoteSha** b463fc906c3610f8cb003b8563f6cd10e065b1b4

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hoardeR-package	<i>Collect and Retrieve Annotation Data for Various Genomic Data Using Different Web Services.</i>
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## Description

The hoardeR package is designed for collecting, retrieving and transforming data from various sources. The current main focus is on setting up a connection to the NCBI Blast service. Also, the gene information for Ensembl Genes can be retrieved from NCBI. Methods for visualizing the results are also provided. The latest developer version of the package can be downloaded from <https://github.com/fischuu/hoardeR>

## Details

Package:	hoardeR
Type:	Package
Version:	0.10
Date:	2024-04-08
License:	GPL
LazyLoad:	yes

## Author(s)

Daniel Fischer, Anu Sironen

Maintainer: Daniel Fischer <daniel.fischer@luke.fi>

blastSeq

*Sending Genomic Sequences to NCBI Blast service***Description**

This function sends genomic sequences to the NCBI Blast service.

**Usage**

```
blastSeq(seq, n_blast=20, delay_req=10, delay_rid=60, email=NULL,
         xmlFolder=NULL, logFolder=NULL, keepInMemory=FALSE,
         database="refseq_genomes", verbose=TRUE, createLog=TRUE)
```

**Arguments**

seq	The fasta sequence that should be blasted (String).
n_blast	Amount of parallel blast requests, in case seq is a vector.
delay_req	Seconds between the single Blast requests.
delay_rid	Seconds between the single result requests.
email	User email, required information from NCBI (String).
xmlFolder	Path to the result folder.
logFolder	Path to the log folder.
keepInMemory	Logical, shall the results be kept in the memory.
database	The NCBI database to use.
verbose	Shall the program give extensive feedback.
createLog	Create log files, needed for continuing a crashed program.

**Details**

This function sends fasta sequences to the NCBI blast service. The defaults for the delays are required by NCBI and must not be smaller than the default values. Also, NCBI asks the user to provide an email address.

The input seq can be a vector of strings. In that case the sequences are one after another processed. The option n\_blast sets then the upper threshold of how many blast requests are send to the NCBI Blast service at a time and kept running there parallel. It is here in the users obligation not to misuse the service with too many parallel requests.

The xmlFolder parameter specifies the folder to where the XML results will be stored. In case the folder does not exist, R will create it.

In case the option keepInMemory is set to TRUE the Blast results will be kept in memory, otherwise they will be just written to the HDD. Especially if many sequences are send to the blast service it is recommended to drop the result from the memory, meaning to set the option keepInMemory=FALSE. The option keepInMemory=TRUE is currently still under development and should not be used.

If log files should be written (createLog=TRUE) a log path should be given in logPath. However, if a xmlPath is given and the option createLog=TRUE is set, then the log folder will be automatically created in the parental folder of the xmlFolder and is called logs.

**Value**

An xml file that contains the the NCBI result.

**Author(s)**

Daniel Fischer

**Examples**

```
## Not run:
blastSeq("ACGTGCATCGACTAGCTACGACTACGACTATC", email="my.name@somewhere.com")

## End(Not run)
```

---

coverageDensity	<i>Calculation of the coverage density</i>
-----------------	--

---

**Description**

Calculates the coverage density.

**Usage**

```
coverageDensity(folder, chr=c(1:22,"X","Y","MT"), chr.length=NULL,
                posneg=FALSE, verbose=TRUE, use.sqrt=FALSE,
                kernel.package="slideWindowSum",
                step.size=50000, window.size=100000, bw=100)
```

**Arguments**

folder	folder with bam files
chr	Chromosome names to be plotted.
chr.length	Length of chromosome
posneg	Logical, plot pos and neg strand
verbose	Logical, verbose output
use.sqrt	Logical, apply sqrt transformation
kernel.package	Class of kernel smoother
step.size	Step size in bases
window.size	Window size in bases
bw	Bandwidth parameter

**Details**

This function calculates the coverage of bam-files

**Author(s)**

Daniel Fischer

---

`findSpecies`                      *Search in the species' Object.*

---

**Description**

This function output rows from the `species` object that contain a certain string.

**Usage**

```
findSpecies(string)
```

**Arguments**

`string`                      Search string.

**Details**

This function output rows from the `species` object that contain a certain string. It uses the `grep1` function to identify the corresponding rows.

**Value**

A `data.frame`.

**Author(s)**

Daniel Fischer

**See Also**

[species](#), [grep1](#)

**Examples**

```
findSpecies("cattle")
```

---

getAnnotation	<i>Downloading or Importing of Annotation Data</i>
---------------	--

---

### Description

This function downloads (if needed) the annotation file from a given species from NCBI and loads it into the namespace.

### Usage

```
getAnnotation(species=NULL, assembly=NULL, annotationFolder=NULL,  
              type="gff3", verbose)
```

### Arguments

species	The scientific name of the species (String).
assembly	The NCBI assembly version.
annotationFolder	The folder where the file will be stored.
type	The file extension/format of the annotation file.
verbose	Logical, if function gives feedback.

### Details

This function downloads for a given species the annotation file, as provided from NCBI. The main parameters basically define the URL, where the file is located. The file is then downloaded into the folder, provided in `annotationFolder` and then imported to the namespace.

If a file has been downloaded previously, it will be loaded directly from that folder. In case the user wants to use an annotation that is not provided by NCBI, the corresponding files can also be placed into the same folder, following the naming scheme as suggested from the function and the function will load it from there.

### Value

A `data.table` with the annotation information.

### Author(s)

Daniel Fischer

**Examples**

```
## Not run:
susScrofa <- getAnnotation(species = "Sus scrofa",
                           annotationFolder="/home/user/annotation")

homoSapiens <- getAnnotation(species = "Homo sapiens",
                             annotationFolder="/home/user/annotation")

## End(Not run)
```

---

`getEnsgInfo`*Retrieve Gene Information From the NCBI Database.*

---

**Description**

This function retrieves for a given Ensembl Number the corresponding information from the NCBI database.

**Usage**

```
getEnsgInfo(ensg)
```

**Arguments**

`ensg` Ensembl ID (String).

**Details**

This function retrieves for a given Ensembl Number the corresponding information from the NCBI database. The object `ensg` can also be a vector of Ensembl IDs.

**Value**

A matrix with information.

**Author(s)**

Daniel Fischer

**Examples**

```
## Not run:
ensg <- c("ENSG00000174482", "ENSG00000113494")
getEnsgInfo(ensg)

## End(Not run)
```

---

`getFastaFromBed`*Get fasta information based on locations in bed-format*

---

**Description**

For a given fasta and a bed file this function can extract the nucleotide sequences and stores them as fasta file.

**Usage**

```
getFastaFromBed.bed, species=NULL, assembly = NULL, fastaFolder=NULL,  
verbose=TRUE, export=NULL, fileName=NULL)
```

**Arguments**

<code>bed</code>	The location in bed format, see details.
<code>species</code>	Define the species.
<code>assembly</code>	Assembly identifier.
<code>fastaFolder</code>	Location of the fasta files.
<code>verbose</code>	Logical, should informative status updates be given.
<code>export</code>	Foldername.
<code>fileName</code>	Filename to store the FA object.

**Details**

Function expects as an input a data.frame in bed format. This means, the first column should contain the chromosome, the second the start-coordinates, the third the end-coordinates. The fourth column contains the ID of the loci.

If a standard species is used (as defined in the species data frame), the function automatically downloads the required files from NCBI, takes the loci and extracts then the nucleotide sequences from it. If the corresponding assembly is not available from NCBI an own fasta file can be provided. For that the fa-file needs to be in the fastaFolder and follow the same naming system as the NCBI files are labelled. In that case, the function suggests the correct filename for an unknown assembly.

The export function, specifies then a folder to where the fasta file should be stored. If no filename is provided, the filename is then the object name passed to the bed function.

**Value**

An fa object containing the nucleotide sequences in fasta format.

**Author(s)**

Daniel Fischer



**Examples**

```
## Not run:

myBed <- data.frame(chr=c(1,2),
                    start=c(235265,12356742),
                    end=c(435265,12386742),
                    gene=c("LOC1", "LOC2"))

myFA <- getFastaFromBed(myBed, species="Homo sapiens", fastaFolder="/home/user/fasta/", export=TRUE)

## End(Not run)
```

---

getGeneLocation	<i>Extracting Gene Locations</i>
-----------------	----------------------------------

---

**Description**

This function extracts the gene locations from an imported gtf file.

**Usage**

```
getGeneLocation(gtf)
```

**Arguments**

gtf                    An imported gtf object.

**Details**

This function extracts the information from an imported gtf object.

**Value**

A matrix.

**Author(s)**

Daniel Fischer

**Examples**

```
## Not run:
getGeneLocation(gtf)

## End(Not run)
```

---

`getGeneSeq`*Extracting a gene sequence from NCBI database.*

---

**Description**

This function retrieves a gene sequence from the NCBI database.

**Usage**

```
getGeneSeq(chr, start, end, organism)
```

**Arguments**

<code>chr</code>	Chromosome number, numeric/string
<code>start</code>	Start position, numeric
<code>end</code>	End position, numeric
<code>organism</code>	Name of the organism, string

**Details**

Extracting a gene sequence from NCBI database. For a list of available organism, visit <http://genome.ucsc.edu/cgi-bin/das/dsn>. All id="." field are available.

**Value**

A string that contains the genomic sequence.

**Author(s)**

Daniel Fischer

**Examples**

```
## Not run:  
# Extracting for Sus Scrofa, build version 3:  
getGeneSeq(1,2134,14532,"susScr3")  
getGeneSeq(10,1233312,1233350,"hg38")  
  
## End(Not run)
```

---

getSequenceFromNCBI     *Extracts a sequence from the NCBI webpage*

---

**Description**

Retrieve a sequence from the NCBI webpage

**Usage**

```
getSequenceFromNCBI(id, file=NULL)
```

**Arguments**

id	The gene identifier
file	File name to where the sequence shall be stored

**Details**

This function extracts the sequence for a given identifier and then stores, if requested the sequence to the HDD.

**Author(s)**

Daniel Fischer

---

intersectXMLAnnot     *Intersect XML object with annotation object*

---

**Description**

For an annotation object this function intersects the loci of it with the output of the tableSpecies function.

**Usage**

```
intersectXMLAnnot(tabSpecies, annot, level="gene", flanking=NULL)
```

**Arguments**

tabSpecies	The table with locations from tableSpecies.
annot	The annotation object.
level	The level of intersection.
flanking	Allowed flanking space for intersection.

## Details

Function expects as an input table from `tableSpecies` with the option `locations=TRUE`. Further, it needs an annotation object, as provided by the `getAnnotation` function. With that it intersects then the loci on the level as specified in `level`. Currently only "gene" is supported.

The `flanking` option allows for flanking space up- and down-stream of the genes. This is especially then useful if the novel gene candidates are in the extension of known genes (e.g. responsible for regulation or if they are novel exons.)

## Value

A table with intersection loci.

## Author(s)

Daniel Fischer

## Examples

```
## Not run:

pigHits <- tableSpecies(xmls, species="Sus scrofa", locations = TRUE)
ssannot <- getAnnotation(species = "Sus scrofa", annotationFolder="/home/user/annotation")
pigInter <- list()
for(i in 1:nrow(pigHits)){
  pigInter[[i]] <- intersectXMLAnnot(pigHits[i,], ssannot)
}

## End(Not run)
```

---

plotCoverage

*Plots a coverage density object*

---

## Description

Plots a coverage density object.

## Usage

```
plotCoverage(x, use.sqrt=TRUE)
```

## Arguments

x	A coverage density object
use.sqrt	Logical, use sqrt scale?

**Details**

This function plots the coverage of bam-files

**Author(s)**

Daniel Fischer

---

plotHit

*Visualization of a cross-species hit*

---

**Description**

For each cross-species hit the function plots the similarity within that area together with an optional annotation and coverage track.

**Usage**

```
plotHit(hits, flanking=1, window=NULL, annot=TRUE, coverage=FALSE,
        smoothPara=NULL, diagonal=0.25, verbose=TRUE, output=FALSE,
        hitSpecies=NULL, hitSpeciesAssembly=NULL, origSpecies=NULL,
        origSpeciesAssembly=NULL, fastaFolder=NULL, origAnnot=NULL,
        hitAnnot=NULL, nTick=5, which=NULL, figureFolder=NULL,
        figurePrefix=NULL, indexOffset=0, bamFolder=NULL, bamFiles=NULL,
        groupIndex=NULL, groupColor=NULL, countWindow=NULL)
```

**Arguments**

hits	The hit object to be plotted.
flanking	Allowed flanking site in Mb.
window	Moving window size of similarity measure.
annot	Logical, add annotation track
coverage	Logical, add coverage track
smoothPara	Smoothing parameter for coverage
diagonal	Threshold for allowed diagonal similarity
verbose	Logical, shall the function give status updates
output	Logical, shall numerical results be given
hitSpecies	Scientific identifier of the hit species.
hitSpeciesAssembly	Version of the hit species assembly
origSpecies	Scientific name of the original species
origSpeciesAssembly	Version of the original species
fastaFolder	Location of the fasta files

origAnnot	Annotation object of the original species
hitAnnot	Annotation object of the hit species
nTick	Number of ticks on the annotation track
which	Which hits should be plotted
figureFolder	Folder where Figures should be stored
figurePrefix	Prefix of the figure filenames
indexOffset	Offset of the running index of the filenames
bamFolder	Folder with the bam-files
bamFiles	Filenames of the bam-files
groupIndex	Index of subgroups in the bamfiles
groupColor	Vector with colors, one for each subgroup
countWindow	Window size to count the reads from bam-files.

## Details

This function is the workhorse of hoardeR and visualizes the findings of the blast and intersection runs. It is really flexibel to handle the hits and hence there are many different options. The required options are hits, hitSpecies, origSpecies and fastaFolder.

The hit object is an object as provided by intersectXMLAnnot and contains all intersections of interest (=intersections that are in close proximity of a gene in the hit species). Naturally the hit and the original species have to be specified as well as the folder, where the required fasta files are stored, or to where they should be downloaded. If the species are the default species from Ensembl (as can be seen in the data.frame species), the annotation and assembly will be automatically downloaded to the specified location on the harddrive. Changes from that version can be adjusted with the hitSpeciesAssembly and origSpeciesAssembly options, but the filenames have still to match the convention, as they are provided by NCBI.

If in addition to the similarity also a coverage track should be added, the option coverage has to be set to TRUE. The option smoothPara sets then the level of smoothing of the coverage. By default no smoothing will be applied.

In case an annotation track is requested (annot=TRUE), the annotation objects need to be provided to the origAnnot and hitAnnot options.

The option diagonal defines the minimum level of similarity so that a (diagonal) match will be plotted. The colors are then towards green for total similarity and towards red for total disagree, based on a nucleotide mismatch matrix.

If the option verbose=TRUE is set, the function gives a verbose output while running. Further, if output=TRUE then, in addition to the figure also a data.frame with the numerical results is provided.

In case that hits contains more than one hit, the plotHit function plots for each hit a figure. In that case a folder should be provided to where the figures should be stored, this can be done with the figureFolder and figurePrefix options. In case only asserted hits of hits shall be plotted, they can be selected with the which option.

The function can also plot a coverage track over the similarity. For that, the option coverage=TRUE has to be set and a folder that contains the necessary bam-files has to be specified in bamFolder. By default all bam files in that folder are used, if only a subset is requested, the filenames can be

specified in `bamFiles`. In case several bam-files are given, the average coverage at each loci is used. Further, if the data contains subgroups (e.g. case/control), the vector `groupIndex` gives the group labels. Naturally its length should be similar to `bamFiles` (or similar to the total amount of files in the bam-folder). In case that more than one group is plotted in the coverage track, their colors can be defined in `groupColor`. Of course, this vector has to be as long as the number of groups are defined. The option `countWindow` controls the moving window length in which the number of counts is calculated. The default is the same length as the hit.

### Value

Optional, a table with intersection loci.

### Author(s)

Daniel Fischer

### Examples

```
## Not run:
pigInter.flank <- list()
for(i in 1:nrow(pigHits)){
  pigInter.flank[[i]] <- intersectXMLAnnot(pigHits[i,], ssannot, flanking=100)
}
# Basic usage:
plotHit(hits=pigInter.flank,
        flanking=100,
        hitSpecies = "Sus scrofa",
        origSpecies = "Bos taurus",
        fastaFolder = "/home/user/fasta/",
        figureFolder = "/home/user/figures/")

# Annotation tracks added:
plotHit(hits=pigInter.flank,
        flanking=100,
        hitSpecies = "Sus scrofa",
        origSpecies = "Bos taurus",
        fastaFolder = "/home/user/fasta/",
        figureFolder = "/home/user/figures/",
        origAnnot=btannot,
        hitAnnot=ssannot)

# Annotation and coverage added:
plotHit(hits=pigInter.flank,
        flanking=100,
        hitSpecies = "Sus scrofa",
        origSpecies = "Bos taurus",
        fastaFolder = "/home/daniel/fasta/",
        figureFolder = "/home/user/figures/",
        origAnnot=btannot,
        hitAnnot=ssannot,
        coverage=TRUE,
        bamFolder = "/home/users/bams/")
```

```
## End(Not run)
```

---

```
print.fa          Print an fa Object
```

---

### Description

Prints an fa object.

### Usage

```
## S3 method for class 'fa'
print(x, n=2, seq.out=50, ...)
```

### Arguments

x	Object of class fa.
n	Amount of elements to be displayed, numeric.
seq.out	Length of each element to be displayed, numeric..
...	Additional parameters.

### Details

The print function displays an fa object. By default just the first two elements with their first 50 bases are displayed. To display the full sequence, set seq.out=NULL.

### Author(s)

Daniel Fischer

---

```
species          Available species at NCBI
```

---

### Description

This is a list of all organisms/species that are provided by NCBI and hence could end up in the Blast run. Further, it defines the default versions of the assemblies that will be downloaded if no further version is specified in plotHit, getAnnotation or getFastaFromBed.

### Format

A data frame with 348 species.



**Source**

As downloaded on 05.10.2016 from  
<ftp://ftp.ncbi.nlm.nih.gov/genomes/>

**Examples**

```
data(species)
summary(species)
```

---

subDose	<i>Rewrite the Dose File from a Beagle Output</i>
---------	---

---

**Description**

This function takes a Dose Beagle output and rewrites the output.

**Usage**

```
subDose(file=NULL, vmmk=NULL, out=NULL, removeInsertions=TRUE, verbose=TRUE)
```

**Arguments**

file	Location of the original Beagle file (String).
vmmk	Location of the Variant Map Master key (String).
out	Name and location of the output file (String).
verbose	The function gives feedback.
removeInsertions	All Indels will be removed..

**Details**

This function takes a Beagle Dose file and rewrites the alleles from numerical to character, based on the information provided in a variant map master key.

**Value**

A rewritten beagle phased file.

**Author(s)**

Daniel Fischer

---

`subGprobs`*Rewrite the Gprobs File from a Beagle Output*

---

**Description**

This function takes a Gprobs Beagle output and rewrites the output.

**Usage**

```
subGprobs(file=NULL, vmmk=NULL, out=NULL, chunkSize=100000, removeInsertions=TRUE,
          verbose = TRUE, writeOut=TRUE)
```

**Arguments**

<code>file</code>	Location of the original Beagle file (String).
<code>vmmk</code>	Location of the Variant Map Master key (String).
<code>out</code>	Name and location of the output file (String).
<code>chunkSize</code>	For large Beagle files, the chunk size.
<code>removeInsertions</code>	All Indels will be removed.
<code>verbose</code>	The function gives feedback.
<code>writeOut</code>	Logical, write the output back to the HDD.

**Details**

This function takes a Beagle Gprobs file and rewrites the alleles from numerical to character, based on the information provided in a variant map master key. For larger files the function can process the rewriting in chunks in order to save memory.

**Value**

A rewritten beagle Gprobs file.

**Author(s)**

Daniel Fischer

---

subPhased	<i>Rewrite the Phased File from a Beagle Output</i>
-----------	---

---

### Description

This function takes a phased Beagle output and rewrites the output.

### Usage

```
subPhased(file=NULL, vmmk = NULL, out=NULL, chunkSize=100000, verbose=TRUE,
          removeInsertions=TRUE)
```

### Arguments

file	Location of the original Beagle file (String).
vmmk	Location of the Variant Map Master key (String).
out	Name and location of the output file (String).
chunkSize	For large Beagle files, the chunk size.
verbose	The function gives feedback.
removeInsertions	All Indels will be removed.

### Details

This function takes a Beagle phased file and rewrites the alleles from numerical to character, based on the information provided in a variant map master key. For larger files the function can process the rewriting in chunks in order to save memory.

### Value

A rewritten beagle phased file.

### Author(s)

Daniel Fischer

---

summary.fa	<i>Summarize an fa Object</i>
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---

**Description**

Summarizes and prints an fa object in an informative way.

**Usage**

```
## S3 method for class 'fa'
summary(object, ...)
```

**Arguments**

object	Object of class fa.
...	Additional parameters.

**Details**

Summary for a fa object, providing the amount of sequences, the minimum and maximum length as well as the average length.

**Author(s)**

Daniel Fischer

---

tableSpecies	<i>Tables the species in xml file</i>
--------------	---------------------------------------

---

**Description**

Tables the species in xml file

**Usage**

```
tableSpecies(xml, species=NULL, type="chr", minOutput=TRUE, exclude="",
             locations=FALSE)
```

**Arguments**

xml	The xml file.
species	Restrict species to a certain set.
type	Filter option.
minOutput	Logical, should the output be minimal.
exclude	Names of species to exclude.
locations	Logical, shall the hit locations be given as well.

### Details

Function provides a table of identified species. This table can e.g. be put into the barplot function to visualize the findings.

Further, if the option `locations` is set to `TRUE` the function not only tables the species, but also the individual locations of the hits. This output is required for the further steps. Hence, this function plays a important role in the identification pipeline.

Be default the option `type="chr"` is set so that only hits in species will full genomes will be reported. Further, the species names are intersected with the species data frame and only those that appear there are reported.

### Value

A table with the species from the XML file

### Author(s)

Daniel Fischer

### Examples

```
## Not run:
tableSpecies(xmls)
pigHits <- tableSpecies(xmls, species="Sus scrofa", locations = TRUE)

## End(Not run)
```

---

targetScan

*Retrieving miRNA target information from targetscan.org*

---

### Description

This function requests from the webpage [targetscan.org](http://targetscan.org) the stored information for mirnas.

### Usage

```
targetScan(mirna=NULL, species=NULL, release="7.1", maxOut=NULL)
```

### Arguments

<code>mirna</code>	The name of the mirna (String).
<code>species</code>	The species identifier, see details (String).
<code>release</code>	The release version of targetscan.org.
<code>maxOut</code>	The amount of target genes, default (NULL) is all.

**Details**

This function sends a miRNA name to the [targetscan.org](http://targetscan.org) webpage, retrieves the information and gives it back as a data.frame. Options for species are "Human", "Mouse", "Rat", "Chimpanzee", "Rhesus", "Cow", "Dog", "Opossum", "Chicken", "Frog".

**Value**

A data.frame with the following columns

Ortholog	The ortholog name of the target gene.
geneName	The long description of the target gene.
consSites	The total number of conserved sites.
poorlySites	The total number of poorly conserved sites.

**Author(s)**

Daniel Fischer

**References**

V. Agarwal, G. Bell, J.Nam, et al. (2015): *Predicting effective microRNA target sites in mammalian mRNAs*. *eLife*, 4, pages 1-38, doi: 10.7554/eLife.05005

**Examples**

```
## Not run:  
targetScan(mirna="miR-9-5p", species="Cow", maxOut=5)  
  
## End(Not run)
```

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