

# Package ‘UCSC.utils’

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**Title** Low-level utilities to retrieve data from the UCSC Genome Browser

**Description** A set of low-level utilities to retrieve data from the UCSC Genome Browser. Most functions in the package access the data via the UCSC REST API but some of them query the UCSC MySQL server directly. Note that the primary purpose of the package is to support higher-level functionalities implemented in downstream packages like GenomeInfoDb or txdbmaker.

**biocViews** Infrastructure, GenomeAssembly, Annotation, GenomeAnnotation, DataImport

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**BugReports** <https://github.com/Bioconductor/UCSC.utils/issues>

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fetch\_UCSC\_track\_data *Fetch UCSC track data*

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### Description

Fetch the track data for a given UCSC genome/track.

### Usage

```
fetch_UCSC_track_data(genome, primary_table, api.url=UCSC.api.url())
```

### Arguments

genome	A single string specifying the name of a UCSC genome e.g. "hs1", "danRer11", or "wuhCor1". See <a href="#">?list_UCSC_genomes</a> for how to get the list of valid UCSC genome names.
primary_table	A single string specifying the name of the primary table associated with the track from which to fetch the data. See <a href="#">?list_UCSC_tracks</a> for how to get the list of tracks and associated primary tables for a given UCSC genome.
api.url	The URL of the UCSC API. By default, the URL returned by <code>UCSC.api.url()</code> is used. Note that what <code>UCSC.api.url()</code> returns is controlled by a global option. See <a href="#">?UCSC.api.url</a> for more information.

### Value

A data frame.

### See Also

- [list\\_UCSC\\_genomes](#) to get the list of UCSC genomes.
- [list\\_UCSC\\_tracks](#) to get the list of tracks and associated primary tables for a given UCSC genome.
- [UCSC\\_dbselect](#) for a more efficient and more flexible way to retrieve data directly from the UCSC MariaDB server.

**Examples**

```

gorGor6_gap_data <- fetch_UCSC_track_data("gorGor6", "gap")
head(gorGor6_gap_data)

## --- Comparison with UCSC_dbselect() ---

gorGor6_gap_data2 <- UCSC_dbselect("gorGor6", "gap")

## Easy sanity checks.
stopifnot(
  identical(dim(gorGor6_gap_data), dim(gorGor6_gap_data2)),
  identical(colnames(gorGor6_gap_data), colnames(gorGor6_gap_data2))
)

## But the two data frames are not identical:
identical(gorGor6_gap_data, gorGor6_gap_data2) # FALSE!

## However, they have the same content. Ony reason they're not
## identical is because their rows are not in the same order.
## Let's reorder the rows in the two data frames by genomic location,
## that is, first by chromosome, then by chromStart, and finally
## by chromEnd:
sort_rows <- function(gap_data) {
  oo <- order(gap_data$chrom, gap_data$chromStart, gap_data$chromEnd)
  gap_data <- gap_data[oo, ]
  rownames(gap_data) <- NULL
  gap_data
}

gorGor6_gap_data <- sort_rows(gorGor6_gap_data)
gorGor6_gap_data2 <- sort_rows(gorGor6_gap_data2)

## Now the two data frames are identical:
stopifnot(identical(gorGor6_gap_data, gorGor6_gap_data2))

```

---

get\_UCSC\_chrom\_sizes *List UCSC chromosome sizes*

---

**Description**

Get the chromosome sizes of a given UCSC genome.

**Usage**

```
get_UCSC_chrom_sizes(genome, api.url=UCSC.api.url(), recache=FALSE)
```

**Arguments**

genome	A single string specifying the name of a UCSC genome e.g. "hs1", "mm39", or "sacCer3". See <a href="#">?list_UCSC_genomes</a> for how to get the list of valid UCSC genome names.
api.url	The URL of the UCSC API. By default, the URL returned by <code>UCSC.api.url()</code> is used. Note that what <code>UCSC.api.url()</code> returns is controlled by a global option. See <a href="#">?UCSC.api.url</a> for more information.
recache	<code>get_UCSC_chrom_sizes()</code> uses a cache mechanism so the information retrieved for a given genome only gets downloaded once during the current R session (note that the caching is done in memory so cached information does NOT persist across sessions). Setting <code>recache</code> to TRUE forces a new download (and re-caching) of the chromosome sizes for the specified genome.

**Value**

A named numeric vector. The names on the vector are the UCSC chromosomes/sequences. The vector values are the corresponding lengths.

Note that the vector is not sorted in any particular order. In particular there's not guarantee that the chromosomes will precede the scaffolds.

**See Also**

- [list\\_UCSC\\_genomes](#) to get the list of UCSC genomes.
- The [Seqinfo](#) constructor function in the **GenomeInfoDb** package for an alternate (higher level) way of retrieving the chromosome information of a given NCBI assembly or UCSC genome.
- [list\\_UCSC\\_tracks](#) to get the list of tracks and associated primary tables for a given UCSC genome.
- [UCSC.api.url](#) for how to use an alternative UCSC API URL by default.

**Examples**

```
get_UCSC_chrom_sizes("ce2")
get_UCSC_chrom_sizes("hg38")
```

---

list_UCSC_genomes	<i>List UCSC genomes</i>
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**Description**

Get the list of UCSC genomes.

**Usage**

```
list_UCSC_genomes(organism=NA, api.url=UCSC.api.url(), recache=FALSE)

get_organism_for_UCSC_genome(genome, api.url=UCSC.api.url(), recache=FALSE)
```

**Arguments**

organism	By default all UCSC genomes are returned (in a data frame). When organism is specified, <code>list_UCSC_genomes()</code> will only return the rows associated with the specified organism. <code>organism</code> must be supplied as a single string that will be used to perform a search (with <code>grep()</code> ) on the <code>organism</code> and <code>common_name</code> columns of the data frame to return. The search is case-insensitive.
api.url	The URL of the UCSC API. By default, the URL returned by <code>UCSC.api.url()</code> is used. Note that what <code>UCSC.api.url()</code> returns is controlled by a global option. See <a href="#">?UCSC.api.url</a> for more information.
recache	<code>list_UCSC_genomes()</code> uses a cache mechanism so the list of genomes only gets downloaded once during the current R session (note that the caching is done in memory so cached information does NOT persist across sessions). Setting <code>recache</code> to TRUE forces a new download (and recaching) of the list of genomes. In the case of <code>get_organism_for_UCSC_genome()</code> , the supplied <code>recache</code> value is just passed down to the internal call to <code>list_UCSC_genomes()</code> .
genome	A character vector of valid UCSC genomes e.g. "hg38", "mm39", or "sacCer3".

**Value**

For `list_UCSC_genomes`: A data frame with 1 row per genome and 5 columns: `organism`, `genome`, `common_name`, `tax_id`, `description`.

For `get_organism_for_UCSC_genome`: A named character of the same length as the input containing the scientific names of the organisms associated with the supplied UCSC genomes.

**See Also**

- <https://genome.ucsc.edu/FAQ/FAQreleases.html> for the online HTML page that lists all UCSC genome releases, including archived ones.
- [registered\\_UCSC\\_genomes](#) in the **GenomeInfoDb** package for a similar function that returns only UCSC genomes registered in the **GenomeInfoDb** package.
- [get\\_UCSC\\_chrom\\_sizes](#) to get the chromosome sizes of a given UCSC genome.
- [list\\_UCSC\\_tracks](#) to get the list of tracks and associated primary tables for a given UCSC genome.
- [UCSC.api.url](#) for how to use an alternative UCSC API URL by default.

**Examples**

```
list_UCSC_genomes("human")

list_UCSC_genomes("pacos")

get_organism_for_UCSC_genome(c("ce11", "xenTro10", "mpxvRivers"))
```

---

list_UCSC_tracks	<i>List UCSC tracks and associated primary tables</i>
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---

### Description

Get the list of tracks and associated primary tables for a given UCSC genome.

### Usage

```
list_UCSC_tracks(genome, group=NULL,
                 api.url=UCSC.api.url(), recache=FALSE)
```

### Arguments

genome	A single string specifying the name of a UCSC genome e.g. "hg38", "mm39", or "sacCer3". See <a href="#">?list_UCSC_genomes</a> for how to get the list of valid UCSC genome names.
group	NULL or a single string specifying the group of tracks to return. By default, all tracks are returned. Passing group=NA is accepted and will return only rows associated with tracks that don't belong to any group.
api.url	The URL of the UCSC API. By default, the URL returned by <code>UCSC.api.url()</code> is used. Note that what <code>UCSC.api.url()</code> returns is controlled by a global option. See <a href="#">?UCSC.api.url</a> for more information.
recache	<code>list_UCSC_tracks()</code> uses a cache mechanism so the information retrieved for a given genome only gets downloaded once during the current R session (note that the caching is done in memory so cached information does NOT persist across sessions). Setting <code>recache</code> to TRUE forces a new download (and re-caching) of the list of tracks for the specified genome.

### Value

A data frame with 1 row per track and 5 columns: track, primary\_table, type, group, composite\_track.

Note that columns group and composite\_track can contain NAs.

### See Also

- [list\\_UCSC\\_genomes](#) to get the list of UCSC genomes.
- [fetch\\_UCSC\\_track\\_data](#) to fetch the track data for a given UCSC genome/track.
- [get\\_UCSC\\_chrom\\_sizes](#) to get the chromosome sizes of a given UCSC genome.
- [UCSC.api.url](#) for how to use an alternative UCSC API URL by default.

## Examples

```
## List all tracks for ce2 genome:
list_UCSC_tracks("ce2")

## List tracks in the "rna" group only:
list_UCSC_tracks("ce2", group="rna")

## Note that some tracks don't belong to any group:
list_UCSC_tracks("hg38", group=NA)
```

---

UCSC.api.url

*Get or set the default UCSC API URL*

---

## Description

Convenience helper for getting or setting global option `UCSC.api.url`.

## Usage

```
UCSC.api.url(api.url=NULL)
```

## Arguments

`api.url` A single string containing the URL to the alternative UCSC API to use by default.

For convenience, `api.url` can also be one of the following aliases:

- "primary": alias for "https://api.genome.ucsc.edu" (primary URL, US West Coast);
- "": same as "primary";
- "europe": alias for "https://genome-euro.ucsc.edu/cgi-bin/hubApi" (Europe mirror);
- "asia": alias for "https://genome-asia.ucsc.edu/cgi-bin/hubApi" (Asia mirror).

## Details

Various functions in the **UCSC.utils** package query the UCSC REST API. This is the case for example for `list_UCSC_genomes`, `get_UCSC_chrom_sizes`, `list_UCSC_tracks` and more.

Global option `UCSC.api.url` controls the UCSC API URL that these functions use by default. The option is set to "https://api.genome.ucsc.edu" (primary URL, US West Coast) at package startup.

`UCSC.api.url()` and `UCSC.api.url(some_url)` are provided as convenient ways of doing `getOption("UCSC.api.url")` and `options(UCSC.api.url=some_url)`, respectively.

**Value**

When called with no argument, `UCSC.api.url()` returns `getOption("UCSC.api.url")`.

When passed an URL, `UCSC.api.url(some_url)` returns the *previous* URL, that is, the UCSC API URL that was previously used by default. Note that the previous URL is returned invisibly.

**See Also**

- <https://genome.ucsc.edu/goldenPath/help/api.html#Mirrors> a list of alternative UCSC API URLs.
- [list\\_UCSC\\_genomes](#) to get the list of UCSC genomes.

**Examples**

```
UCSC.api.url() # current default value of the UCSC API URL
get_UCSC_chrom_sizes("ce11", recache=TRUE)

## Temporarily use the mirror in Europe:
previous_url <- UCSC.api.url("europe")
UCSC.api.url() # new default value of the UCSC API URL

get_UCSC_chrom_sizes("ce11", recache=TRUE)

## Restore previous default value:
UCSC.api.url(previous_url)
UCSC.api.url()
```

---

UCSC\_dbselect

*Conveniently retrieve data from the UCSC MariaDB server*


---

**Description**

A convenience utility to retrieve data from the UCSC MariaDB server via simple SQL SELECT statements.

Requires the **RMariaDB** package!

**Usage**

```
UCSC_dbselect(dbname, from, columns=NULL, where=NULL, MoreSQL=NULL,
              host="genome-mysql.soe.ucsc.edu", port=3306,
              try.blob.as.list=FALSE)
```

**Arguments**

`dbname` A single string specifying the name of the database to connect to. This is usually the name of a valid UCSC genome e.g. "hg38", "mm39", or "sacCer3". See [?list\\_UCSC\\_genomes](#) for how to get the list of valid UCSC genome names.

from	A single string specifying the <i>input data</i> , that is, the FROM clause of the SELECT statement. This is typically the name of an SQL table, or a join-clause, or a sub-query.
columns	NULL (the default), or a character vector specifying the columns to return. By default all columns are returned.
where	NULL (the default), or a single string specifying the WHERE clause of the SELECT statement.
MoreSQL	NULL (the default), or a character vector containing additional SQL clauses e.g. GROUP BY, ORDER BY, and/or LIMIT clauses.
host	A single string specifying the name of the server's host
port	The TCP/IP port to use to connect to the server.
try.blob.as.list	By default, any column of type blob returned by <b>RMariaDB</b> is converted to a character vector. If <code>try.blob.as.list</code> is set to TRUE, <code>UCSC_dbselect()</code> first tries to convert any blob column to a list of integers, and falls back to the character vector representation if that didn't work.

### Details

`UCSC_dbselect` is an alternative to [fetch\\_UCSC\\_track\\_data](#) that is more efficient and gives the user more control on what data to retrieve exactly from the server.

However, the downside is that `UCSC_dbselect` does not work with all tracks! This is because, not all track data are stored in a database: some tracks are actually file-based (e.g. bigBed tracks). See below for an example.

### Value

A data frame.

### See Also

- [list\\_UCSC\\_genomes](#) to get the list of UCSC genomes.
- [list\\_UCSC\\_tracks](#) to get the list of tracks and associated primary tables for a given UCSC genome.
- [fetch\\_UCSC\\_track\\_data](#) to fetch the track data for a given UCSC genome/track.
- <https://genome.ucsc.edu/goldenpath/help/mysql.html> for more information about downloading data from UCSC MariaDB databases.

### Examples

```
### Retrieve full "ncbiGene" table ("NCBI Genes" track) for eboVir3:
df <- UCSC_dbselect("eboVir3", "ncbiGene")
df
```

```
### The exonStarts and exonEnds columns are character vectors:
sapply(df, class)
df$exonStarts
```

```
df$exonEnds

### Use 'try.blob.as.list=TRUE' to have them returned as lists of
### integers:
df2 <- UCSC_dbselect("eboVir3", "ncbiGene", try.blob.as.list=TRUE)
sapply(df2, class)
df2$exonStarts
df2$exonEnds

### Retrieve subset of "sgpGene" table ("SGP Genes" track) for hg38:
UCSC_dbselect("hg38", "sgpGene", where="chrom='chrM'")

### Retrieve subset of "gap" table for gorGor6:
columns <- c("chrom", "chromStart", "chromEnd", "type")
where <- "chrom='chrX' AND type='contig'"
UCSC_dbselect("gorGor6", "gap", columns=columns, where=where)

### With a LEFT JOIN and a LIMIT clause:
from <- paste("ncbiRefSeq LEFT JOIN ncbiRefSeqLink",
              "ON ncbiRefSeq.name=ncbiRefSeqLink.id")
res <- UCSC_dbselect("sacCer3", from, MoreSQL="LIMIT 5")
dim(res)

## WARNING: UCSC_dbselect() does not work with all tracks!
## For example it does not work with bigBed tracks:
subset(list_UCSC_tracks("hg38", group="genes"), grepl("bigBed 9", type))
## Not run:
UCSC_dbselect("hg38", "crisprAllTargets") # table doesn't exist!

## End(Not run)
```

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