Package 'biomaRt'

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Title Interface to BioMart databases (i.e. Ensembl)

Version 2.65.16

Description In recent years a wealth of biological data has become available in public data repositories. Easy access to these valuable data resources and firm integration with data analysis is needed for comprehensive bioinformatics data analysis. biomaRt provides an interface to a growing collection of databases implementing the BioMart software suite

(<https://www.ensembl.org/info/data/biomart/index.html>). The package enables retrieval of large amounts of data in a uniform way without the need to know the underlying database schemas or write complex SQL queries. The most prominent examples of BioMart databases are maintained by Ensembl, which provides biomaRt users direct access to a diverse set of data and enables a wide range of powerful online queries from gene annotation to database mining.

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```
URL https://github.com/Huber-group-EMBL/biomaRt,
    https://huber-group-embl.github.io/biomaRt/
```

BugReports https://github.com/Huber-group-EMBL/biomaRt/issues

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Description

In recent years a wealth of biological data has become available in public data repositories. Easy access to these valuable data resources and firm integration with data analysis is needed for comprehensive bioinformatics data analysis. biomaRt provides an interface to a growing collection of databases implementing the BioMart software suite (https://www.ensembl.org/info/data/biomart/index.html). The package enables retrieval of large amounts of data in a uniform way without the need to know the underlying database schemas or write complex SQL queries. The most prominent examples of BioMart databases are maintained by Ensembl, which provides biomaRt users direct access to a diverse set of data and enables a wide range of powerful online queries from gene annotation to database mining.

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See Also

Useful links:

- https://github.com/Huber-group-EMBL/biomaRt
- https://huber-group-embl.github.io/biomaRt/
- Report bugs at https://github.com/Huber-group-EMBL/biomaRt/issues

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attributePages

Gives a summary of the attribute pages

Description

Attributes in BioMart databases are grouped together in attribute pages. The attributePages() function gives a summary of the attribute categories and groups present in the BioMart. These page names can be used to display only a subset of the available attributes in the listAttributes() function.

Usage

```
attributePages(mart)
```

Arguments

mart

object of class Mart, created with the useMart() function.

Author(s)

Steffen Durinck

Examples

```
mart <- useMart(
   "ENSEMBL_MART_ENSEMBL",
   dataset = "hsapiens_gene_ensembl"
)
attributePages(mart)</pre>
```

biomaRt-deprecated

Deprecated and defunct functions in package biomaRt

Description

These functions have been removed from biomaRt and replaced with alternatives.

Details

The following functions are defunct and no longer work; use the replacement indicated below:

```
• filterOptions: listFilterOptions()
```

• listFilterValues: listFilterOptions()

• searchFilterValues: searchFilterOptions()

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biomartCache	biomaRt result caching

Description

biomaRt makes use of a results cache to speedup execution of queries that have been run before. These functions provide details on the status of this cache, and allow it to be deleted.

Usage

```
biomartCacheClear()
biomartCacheInfo()
```

Value

These functions do not return anything and are called for their side effects. biomartCacheInfo() prints the location of the cache, along with the number of files and their total size on disk. biomartCacheClear() will delete the current contents of the cache.

Author(s)

Mike Smith

exportFASTA

Exports getSequence results to FASTA format

Description

Exports getSequence results to FASTA format

Usage

```
exportFASTA(sequences, file)
```

Arguments

sequences A data.frame that was the output of the getSequence() function

file File to which you want to write the data

Author(s)

Steffen Durinck Hugo Gruson 6 filterType

Examples

```
mart <- useMart("ENSEMBL_MART_ENSEMBL", dataset = "hsapiens_gene_ensembl")

seq <- getSequence(
  id = "BRCA1",
    type = "hgnc_symbol",
    seqType = "cdna",
    mart = mart
)
exportFASTA(seq, file = "test.fasta")</pre>
```

filterType

Displays the filter type

Description

Displays the type of the filer given a filter name.

Usage

```
filterType(filter, mart)
```

Arguments

filter A valid filter name. Valid filters are given by the listFilters() function mart object of class Mart, created using the useMart() function

Author(s)

Steffen Durinck

```
mart <- useMart("ENSEMBL_MART_ENSEMBL", dataset = "hsapiens_gene_ensembl")
filterType("chromosome_name", mart)</pre>
```

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getBM

Retrieves information from the BioMart database

Description

This function is the main biomaRt query function. Given a set of filters and corresponding values, it retrieves the user specified attributes from the BioMart database one is connected to.

Usage

```
getBM(
  attributes,
  filters = "",
  values = "",
  mart,
  checkFilters = TRUE,
  verbose = FALSE,
  uniqueRows = TRUE,
  bmHeader = FALSE,
  quote = "\"",
  useCache = TRUE
)
```

Arguments

attributes	Attributes you want to retrieve. A	a possible list of attributes can be retrieved
		45

using the function listAttributes().

filters Filters (one or more) that should be used in the query. A possible list of filters

can be retrieved using the function listFilters().

values Values of the filter, e.g. vector of affy IDs. If multiple filters are specified then

the argument should be a list of vectors of which the position of each vector

corresponds to the position of the filters in the filters argument.

mart object of class Mart, created with the useMart() function.

checkFilters Sometimes attributes where a value needs to be specified, for example upstream flank

with value 20 for obtaining upstream sequence flank regions of length 20bp, are treated as filters in BioMarts. To enable such a query to work, one must specify the attribute as a filter and set checkFilters = FALSE for the query to work.

verbose When using biomaRt in webservice mode and setting verbose to TRUE, the

XML query to the webservice will be printed.

uniqueRows If the result of a query contains multiple identical rows, setting this argument to

TRUE (default) will result in deleting the duplicated rows in the query result at

the server side.

bmHeader Boolean to indicate if the result retrieved from the BioMart server should include

the data headers or not, defaults to FALSE. This should only be switched on if the default behavior results in errors, setting to on might still be able to retrieve

your data in that case

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quote Sometimes parsing of the results fails due to errors in the Ensembl data fields

such as containing a quote, in such cases you can try to change the value of

quote to try to still parse the results.

useCache Boolean indicating whether the results cache should be used. Setting to FALSE

will disable reading and writing of the cache. This argument is likely to disap-

pear after the cache functionality has been tested more thoroughly.

Value

A data frame. There is no implicit mapping between its rows and the function arguments (e.g. filters, values), therefore make sure to have the relevant identifier(s) returned by specifying them in attributes. See Examples.

Author(s)

Steffen Durinck

Examples

getGene

Retrieves gene annotation information given a vector of identifiers

Description

This function retrieves gene annotations from Ensembl given a vector of identifiers. Annotation includes chromosome name, band, start position, end position, gene description and gene symbol. A wide variety of identifiers is available in Ensembl, these can be found with the listFilters function.

Usage

```
getGene(id, type, mart)
```

Arguments

id		e wants to	

type of identifier, possible values can be obtained by the listFilters function. Ex-

amples are entrezgene_id, hgnc_symbol (for hugo gene symbol), ensembl_gene_id,

unigene, agilentprobe, affy_hg_u133_plus_2, refseq_dna, etc.

mart object of class Mart, containing connections to the BioMart databases. You can

create such an object using the function useMart().

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Author(s)

Steffen Durinck

Examples

```
mart <- useMart("ENSEMBL_MART_ENSEMBL", dataset = "hsapiens_gene_ensembl")

# example using affy id
g <- getGene(id = "1939_at", type = "affy_hg_u95av2", mart = mart)
show(g)

# example using Entrez Gene id
g <- getGene(id = "100", type = "entrezgene_id", mart = mart)
show(g)</pre>
```

getHomologs

List homologous genes between two species.

Description

This function simplifies the querying of the Ensembl BioMart if you're trying to return the homologs for one or more gene IDs between two species.

Usage

```
getHomologs(ensembl_gene_ids, species_from, species_to)
```

Arguments

```
ensembl_gene_ids
```

Character vector. This contains the Ensembl Gene IDs that you want to find the homologs for.

```
species_from, species_to
```

Character vectors of length 1. These arguments specify the species the input IDs belong to (species_from) and the species you want to find the homologs in (species_to). These can be Ensembl genomes names e.g. "homo_sapiens" or "canis_lupus_familiaris" or common names e.g. "human" or "dog". The function will do it's best to parse common names, and will report and error if no match to an Ensembl genome can be made.

Author(s)

Mike Smith

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getLDS

Retrieves information from two linked datasets

Description

This function is the main biomaRt query function that links 2 datasets and retrieves information from these linked BioMart datasets. In Ensembl this translates to homology mapping.

Usage

```
getLDS(
  attributes,
  filters = "",
  values = "",
  mart,
  attributesL,
  filtersL = "",
  valuesL = "",
  wartL,
  verbose = FALSE,
  uniqueRows = TRUE,
  bmHeader = TRUE
)
```

Arguments

attributes	Attributes you want to retrieve of primary dataset. A possible list of attributes can be retrieved using the function listAttributes().
filters	Filters that should be used in the query. These filters will be applied to primary dataset. A possible list of filters can be retrieved using the function listFilters().
values	Values of the filter, e.g. list of affy IDs
mart	object of class Mart created with the useMart() function.
attributesL	Attributes of linked dataset that needs to be retrieved
filtersL	Filters to be applied to the linked dataset
valuesL	Values for the linked dataset filters
martL	Mart object representing linked dataset
verbose	When using biomaRt in webservice mode and setting verbose to TRUE, the XML query to the webservice will be printed. Alternatively in MySQL mode the MySQL query will be printed.
uniqueRows	Logical to indicate if the BioMart web service should return unique rows only or not. Has the value of either TRUE or FALSE
bmHeader	Boolean to indicate if the result retrieved from the BioMart server should include the data headers or not, defaults to TRUE. This should only be switched off if the default behavior results in errors, setting to off might still be able to retrieve your data in that case

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Author(s)

Steffen Durinck

Examples

```
human <- useMart(</pre>
  "ENSEMBL_MART_ENSEMBL",
  dataset = "hsapiens_gene_ensembl",
  host = "https://dec2021.archive.ensembl.org"
)
mouse <- useMart(</pre>
  "ENSEMBL_MART_ENSEMBL",
  dataset = "mmusculus_gene_ensembl",
  host = "https://dec2021.archive.ensembl.org"
getLDS(
  attributes = c("hgnc_symbol","chromosome_name", "start_position"),
  filters = "hgnc_symbol",
  values = "TP53",
  mart = human,
  attributesL = c("chromosome_name", "start_position"),
  martL = mouse
)
```

getSequence

Retrieves sequences

Description

This function retrieves sequences given the chromosome, start and end position or a list of identifiers. Using getSequence in web service mode (default) generates 5' to 3' sequences of the requested type on the correct strand.

Usage

```
getSequence(
  chromosome,
  start,
  end,
  id,
  type,
  seqType,
  upstream,
  downstream,
  mart,
  useCache = TRUE,
  verbose = FALSE
)
```

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Arguments

chromosome	Chromosome name
start	start position of sequence on chromosome
end	end position of sequence on chromosome
id	An identifier or vector of identifiers.
type	The type of identifier used. Supported types are hugo, ensembl, embl, entrezgene, refseq, ensemblTrans and unigene. Alternatively one can also use a filter to specify the type. Possible filters are given by the listFilters() function.
seqType	Type of sequence that you want to retrieve. Allowed seqTypes are given in the details section.
upstream	To add the upstream sequence of a specified number of basepairs to the output.
downstream	To add the downstream sequence of a specified number of basepairs to the output.
mart	object of class Mart created using the useEnsembl() function
useCache	If useCache = TRUE then biomaRt will try to store succesful query results on disk, and will load these if a query is run again, rather than contacting the Ensembl server.
verbose	If 'verbose = TRUE" then the XML query that was send to the webservice will be displayed.

Details

The type of sequence returned can be specified by the seqType argument which takes the following values:

- 'cdna': for nucleotide sequences
- 'peptide': for protein sequences
- '3utr': for 3' UTR sequences
- '5utr': for 5' UTR sequences
- 'gene_exon': for exon sequences only
- 'transcript_exon_intron': gives the full unspliced transcript, that is exons + introns
- 'gene_exon_intron' gives the exons + introns of a gene;'coding' gives the coding sequence only
- 'coding_transcript_flank': gives the flanking region of the transcript including the UTRs, this must be accompanied with a given value for the upstream or downstream attribute
- 'coding_gene_flank': gives the flanking region of the gene including the UTRs, this must be accompanied with a given value for the upstream or downstream attribute
- 'transcript_flank': gives the flanking region of the transcript excluding the UTRs, this must be accompanied with a given value for the upstream or downstream attribute
- 'gene_flank': gives the flanking region of the gene excluding the UTRs, this must be accompanied with a given value for the upstream or downstream attribute

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Author(s)

Steffen Durinck, Mike Smith

Examples

```
mart <- useEnsembl("ENSEMBL_MART_ENSEMBL", dataset = "hsapiens_gene_ensembl")

seq <- getSequence(
   id = "BRCA1",
    type = "hgnc_symbol",
    seqType = "peptide",
    mart = mart
)

show(seq)

seq <- getSequence(
   id = "1939_at",
    type = "affy_hg_u95av2",
   seqType = "gene_flank",
   upstream = 20,
   mart = mart
)
show(seq)</pre>
```

listAttributes

lists the attributes available in the selected dataset

Description

Attributes are the outputs of a biomaRt query, they are the information we want to retrieve. For example if we want to retrieve all EntrezGene identifiers of genes located on chromosome X, entrezgene_id will be the attribute we use in the query. The listAttributes function lists the available attributes in the selected dataset.

Usage

```
listAttributes(mart, page, what = c("name", "description", "page"))
searchAttributes(mart, pattern = ".*")
```

Arguments

mart	object of class Mart created using the useMart() function
page	Show only the attributes that belong to the specified attribute page.
what	vector of types of information about the attributes that need to be displayed. Can have values like name, description, fullDescription, page
pattern	Character vector defining the regular expression (regex) to be used for the search. If left blank the default is to use ".*" which will match everything.

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Author(s)

Steffen Durinck, Mike Smith

Examples

```
## list the available Ensembl marts and use Ensembl Genes
listEnsembl()
ensembl <- useEnsembl(
    biomart = "ENSEMBL_MART_ENSEMBL",
    dataset = 'hsapiens_gene_ensembl'
)

## list the available datasets in this Mart
listAttributes(mart = ensembl)

## the list of attributes is very long and gets truncated by R

## we can search for a term of interest to filter this e.g. 'start'
searchAttributes(mart = ensembl, pattern = "start")

## filter the attributes to give only entries containing 'entrez' or 'hgnc'
searchAttributes(mart = ensembl, 'entrez|hgnc')</pre>
```

listDatasets

List or search the datasets available in the selected BioMart database

Description

Lists or search the datasets available in the selected BioMart database

Usage

```
listDatasets(mart, verbose = FALSE)
searchDatasets(mart, pattern = ".*")
```

Arguments

mart object of class Mart created with the useMart function

verbose Give detailed output of what the method is doing, for debugging purposes

pattern Character vector defining the regular expression (regex) to be used for the search.

If left blank the default is to use ".*" which will match everything and return the

same as listDatasets().

Author(s)

Steffen Durinck, Mike Smith

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Examples

```
## list the available Ensembl marts and use Ensembl Genes
listEnsembl()
ensembl <- useEnsembl(biomart = "ENSEMBL_MART_ENSEMBL")

## list the available datasets in this Mart
listDatasets(mart = ensembl)

## the list of Ensembl datasets grows ever larger (101 as of Ensembl 93)

## we can search for a term of interest to reduce the length e.g. 'sapiens'
searchDatasets(mart = ensembl, pattern = "sapiens")

## search for any dataset containing the word Rat or rat
searchDatasets(mart = ensembl, pattern = "(R|r)at")</pre>
```

listEnsembl

lists the available BioMart databases hosted by Ensembl

Description

This function returns a list of BioMart databases hosted by Ensembl. To establish a connection use the useEnsembl() function.

Usage

```
listEnsembl(
  mart = NULL,
  version = NULL,
  GRCh = NULL,
  mirror = NULL,
  verbose = FALSE
)

listEnsemblGenomes(includeHosts = FALSE, host = NULL)
```

Arguments

mart	mart object created with the useEnsembl function. This is optional, as you usually use listMarts() to see which marts there are to connect to.
version	Ensembl version to connect to when wanting to connect to an archived Ensembl version
GRCh	GRCh version to connect to if not the current GRCh38, currently this can only be 37
mirror	Specify an Ensembl mirror to connect to. The valid options here are 'www', 'useast', 'asia'. If no mirror is specified the primary site at www.ensembl.org will be used.

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verbose Give detailed output of what the method is doing, for debugging purposes

includeHosts If this option is set to TRUE a more detailed output is produced, including the

URL used to access the corresponding mart.

host Host to connect to. Use this argument to specify and archive site for listEnsemblGenomes()

to work with.

Author(s)

Steffen Durinck, Mike L. Smith

Examples

```
listEnsembl()
## list the default Ensembl Genomes marts
listEnsemblGenomes()
## list only the marts available in the Ensmbl Plants 56 archive
listEnsemblGenomes(host = "https://eg56-plants.ensembl.org/")
```

listEnsemblArchives

Lists the available archived versions of Ensembl

Description

Returns a table containing the available archived versions of Ensembl, along with the dates they were created and the URL used to access them.

Usage

listEnsemblArchives()

Author(s)

Mike Smith

Examples

listEnsemblArchives()

listFilters 17

listFilters

List or search the filters available in the selected dataset

Description

Filters are what we use as inputs for a biomaRt query. For example, if we want to retrieve all EntrezGene identifiers on chromosome X, chromosome will be the filter, with corresponding value X.

Usage

```
listFilters(mart, what = c("name", "description"))
searchFilters(mart, pattern = ".*")
```

Arguments

mart object of class Mart created using the useMart() function

what character vector indicating what information to display about the available fil-

ters. Valid values are name, description, options, fullDescription, filters,

type, operation, filters8, filters9.

pattern Character vector defining the regular expression (regex) to be used for the search.

If left blank the default is to use ".. *"" which will match everything.

Author(s)

Steffen Durinck, Mike Smith

```
## list the available Ensembl marts and use Ensembl Genes
listEnsembl()
ensembl <- useEnsembl(
    biomart = "ENSEMBL_MART_ENSEMBL",
    dataset = "hsapiens_gene_ensembl"
)

## list the available datasets in this Mart
listFilters(mart = ensembl)

## the list of filters is long and not easy to read
## we can search for a term of interest to reduce this e.g. 'gene'
searchFilters(mart = ensembl, pattern = "gene")

## search the available filters to find entries containing 'entrez' or 'hgnc'
searchFilters(mart = ensembl, 'entrez|hgnc')</pre>
```

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listMarts

lists the available BioMart databases

Description

This function returns a list of BioMart databases to which biomaRt can connect. By default the Ensembl BioMart databases are displayed. To establish a connection use the useMart() function.

Usage

```
listMarts(
  mart = NULL,
  host = "https://www.ensembl.org",
  path = "/biomart/martservice",
  port,
  includeHosts = FALSE,
  archive = FALSE,
  http_config = list(),
  verbose = FALSE
)
```

Arguments

mart	mart object created with the useMart() function. This is optional, as you usually use listMarts() to see which marts there are to connect to.
host	Host to connect to. Defaults to www.ensembl.org
path	path to martservice that should be pasted behind the host to get to web service URL
port	port to use in HTTP communication
includeHosts	boolean to indicate if function should return host of the BioMart databases
archive	Boolean to indicate if you want to access archived versions of BioMart database. Note that this argument is now defunct and setting this value to TRUE will produce an error. A better alternative is to specify the url of the archived BioMart you want to access. For Ensembl you can view the list of archives using listEnsemblArchives()
http_config	Some hosts require specific HTTP settings to be used when connecting. This argument takes the output of httr::config() and will be used when connecting to host. Can be ignored if you experience no problems accessing host.
verbose	Give detailed output of what the method is doing, for debugging purposes.

Details

If you receive an error message saying 'Unexpected format to the list of available marts', this is often because there is a problem with the BioMart server you are trying to connect to, and something other than the list of available marts is being returned - often some like a 'down for maintenance' page. If you browse to the provided URL and find a page that starts with '<MartRegistry>' this is the correct listing and you should report the issue on the Bioconductor support site: https://support.bioconductor.org

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Author(s)

Steffen Durinck, Mike Smith

Examples

listMarts()

NP2009code

Display the analysis code from the 2009 Nature protocols paper

Description

This function opens an editor displaying the analysis code of the Nature Protocols 2009 paper

Usage

```
NP2009code()
```

Details

The edit() function uses getOption("editor") to select the editor. Use, for instance, options(editor="emacs") to set another editor.

Author(s)

Steffen Durinck, Wolfgang Huber

See Also

edit()

Examples

NP2009code()

20 searchFilterOptions

searchFilterOptions

List or search the options available for a specified filter.

Description

Some filters have a predefined list of values that can be used to search them. These functions give access to this list of options for a named filter, so you can check in the case where your biomaRt query is not finding anything.

Usage

```
searchFilterOptions(mart, filter, pattern = ".*")
listFilterOptions(mart, filter)
```

Arguments

mart object of class Mart created using the useMart(), or useEnsembl() functions

filter The name of the filter whose options should be listed or searched. You can list available filters via listFilters()

pattern Character vector defining the regular expression (regex) to be used for the search.

If left blank the default is to use ".*" which will match everything.

Author(s)

Mike Smith

See Also

```
listFilters()
```

```
## Use the Ensembl human genes dataset
ensembl <- useEnsembl(
   biomart = "ENSEMBL_MART_ENSEMBL",
   dataset = "hsapiens_gene_ensembl"
)

## we can search for the name of a filter we're interested in e.g. 'phenotype'
## we need to use the name of the filter in the next function
searchFilters(ensembl, pattern = "phenotype")

## list all the options available to the 'phenotype_source' filter
listFilterOptions(mart = ensembl, filter = "phenotype_source")

## search the 'phenotype_description' filter for the term 'crohn'
searchFilterOptions(</pre>
```

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```
mart = ensembl,
filter = "phenotype_description",
pattern = "crohn"
)
```

select-methods

Retrieve information from the BioMart databases

Description

select, columns and keys are used together to extract data from a Mart object. These functions work much the same as the classic biomaRt functions such as getBM() etc. and are provide here to make this easier for people who are comfortable using these methods from other Annotation packages. Examples of other objects in other packages where you can use these methods include (but are not limited to): ChipDb, OrgDb GODb, InparanoidDb and ReactomeDb.

Usage

```
## S4 method for signature 'Mart'
keys(x, keytype, ...)
## S4 method for signature 'Mart'
keytypes(x)
## S4 method for signature 'Mart'
columns(x)
## S4 method for signature 'Mart'
select(x, keys, columns, keytype, ...)
```

Arguments

Χ

the Mart object. The dataset of the Mart object must already be specified for all of these methods.

keytype

the keytype that matches the keys used. For the select methods, this is used to indicate the kind of ID being used with the keys argument. For the keys method this is used to indicate which kind of keys are desired from keys

. . .

other arguments. These include:

pattern: the pattern to match (used by keys)

column: the column to search on. This is used by keys and is for when the thing you want to pattern match is different from the keytype, or when you want to simply want to get keys that have a value for the thing specified by the column argument.

fuzzy: TRUE or FALSE value. Use fuzzy matching? (this is used with pattern by the keys method)

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keys the keys to select records for from the database. Keys for some keytypes can be

extracted by using the keys method.

columns the columns or kinds of things that can be retrieved from the database. As with

keys, all possible columns are returned by using the columns method.

Details

columns shows which kinds of data can be returned from the Mart object.

keytypes allows the user to discover which keytypes can be passed in to select or keys as the keytype argument.

keys returns keys from the Mart of the type specified by it's keytype argument.

select is meant to be used with these other methods and has arguments that take the kinds of values that these other methods return. select will retrieve the results as a data frame based on parameters for selected keys and columns and keytype arguments.

Value

keys, columns and keytypes each return a character vector or possible values. select returns a data frame.

Author(s)

Marc Carlson

```
## 1st create a Mart object and specify the dataset
mart <- useEnsembl(</pre>
 biomart = "ENSEMBL_MART_ENSEMBL",
 dataset = "hsapiens_gene_ensembl"
)
## you can list the keytypes
keytypes(mart)
## you can list the columns
columns(mart)
## And you can extract keys when this is supported for your keytype of interest
k <- keys(mart, keytype="chromosome_name")</pre>
head(k)
## You can even do some pattern matching on the keys
k <- keys(mart, keytype="chromosome_name", pattern="LRG")</pre>
head(k)
## Finally you can use select to extract records for things that you are
## interested in.
affy <- c("202763_at", "209310_s_at", "207500_at")
select(mart, keys=affy, columns=c('affy_hg_u133_plus_2','entrezgene_id'),
 keytype='affy_hg_u133_plus_2')
```

setEnsembISSL 23

setEnsemb1SSL

Save system specific SSL settings for contacting Ensembl

Description

On some systems specific SSL settings have to be applied to allow https connections to the Ensembl servers. This function allows these to be saved in the biomaRt cache, so they will be retrieved each time they are needed. biomaRt will try to determine them automatically, but this function can be used to set them manually if required.

Usage

```
setEnsemblSSL(settings)
```

Arguments

settings

A named list. Each entry should be a valid curl option, as found in curl::curl_options().

Author(s)

Mike Smith

Examples

```
## Not run:
ssl_settings <- list(
    "ssl_cipher_list" = "DEFAULT@SECLEVEL=1",
    "ssl_verifypeer" = FALSE
)
setEnsemblSSL(ssl_settings)
## End(Not run)</pre>
```

show, Mart-method

Class Mart

Description

Represents a Mart class, containing connections to different BioMarts

Usage

```
## S4 method for signature 'Mart'
show(object)
```

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Arguments

object An object of class Mart

Methods

show Print summary of the object

Author(s)

Steffen Durinck

useDataset

Select a dataset to use and updates Mart object

Description

This function selects a dataset and updates the Mart object

Usage

```
useDataset(dataset, mart, verbose = FALSE)
```

Arguments

dataset Dataset you want to use. List of possible datasets can be retrieved using the

function listDatasets()

mart Mart object created with the useMart() function

verbose Give detailed output of what the method is doing, for debugging

Author(s)

Steffen Durinck

```
mart <- useMart("ENSEMBL_MART_ENSEMBL")
mart <- useDataset("hsapiens_gene_ensembl", mart = mart)</pre>
```

useEnsembl 25

useEnsembl	Connects to the selected BioMart database and dataset hosted by Ensembl
	semoi

Description

A first step in using the biomaRt package is to select a BioMart database and dataset to use. The useEnsembl() function enables one to connect to a specified BioMart database and dataset hosted by Ensembl without having to specify the Ensembl URL. To know which BioMart databases are available see the listEnsembl() and listEnsemblGenomes() functions. To know which datasets are available within a BioMart database, first select the BioMart database using useEnsembl() and then use the listDatasets() function on the selected Mart object.

Usage

```
useEnsembl(
  biomart,
  dataset,
  host,
  version = NULL,
  GRCh = NULL,
  mirror = NULL,
  verbose = FALSE
)
useEnsemblGenomes(biomart, dataset, host = NULL)
```

Arguments

biomart	BioMart database name you want to connect to. Possible database names can be retrieved with the function listEnsembl()
dataset	Dataset you want to use. To see the different datasets available within a biomaRt you can e.g. do: mart = useEnsembl('genes'), followed by listDatasets(mart).
host	Host to connect to. Only needs to be specified if different from www.ensembl.org. For useEnsemblGenomes() this argument can be used to specify an archive site.
version	Ensembl version to connect to when wanting to connect to an archived Ensembl version
GRCh	GRCh version to connect to if not the current GRCh38, currently this can only be 37
mirror	Specify an Ensembl mirror to connect to. The valid options here are 'www', 'useast', 'asia'. If no mirror is specified the primary site at www.ensembl.org will be used. Mirrors are not available for the Ensembl Genomes databases.
verbose	Give detailed output of what the method is doing while in use, for debugging

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Details

The mirror argument can be considered as a "preferred choice" when connecting to Ensembl. If the argument is provided then connectivity to that mirror will be tested. If it responds positively then the requested mirror will be used. If the response is a failure each of the remaining mirrors will be selected at random and tested until a working server is found. Once identified that Ensembl server will be associated with the returned Mart object and will be used for all queries.

Author(s)

Steffen Durinck & Mike Smith

Examples

```
mart <- useEnsembl("ENSEMBL_MART_ENSEMBL")

## using the US East mirror
us_mart <- useEnsembl(biomart = "ENSEMBL_MART_ENSEMBL", mirror = "useast")

## using the Arabidopsis thaliana genes dataset in Ensembl Plants
plants_mart <- useEnsemblGenomes(
   biomart = "plants_mart",
   dataset = "athaliana_eg_gene"
)

## using the Cucumis melo genes dataset in the Ensembl Plants 56 archive
plants_mart <- useEnsemblGenomes(
   biomart = "plants_mart",
   dataset = "cmelo_eg_gene",
   host = "https://feb2023-plants.ensembl.org/"
)</pre>
```

useMart

Connects to the selected BioMart database and dataset

Description

A first step in using the biomaRt package is to select a BioMart database and dataset to use. The useMart function enables one to connect to a specified BioMart database and dataset within this database. To know which BioMart databases are available see the <code>listMarts()</code> function. To know which datasets are available within a BioMart database, first select the BioMart database using <code>useMart()</code> and then use the <code>listDatasets()</code> function on the selected BioMart, see <code>listDatasets()</code> function.

Usage

```
useMart(
  biomart,
```

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```
dataset,
host = "https://www.ensembl.org",
path = "/biomart/martservice",
port,
archive = FALSE,
version,
verbose = FALSE
)
```

Arguments

biomart	BioMart database name you want to connect to. Possible database names can be retrieved with the function listMarts()
dataset	Dataset you want to use. To see the different datasets available within a biomaRt you can e.g. do: mart = useMart(), followed by listDatasets().
host	Host to connect to. Defaults to www.ensembl.org
path	Path that should be pasted after to host to get access to the web service URL
port	port to connect to, will be pasted between host and path
archive	Boolean to indicate if you want to access archived versions of BioMart databases. Note that this argument is now deprecated and will be removed in the future. A better alternative is to leave archive = FALSE and to specify the url of the archived BioMart you want to access. For Ensembl you can view the list of archives using listEnsemblArchives()
version	Use version name instead of biomart name to specify which BioMart you want to use

Give detailed output of what the method is doing while in use, for debugging

Author(s)

verbose

Steffen Durinck, Mike L. Smith

```
mart <- useMart("ENSEMBL_MART_ENSEMBL")
mart <- useMart(
  biomart = "ENSEMBL_MART_ENSEMBL",
  dataset = "hsapiens_gene_ensembl"
)</pre>
```

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