

Package ‘DeepBlueR’

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Title DeepBlueR

Type Package

Description Accessing the DeepBlue Epigenetics Data Server through R.

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Author Felipe Albrecht, Markus List

Maintainer Felipe Al-

brecht <felipe.albrecht@mpi-inf.mpg.de>, Markus List <markus.list@wzw.tum.de>, Quirin Manz <quirin.manz@t

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Imports GenomicRanges, data.table, stringr, diffr, dplyr, methods,
rjson, utils, R.utils, foreach, withr, rtracklayer,
GenomeInfoDb, settings, filehash

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DeepBlueCommand-class *DeepBlueCommand class*

Description

An S4 class returned when calling a DeepBlue-R function. It holds information about the original call, the query / request status, previous commands, the user_key, and results in GRanges format once a request is downloaded.

Arguments

call	language
status	character
query_id	character
previous_commands	list
user_key	character
result	GRanges

Value

class for managin DeepBlue commands

deepblue_aggregate *aggregate*

Description

Summarize the data_id content using the regions specified in ranges_id as boundaries. Use the fields @AGG.MIN, @AGG.MAX, @AGG.SUM, @AGG.MEDIAN, @AGG.MEAN, @AGG.VAR, @AGG.SD, @AGG.COUNT in 'get_regions' command 'format' parameter to retrieve the computed values minimum, maximum, median, mean, variance, standard deviation and number of regions, respectively.

Usage

```
deepblue_aggregate(data_id = NULL, ranges_id = NULL, column = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

data_id	- A string (id of the query with the data)
ranges_id	- A string (id of the query with the regions range)
column	- A string (name of the column that will be used in the aggregation)
user_key	- A string (users token key)

Value

regions - A string (query id of this aggregation operation)

See Also

Other Operating on the data regions: [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(
    annotation_name="CpG Islands",
    genome="hg19", chromosome="chr1")
data_id = deepblue_select_experiments(
    experiment_name="E002-H3K9ac.narrowPeak.bed")
deepblue_aggregate(
    data_id = data_id,
    ranges_id=annotation_id,
    column = "SCORE")
```

```
deepblue_batch_export_results
    batch_export_results
```

Description

Write results from DeepBlue to disk as they become available

Usage

```
deepblue_batch_export_results(requests, target.directory = NULL,
    suffix = "result", prefix = "DeepBlue", sleep.time = 1,
    bed.format = TRUE, user_key = deepblue_options("user_key"))
```

Arguments

requests	A list of request objects
target.directory	Where the results should be saved
suffix	File names suffix
prefix	File names prefix
sleep.time	How long this function will wait after the requests verification

bed.format whether to store the results as BED files or tab delimited.
 user_key A string used to authenticate the user

Value

A list containing the requests IDs data

Examples

```
data_id = deepblue_select_experiments(
  experiment_name="E002-H3K9ac.narrowPeak.bed", chromosome="chr1")
request_id = deepblue_get_regions(query_id =data_id,
  output_format = "CHROMOSOME,START,END")
request_data = deepblue_batch_export_results(list(request_id))
```

deepblue_binning *binning*

Description

Bin results according to counts.

Usage

```
deepblue_binning(query_data_id = NULL, column = NULL, bins = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

query_data_id - A string (query data that will made by the binning.)
 column - A string (name of the column that will be used in the aggregation)
 bins - A int (number of of bins)
 user_key - A string (users token key)

Value

request_id - A string (Request ID - Use it to retrieve the result with info() and get_request_data())

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
experiment_id = deepblue_select_experiments(  
    experiment_name="S00XDKH1.ERX712765.H3K27ac.bwa.GRCh38.20150527.bed")  
deepblue_binning (query_data_id=experiment_id,  
    column="SIGNAL_VALUE",  
    bins=40)
```

deepblue_cache_status *Report on the cache size and status*

Description

Report on the cache size and status

Usage

```
deepblue_cache_status()
```

Value

cache size in byte

Examples

```
deepblue_cache_status()
```

deepblue_cancel_request
cancel_request

Description

Stop, cancel, and remove request data. The request processed data is remove if its processing was finished.

Usage

```
deepblue_cancel_request(id = NULL, user_key = deepblue_options("user_key"))
```

Arguments

id - A string (Request ID to be canceled, stopped or removed.)
user_key - A string (users token key)

Value

id - A string (ID of the canceled request)

See Also

Other Commands for all types of data: [deepblue_info](#), [deepblue_is_biosource](#), [deepblue_list_in_use](#), [deepblue_name_to_id](#), [deepblue_search](#)

Examples

```
deepblue_cancel_request(id = "r12345")
```

deepblue_chromosomes *chromosomes*

Description

List the chromosomes of a given Genome.

Usage

```
deepblue_chromosomes(genome = NULL, user_key = deepblue_options("user_key"))
```

Arguments

genome - A string (the target genome)
user_key - A string (users token key)

Value

chromosomes - A array (A list containing all chromosomes, with their names and sizes)

See Also

Other Inserting and listing genomes: [deepblue_list_genomes](#), [deepblue_list_similar_genomes](#)

Examples

```
deepblue_chromosomes(genome = "g1")
```

deepblue_clear_cache *Clear cache*

Description

Clear cache

Usage

```
deepblue_clear_cache()
```

Value

TRUE if successful

Examples

```
deepblue_clear_cache()
```

deepblue_collection_experiments_count
 collection_experiments_count

Description

Count the number of experiments that match the selection criteria in each term of the selected controlled_vocabulary. The selection can be achieved through specifying a list of BioSources, experimental Techniques, Epigenetic Marks, Samples or Projects.

Usage

```
deepblue_collection_experiments_count(controlled_vocabulary = NULL,
  genome = NULL, type = NULL, epigenetic_mark = NULL, biosource = NULL,
  sample = NULL, technique = NULL, project = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

controlled_vocabulary - A string (controlled vocabulary name)

genome - A string or a vector of string (the target genome)

type - A string or a vector of string (type of the experiment: peaks or signal)

epigenetic_mark - A string or a vector of string (name(s) of selected epigenetic mark(s))

biosource - A string or a vector of string (name(s) of selected biosource(s))

sample - A string or a vector of string (id(s) of selected sample(s))
technique - A string or a vector of string (name(s) of selected technique(s))
project - A string or a vector of string (name(s) of selected projects)
user_key - A string (users token key)

Value

terms - A array (controlled_vocabulary terms with count)

See Also

Other Inserting and listing experiments: [deepblue_faceting_experiments](#), [deepblue_list_experiments](#), [deepblue_list_recent_experiments](#), [deepblue_list_similar_experiments](#), [deepblue_preview_experiment](#)

Examples

```
deepblue_collection_experiments_count(  
  controlled_vocabulary="epigenetic_marks",  
  genome = "hg19", type = "peaks",  
  biosource = "blood")
```

deepblue_commands *commands*

Description

List all available DeepBlue commands.

Usage

```
deepblue_commands()
```

Value

commands - A struct (command descriptions)

See Also

Other Checking DeepBlue status: [deepblue_echo](#)

Examples

```
deepblue_commands()
```

```
deepblue_count_gene_ontology_terms
    count_gene_ontology_terms
```

Description

Summarize the controlled_vocabulary fields, from experiments that match the selection criteria. It is similar to the 'collection_experiments_count' command, but this command return the summarization for all controlled_vocabulary terms.

Usage

```
deepblue_count_gene_ontology_terms(genes = NULL, go_terms = NULL,
    chromosome = NULL, start = NULL, end = NULL, gene_model = NULL,
    user_key = deepblue_options("user_key"))
```

Arguments

genes	- A string or a vector of string (Name(s) or ENSEMBL ID (ENSGXXXXXXXXXXXXX.X) of the gene(s).)
go_terms	- A string or a vector of string (gene ontology terms - ID or label)
chromosome	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
gene_model	- A string (the gene model)
user_key	- A string (users token key)

Value

faceting - A struct (Map with the mandatory fields of the experiments metadata, where each contains a list of terms that appears.)

See Also

Other Gene models and genes identifiers: [deepblue_list_gene_models](#), [deepblue_list_genes](#), [deepblue_select_genes](#)

Examples

```
gene_names = c('CCR1', 'CD164', 'CD1D', 'CD2', 'CD34', 'CD3G', 'CD44')
deepblue_count_gene_ontology_terms (genes = gene_names, gene_model = "gencode v23")
```

deepblue_count_regions
count_regions

Description

Return the number of genomic regions present in the query.

Usage

```
deepblue_count_regions(query_id = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

query_id - A string (Query ID)
user_key - A string (users token key)

Value

request_id - A string (Request ID - Use it to retrieve the result with info() and get_request_data())

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
data_id = deepblue_select_experiments(  
  experiment_name="E002-H3K9ac.narrowPeak.bed")  
deepblue_count_regions(query_id = data_id)
```

deepblue_coverage	<i>coverage</i>
-------------------	-----------------

Description

Send a request to count the number of regions in the result of the given query.

Usage

```
deepblue_coverage(query_id = NULL, genome = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
genome	- A string (Genome where the coverage will be calculated to)
user_key	- A string (users token key)

Value

request_id - A string (Request ID - Use it to retrieve the result with info() and get_request_data())

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
data_id = deepblue_select_experiments(  
  experiment_name="E002-H3K9ac.narrowPeak.bed")  
deepblue_coverage(query_id = data_id, genome="hg19")
```

deepblue_delete_request_from_cache

Delete a specific request from the cache

Description

Delete a specific request from the cache

Usage

```
deepblue_delete_request_from_cache(request_id)
```

Arguments

request_id the request to delete from the cache

Value

TRUE if the request was successfully deleted, FALSE otherwise

Examples

```
deepblue_delete_request_from_cache("non-existing-request-id")  
# returns FALSE
```

deepblue_diff

diff

Description

A utility command that creates a diff view of info for two DeepBlue ids

Usage

```
deepblue_diff(id1, id2, user_key = deepblue_options("user_key"))
```

Arguments

id1 - A DeepBlue id
id2 - Another DeepBlue id
user_key - A string (users token key)

Value

None

See Also

Other Utilities for information processing: [deepblue_select_column](#)

Examples

```
deepblue_diff(  
  id1 = "e16918",  
  id2 = "e16919")
```

```
deepblue_distinct_column_values  
  distinct_column_values
```

Description

Obtain the distinct values of the field.

Usage

```
deepblue_distinct_column_values(query_id = NULL, field = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
field	- A string (field that is filtered by)
user_key	- A string (users token key)

Value

id - A string (id of filtered query)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
css_experiment <- deepblue_select_experiments ( "wgEncodeBroadHmK562HMM")  
distinct_names_request <- deepblue_distinct_column_values (css_experiment, "NAME")
```

deepblue_download_request_data
deepblue_download_request_data

Description

Returns the requested data as the expected type object. Expects two input parameters; Request information and user key. It depends on outputs from several functions, namely; `deepblue_get_request_data`, `convert_to_df`, and `convert_to_grange`.

Usage

```
deepblue_download_request_data(request_id,  
    user_key = deepblue_options("user_key"),  
    force_download = deepblue_options("force_download"),  
    do_not_cache = deepblue_options("do_not_cache"))
```

Arguments

<code>request_id</code>	- Id of the request that will be downloaded
<code>user_key</code>	A string
<code>force_download</code>	forces DeepBlueR to download the request overwriting any results that might already be in the cache
<code>do_not_cache</code>	whether to use local caching of requests

Value

`grange_regions` Final output in GRanges format or as data frame

Examples

```
data_id = deepblue_select_experiments(  
    experiment_name="E002-H3K9ac.narrowPeak.bed", chromosome="chr1")  
request_id = deepblue_get_regions(query_id =data_id,  
    output_format = "CHROMOSOME,START,END")  
request_data = deepblue_download_request_data(request_id)
```

deepblue_download_request_data,DeepBlueCommand-method
deepblue_download_request_data

Description

Returns the requested data as the expected type object. Expects two input parameters; Request information and user key. It depends on outputs from several functions, namely; `deepblue_get_request_data`, `convert_to_df`, and `convert_to_grange`.

Usage

```
## S4 method for signature 'DeepBlueCommand'  
deepblue_download_request_data(request_id)
```

Arguments

`request_id` DeepBlueCommand object

Value

`grange_regions` Final output in GRanges format

`deepblue_echo` *echo*

Description

Greet the user with the DeepBlue version.

Usage

```
deepblue_echo(user_key = deepblue_options("user_key"))
```

Arguments

`user_key` - A string (users token key)

Value

`message` - A string (echo message including version)

See Also

Other Checking DeepBlue status: [deepblue_commands](#)

Examples

```
deepblue_echo(user_key = "anonymous_key")
```

```
deepblue_enrich_regions_fast  
    enrich_regions_fast
```

Description

Enrich the regions based on regions bitmap signature comparison.

Usage

```
deepblue_enrich_regions_fast(query_id = NULL, genome = NULL,  
    epigenetic_mark = NULL, biosource = NULL, sample = NULL,  
    technique = NULL, project = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
genome	- A string or a vector of string (the target genome)
epigenetic_mark	- A string or a vector of string (name(s) of selected epigenetic mark(s))
biosource	- A string or a vector of string (name(s) of selected biosource(s))
sample	- A string or a vector of string (id(s) of selected sample(s))
technique	- A string or a vector of string (name(s) of selected technique(s))
project	- A string or a vector of string (name(s) of selected projects)
user_key	- A string (users token key)

Value

request_id - A string (Request ID - Use it to retrieve the result with info() and get_request_data()).
The result is a list containing the datasets that overlap with the query_id regions.)

See Also

Other Enrich the genome regions: [deepblue_enrich_regions_go_terms](#), [deepblue_enrich_regions_overlap](#)

deepblue_enrich_regions_go_terms
enrich_regions_go_terms

Description

Enrich the regions based on Gene Ontology terms.

Usage

```
deepblue_enrich_regions_go_terms(query_id = NULL, gene_model = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

query_id - A string (Query ID)
gene_model - A string (the gene model)
user_key - A string (users token key)

Value

request_id - A string (Request ID - Use it to retrieve the result with `info()` and `get_request_data()`.
The result is a list containing the GO terms that overlap with the query_id regions.)

See Also

Other Enrich the genome regions: [deepblue_enrich_regions_fast](#), [deepblue_enrich_regions_overlap](#)

Examples

```
data_id = deepblue_select_experiments(  
    experiment_name="E002-H3K9ac.narrowPeak.bed")  
  
filtered_id = deepblue_filter_regions(query_id = data_id,  
    field = "VALUE",  
    operation = ">",  
    value = "100",  
    type = "number",  
    user_key = "anonymous_key")  
  
deepblue_enrich_regions_go_terms(query_id = filtered_id,  
    gene_model = "gencode v23")
```

```
deepblue_enrich_regions_overlap
    enrich_regions_overlap
```

Description

Enrich the regions based on regions overlap analysis.

Usage

```
deepblue_enrich_regions_overlap(query_id = NULL, background_query_id = NULL,
    datasets = NULL, genome = NULL, user_key = deepblue_options("user_key"))
```

Arguments

query_id - A string (Query ID)
background_query_id - A string (query_id containing the regions that will be used as the background data.)
datasets - A struct (a map where each key is an identifier and the value is a list containing experiment names or query_ids (you can use both together).)
genome - A string (the target genome)
user_key - A string (users token key)

Value

request_id - A string (Request ID - Use it to retrieve the result with info() and get_request_data()).
The result is a list containing the datasets that overlap with the query_id regions.)

See Also

Other Enrich the genome regions: [deepblue_enrich_regions_fast](#), [deepblue_enrich_regions_go_terms](#)

Examples

```
query_id = deepblue_select_experiments(
    experiment_name="S00VEQA1.hypo_meth.bs_call.GRCh38.20150707.bed")

filtered_query_id = deepblue_filter_regions(
    query_id = query_id,
    field = "AVG_METHYL_LEVEL",
    operation = "<",
    value = "0.0025",
    type="number")

rg_10kb_tiling = deepblue_tiling_regions(
    size = 1000,
```

```

    genome = "hg19")

# We could have included more Epigenetic Marks here
epigenetic_marks <- c("h3k27ac", "H3K27me3", "H3K4me3")

histones_datasets = c()
for (i in 1:length(epigenetic_marks)) {
  experiments_list <- deepblue_list_experiments(
    epigenetic_mark=epigenetic_marks[[i]],
    type="peaks",
    genome="grch38",
    project="BLUEPRINT Epigenome");

  experiment_names = deepblue_extract_names(experiments_list)
  histones_datasets[[epigenetic_marks[[i]]]] = experiment_names
}

deepblue_enrich_regions_overlap(
  query_id=filtered_query_id,
  background_query=rg_10kb_tiling,
  datasets=histones_datasets,
  genome="grch38")

```

deepblue_export_bed *Export GenomicRanges result as BED file*

Description

Export GenomicRanges result as BED file

Usage

```
deepblue_export_bed(result, target.directory = "./", file.name,
  score.field = NULL)
```

Arguments

result	A result from a DeepBlue request such as a set of genomic regions.
target.directory	The directory to save the file to
file.name	The name of the file without suffix
score.field	Which column of the results should be used to populate the score column of the BED file (optional)

Value

return value of write.table

Examples

```

query_id = deepblue_select_experiments (
  experiment=c("GC_T14_10.CPG_methylation_calls.bs_call.GRCh38.20160531.wig"),
  chromosome="chr1", start=0, end=50000000)
cpg_islands = deepblue_select_annotations(annotation_name="CpG Islands",
  genome="GRCh38", chromosome="chr1", start=0, end=50000000)
overlapped = deepblue_aggregate (data_id=query_id, ranges_id=cpg_islands,
  column="VALUE" )
request_id = deepblue_get_regions(query_id=overlapped,
  output_format=
    "CHROMOSOME,START,END,@AGG.MIN,@AGG.MAX,@AGG.MEAN,@AGG.VAR")
regions = deepblue_download_request_data(request_id=request_id)
temp_dir = tempdir()
deepblue_export_bed(regions, target.directory = temp_dir,
  file.name = "GC_T14_10.CpG_islands")

```

deepblue_export_meta_data

Export meta data as tab delimited file

Description

Export meta data as tab delimited file

Usage

```

deepblue_export_meta_data(ids, target.directory = "./", file.name,
  user_key = deepblue_options("user_key"))

```

Arguments

ids	an id or a list of DeepBlue ids
target.directory	where the meta data should be stored
file.name	name of the file
user_key	DeepBlue user key

Value

return value of write.table

Examples

```

deepblue_export_meta_data(list("e30035", "e30036"),
  file.name = "test_export",
  target.directory = tempdir())

```

deepblue_export_tab *Export a DeepBlue result as ordinary tab delimited file*

Description

Export a DeepBlue result as ordinary tab delimited file

Usage

```
deepblue_export_tab(result, target.directory = "./", file.name)
```

Arguments

result	A result from a DeepBlue request such as a set of genomic regions.
target.directory	The directory to save the file to
file.name	The name of the file without suffix

Value

return value of write.table

Examples

```
query_id = deepblue_select_experiments (
  experiment=c("GC_T14_10.CPG_methylation_calls.bs_call.GRCh38.20160531.wig"),
  chromosome="chr1", start=0, end=50000000)
cpg_islands = deepblue_select_annotations(annotation_name="CpG Islands",
  genome="GRCh38", chromosome="chr1", start=0, end=50000000)
overlapped = deepblue_aggregate (data_id=query_id, ranges_id=cpg_islands,
  column="VALUE" )
request_id = deepblue_get_regions(query_id=overlapped,
  output_format=
    "CHROMOSOME,START,END,@AGG.MIN,@AGG.MAX,@AGG.MEAN,@AGG.VAR")
regions = deepblue_download_request_data(request_id=request_id)
temp_dir = tempdir()
deepblue_export_tab(regions, target.directory = temp_dir,
  file.name = "GC_T14_10.CpG_islands")
```

deepblue_extend	<i>extend</i>
-----------------	---------------

Description

Extend the genomic regions included in the query. It is possible to extend downstream, upstream or in both directions.

Usage

```
deepblue_extend(query_id = NULL, length = NULL, direction = NULL,
                use_strand = NULL, user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
length	- A int (The new region length)
direction	- A string (The direction that the region will be extended: 'BACKWARD', 'FORWARD', 'BOTH'. (Empty value will be used for both direction.)
use_strand	- A boolean (Use the region column STRAND to define the region direction)
user_key	- A string (users token key)

Value

id - A string (id of the new query)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_region](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(
  annotation_name="CpG Islands",
  genome="hg19", chromosome="chr1")
deepblue_extend(query_id = annotation_id,
  length = 2000, direction = "BOTH",
  use_strand = TRUE)
```

deepblue_extract_ids *extract_ids*

Description

A utility command that returns a list of IDs extracted from a data frame of ID and names.

Usage

```
deepblue_extract_ids(df = NULL)
```

Arguments

df - A array of IDs and names

Value

ids - A vector containing the extracted IDs)

See Also

Other Utilities for connecting operations: [deepblue_extract_names](#)

Examples

```
deepblue_extract_ids(  
  df = data.frame(id = c("a124", "a1235"),  
    name = c("Annotation 1", "Annotation 2")))
```

deepblue_extract_names
extract_names

Description

A utility command that returns a list of names extracted from a list of ID and names.

Usage

```
deepblue_extract_names(df = NULL)
```

Arguments

df - A array of IDs and Names

Value

names - A vector containing the extracted names

See Also

Other Utilities for connecting operations: [deepblue_extract_ids](#)

Examples

```
deepblue_extract_ids(
  df = data.frame(id = c("a124", "a1235"),
    name = c("Annotation 1", "Annotation 2")))
```

```
deepblue_faceting_experiments
  faceting_experiments
```

Description

Summarize the controlled_vocabulary fields, from experiments that match the selection criteria. It is similar to the 'collection_experiments_count' command, but this command return the summarization for all controlled_vocabulary terms.

Usage

```
deepblue_faceting_experiments(genome = NULL, type = NULL,
  epigenetic_mark = NULL, biosource = NULL, sample = NULL,
  technique = NULL, project = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

genome	- A string or a vector of string (the target genome)
type	- A string or a vector of string (type of the experiment: peaks or signal)
epigenetic_mark	- A string or a vector of string (name(s) of selected epigenetic mark(s))
biosource	- A string or a vector of string (name(s) of selected biosource(s))
sample	- A string or a vector of string (id(s) of selected sample(s))
technique	- A string or a vector of string (name(s) of selected technique(s))
project	- A string or a vector of string (name(s) of selected projects)
user_key	- A string (users token key)

Value

faceting - A struct (Map with the mandatory fields of the experiments metadata, where each contains a list of terms that appears.)

See Also

Other Inserting and listing experiments: [deepblue_collection_experiments_count](#), [deepblue_list_experiments](#), [deepblue_list_recent_experiments](#), [deepblue_list_similar_experiments](#), [deepblue_preview_experiment](#)

Examples

```
deepblue_faceting_experiments(genome = "hg19",
                              type = "peaks",
                              biosource = "blood")
```

```
deepblue_filter_regions
                        filter_regions
```

Description

Filter the genomic regions by their content.

Usage

```
deepblue_filter_regions(query_id = NULL, field = NULL, operation = NULL,
                        value = NULL, type = NULL, user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
field	- A string (field that is filtered by)
operation	- A string (operation used for filtering. For 'string' must be '==' or '!=' and for 'number' must be one of these: ==, !=, >, >=, <, <=)
value	- A string (value the operator is applied to)
type	- A string (type of the value: 'number' or 'string')
user_key	- A string (users token key)

Value

id - A string (id of filtered query)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
deepblue_filter_regions(query_id = "q12345",
  field = "VALUE",
  operation = ">",
  value = "100",
  type = "number",
  user_key = "anonymous_key")
```

deepblue_find_motif *find_motif*

Description

Find genomic regions based on a given motif that appears in the genomic sequence.

Usage

```
deepblue_find_motif(motif = NULL, genome = NULL, chromosomes = NULL,
  start = NULL, end = NULL, overlap = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

motif	- A string (motif (PERL regular expression))
genome	- A string (the target genome)
chromosomes	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
overlap	- A boolean (if the matching should do overlap search)
user_key	- A string (users token key)

Value

id - A string (id of the annotation that contains the positions of the given motif)

See Also

Other Inserting and listing annotations: [deepblue_list_annotations](#)

Examples

```
deepblue_find_motif(motif = "C[GT]+C", chromosomes=c("chr11", "chr12"),
  genome = "hg19", overlap = FALSE)
```

deepblue_flank	<i>flank</i>
----------------	--------------

Description

Create a set of genomic regions that flank the query regions. The original regions are removed from the query. Use the merge command to combine flanking regions with the original query.

Usage

```
deepblue_flank(query_id = NULL, start = NULL, length = NULL,  
use_strand = NULL, user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
start	- A int (Number of base pairs after the end of the region. Use a negative number to denote the number of base pairs before the start of the region.)
length	- A int (The new region length)
use_strand	- A boolean (Use the region column STRAND to define the region direction)
user_key	- A string (users token key)

Value

id - A string (id of the new query)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_region](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(  
  annotation_name="CpG Islands",  
  genome="hg19", chromosome="chr1")  
deepblue_flank(query_id = annotation_id,  
  start = 0, length = 2000,  
  use_strand = TRUE)
```

deepblue_format_object_size

Format byte size as human readable units

Description

Format byte size as human readable units

Usage

```
deepblue_format_object_size(x, units = "b")
```

Arguments

x	size in bytes
units	target unit or 'auto'

Value

formatted size

Source

utils:::format.object_size

deepblue_get_biosource_children

get_biosource_children

Description

A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line. These form a hierarchy in which children of a BioSource term can be fetched with this command. Children terms are more specific terms that are defined in the imported ontologies.

Usage

```
deepblue_get_biosource_children(biosource = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

biosource	- A string (biosource name)
user_key	- A string (users token key)

Value

biosources - A array (related biosources)

See Also

Other Set the relationship between different biosources: [deepblue_get_biosource_parents](#), [deepblue_get_biosource_r](#), [deepblue_get_biosource_synonyms](#)

Examples

```
deepblue_get_biosource_children(biosource = "Blood")
```

```
deepblue_get_biosource_parents  
    get_biosource_parents
```

Description

A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line. These form a hierarchy in which the parent of a BioSource term can be fetched with this command. Parent terms are more generic terms that are defined in the imported ontologies.

Usage

```
deepblue_get_biosource_parents(biosource = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

biosource - A string (biosource name)
user_key - A string (users token key)

Value

biosources - A array (parents biosources)

See Also

Other Set the relationship between different biosources: [deepblue_get_biosource_children](#), [deepblue_get_biosource_related](#), [deepblue_get_biosource_synonyms](#)

Examples

```
deepblue_get_biosource_parents(biosource = "Blood")
```

deepblue_get_biosource_related
get_biosource_related

Description

A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line. These form a hierarchy in which the children of a BioSource term and its synonyms can be fetched with this command. Children terms are more specific terms that are defined in the imported ontologies. Synonyms are different aliases for the same biosource.

Usage

```
deepblue_get_biosource_related(biosource = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

biosource - A string (biosource name)
user_key - A string (users token key)

Value

biosources - A array (related biosources)

See Also

Other Set the relationship between different biosources: [deepblue_get_biosource_children](#), [deepblue_get_biosource_parents](#), [deepblue_get_biosource_synonyms](#)

Examples

```
deepblue_get_biosource_related(biosource = "Blood")
```

deepblue_get_biosource_synonyms
get_biosource_synonyms

Description

Obtain the synonyms of the specified biosource. Synonyms are different aliases for the same biosource. A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line.

Usage

```
deepblue_get_biosource_synonyms(biosource = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

biosource - A string (name of the biosource)
user_key - A string (users token key)

Value

synonyms - A array (synonyms of the biosource)

See Also

Other Set the relationship between different biosources: [deepblue_get_biosource_children](#), [deepblue_get_biosource_parents](#), [deepblue_get_biosource_related](#)

Examples

```
deepblue_get_biosource_synonyms(biosource = "prostate gland")
```

deepblue_get_db	<i>Sets up the DeepBlueR cache and returns a filehash db object</i>
-----------------	---------------------------------------------------------------------

Description

Sets up the DeepBlueR cache and returns a filehash db object

Usage

```
deepblue_get_db()
```

Value

A filehash package database

```
deepblue_get_experiments_by_query
    get_experiments_by_query
```

Description

List the experiments and annotations that have at least one genomic region in the final query result.

Usage

```
deepblue_get_experiments_by_query(query_id = NULL,
    user_key = deepblue_options("user_key"))
```

Arguments

```
query_id      - A string (Query ID)
user_key      - A string (users token key)
```

Value

experiments - A array (List containing experiments names and ids)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
deepblue_get_experiments_by_query(query_id = "q12345")
```

```
deepblue_get_regions    get_regions
```

Description

Trigger the processing of the query's genomic regions. The output is a column based format with columns as defined in the 'output_format' parameter. Use the command 'info' for verifying the processing status. The 'get_request_data' command is used to download the regions using the programmatic interface. Alternatively, results can be download using the URL: http://deepblue.mpi-inf.mpg.de/download?r_id=<request_id>&key=<user_key>.

Usage

```
deepblue_get_regions(query_id = NULL, output_format = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

```
query_id      - A string (Query ID)  
output_format - A string (Output format)  
user_key      - A string (users token key)
```

Value

request_id - A string (Request ID - Use it to retrieve the result with info() and get_request_data())

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
data_id = deepblue_select_experiments(  
  experiment_name="E002-H3K9ac.narrowPeak.bed")  
deepblue_get_regions(query_id =data_id,  
  output_format = "CHROMOSOME,START,END")
```

deepblue_get_request_data

get_request_data

Description

Download the requested data. The output can be (i) a string (get_regions, score_matrix, and count_regions), or (ii) a list of ID and names (get_experiments_by_query), or (iii) a struct (coverage).

Usage

```
deepblue_get_request_data(request_id = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

request_id - A string (ID of the request)
 user_key - A string (users token key)

Value

data - A string or a vector of string (the request data)

See Also

Other Requests status information and results: [deepblue_list_requests](#)

Examples

```
data_id = deepblue_select_experiments(
  experiment_name="E002-H3K9ac.narrowPeak.bed",
  chromosome="chr1")
request_id = deepblue_get_regions(
  query_id =data_id,
  output_format = "CHROMOSOME,START,END")
deepblue_get_request_data(request_id = request_id)
```

deepblue_info	<i>info</i>
---------------	-------------

Description

Information about a DeepBlue data identifier (ID). Any DeepBlue data ID can be queried with this command. For example, it is possible to obtain all available information about an Experiment using its ID, to obtain the actual Request processing status or the information about a Sample. A user can obtain information about him- or herself using the value 'me' in the parameter 'id'. Multiple IDs can be queried in the same operation.

Usage

```
deepblue_info(id = NULL, user_key = deepblue_options("user_key"))
```

Arguments

id - A string or a vector of string (ID or an array of IDs)
 user_key - A string (users token key)

Value

information - A array or a vector of array (List of Maps, where each map contains the info of an object.)

See Also

Other Commands for all types of data: [deepblue_cancel_request](#), [deepblue_is_biosource](#), [deepblue_list_in_use](#), [deepblue_name_to_id](#), [deepblue_search](#)

Examples

```
deepblue_info(id = "e30035")
```

```
deepblue_input_regions  
    input_regions
```

Description

Upload a set of genomic regions that can be accessed through a query ID. An interesting use case for this command is to upload a set of custom regions for intersecting with genomic regions in DeepBlue to specifically select regions of interest.

Usage

```
deepblue_input_regions(genome = NULL, region_set = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

genome	- A string (the target genome)
region_set	- A string (Regions in CHROMOSOME START END format)
user_key	- A string (users token key)

Value

id - A string (query id)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
regions_set = "chr1 28735 29810
chr1 135124 135563
chr1 327790 328229
chr1 437151 438164
chr1 449273 450544
chr1 533219 534114
chr1 544738 546649
chr1 713984 714547
chr1 762416 763445
chr1 788863 789211"
deepblue_input_regions(genome = "hg19",
    region_set = regions_set)
```

deepblue_intersection *intersection*

Description

Select genomic regions that intersect with at least one region of the second query. This command is a simplified version of the 'overlap' command.

Usage

```
deepblue_intersection(query_data_id = NULL, query_filter_id = NULL,
    user_key = deepblue_options("user_key"))
```

Arguments

query_data_id - A string (query data that will be filtered.)
query_filter_id - A string (query containing the regions that the regions of the query_data_id must overlap.)
user_key - A string (users token key)

Value

id - A string (id of the new query)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_region](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(  
    annotation_name="CpG Islands",  
    genome="hg19", chromosome="chr1")  
data_id = deepblue_select_experiments(  
    experiment_name="E002-H3K9ac.narrowPeak.bed")  
deepblue_intersection(query_data_id = annotation_id,  
    query_filter_id = data_id)
```

deepblue_is_biosource *is_biosource*

Description

Verify if the name is an existing and valid DeepBlue BioSource name. A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line.

Usage

```
deepblue_is_biosource(biosource = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

biosource - A string (biosource name)
user_key - A string (users token key)

Value

information - A string or a vector of string (A string containing the biosource name)

See Also

Other Commands for all types of data: [deepblue_cancel_request](#), [deepblue_info](#), [deepblue_list_in_use](#), [deepblue_name_to_id](#), [deepblue_search](#)

Examples

```
deepblue_is_biosource(biosource = "blood")
```

deepblue_liftover	<i>Lift over region results between Genome Assemblies used in DeepBlue</i>
-------------------	----------------------------------------------------------------------------

Description

This is a wrapper function for the liftOver function found in the rtracklayer package. For common genome assemblies available in DeepBlue, this function automatically downloads the necessary chain file and calls liftOver.

Usage

```
deepblue_liftover(regions, source = "hg19", target = "hg38",  
collapse = TRUE)
```

Arguments

regions	The GRanges object to lift over to another assembly
source	The source assembly version, e.g. hg38. If NULL, we try to read the genome version from the GRanges object.
target	The target assembly version, e.g. hg19. Required.
collapse	Whether to return a single GRanges object or a list of GRanges (one per region in the input). The latter is the default behavior of liftOver since multiple assignments are possible.

Value

A GRanges object using the target chromosome positions

Examples

```
data_id = deepblue_select_experiments(  
  experiment_name="E002-H3K9ac.narrowPeak.bed", chromosome="chr1")  
request_id = deepblue_get_regions(query_id =data_id,  
                                output_format = "CHROMOSOME,START,END")  
request_data = deepblue_download_request_data(request_id)  
deepblue_liftover(request_data, source = "hg38", target = "hg19")
```

```
deepblue_list_annotations
    list_annotations
```

Description

List all annotations of genomic regions currently available in DeepBlue.

Usage

```
deepblue_list_annotations(genome = NULL,
    user_key = deepblue_options("user_key"))
```

Arguments

genome - A string or a vector of string (the target genome)
user_key - A string (users token key)

Value

annotations - A array (annotations names and IDs)

See Also

Other Inserting and listing annotations: [deepblue_find_motif](#)

Examples

```
deepblue_list_annotations(genome = "hg19")
```

```
deepblue_list_biosources
    list_biosources
```

Description

List BioSources included in DeepBlue. A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line. It is possible to filter the BioSources by their extra_metadata fields content. These fields vary depending on the original data source.

Usage

```
deepblue_list_biosources(extra_metadata = NULL,
    user_key = deepblue_options("user_key"))
```

Arguments

- extra_metadata - A struct (Metadata that must be matched)
- user_key - A string (users token key)

Value

biosources - A array (biosources names and IDS)

See Also

Other Inserting and listing biosources: [deepblue_list_similar_biosources](#)

Examples

```
deepblue_list_biosources(extra_metadata = list(ontology_id = "UBERON:0002485"))
```

deepblue_list_cached_requests
List cached requests

Description

List cached requests

Usage

```
deepblue_list_cached_requests()
```

Value

list of request ids that are cached

Examples

```
deepblue_list_cached_requests()
```

```
deepblue_list_column_types
    list_column_types
```

Description

Lists the ColumnTypes included in DeepBlue.

Usage

```
deepblue_list_column_types(user_key = deepblue_options("user_key"))
```

Arguments

user_key - A string (users token key)

Value

column_types - A array (column types names and IDS)

Examples

```
deepblue_list_column_types()
```

```
deepblue_list_epigenetic_marks
    list_epigenetic_marks
```

Description

List Epigenetic Marks included in DeepBlue. This includes histone marks, DNA methylation, DNA sensitivity, etc. It is possible to filter the Epigenetic Marks by their extra_metadata field content.

Usage

```
deepblue_list_epigenetic_marks(extra_metadata = NULL,
    user_key = deepblue_options("user_key"))
```

Arguments

extra_metadata - A struct (Metadata that must be matched)

user_key - A string (users token key)

Value

epigenetic_marks - A array (epigenetic mark names and IDS)

See Also

Other Inserting and listing epigenetic marks: [deepblue_list_similar_epigenetic_marks](#)

Examples

```
deepblue_list_epigenetic_marks()
```

```
deepblue_list_experiments
      list_experiments
```

Description

List the DeepBlue Experiments that matches the search criteria defined by this command parameters.

Usage

```
deepblue_list_experiments(genome = NULL, type = NULL,
  epigenetic_mark = NULL, biosource = NULL, sample = NULL,
  technique = NULL, project = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

genome	- A string or a vector of string (the target genome)
type	- A string or a vector of string (type of the experiment: peaks or signal)
epigenetic_mark	- A string or a vector of string (name(s) of selected epigenetic mark(s))
biosource	- A string or a vector of string (name(s) of selected biosource(s))
sample	- A string or a vector of string (id(s) of selected sample(s))
technique	- A string or a vector of string (name(s) of selected technique(s))
project	- A string or a vector of string (name(s) of selected projects)
user_key	- A string (users token key)

Value

experiments - A array (experiment names and IDS)

See Also

Other Inserting and listing experiments: [deepblue_collection_experiments_count](#), [deepblue_faceting_experiments](#), [deepblue_list_recent_experiments](#), [deepblue_list_similar_experiments](#), [deepblue_preview_experiment](#)

Examples

```
deepblue_list_experiments(genome = "hg19", type = "peaks",  
  epigenetic_mark = "H3K27ac", biosource = "blood")
```

```
deepblue_list_expressions  
  list_expressions
```

Description

List the Expression currently available in DeepBlue. A expression is a set of data with an identifier and an expression value.

Usage

```
deepblue_list_expressions(expression_type = NULL, sample_id = NULL,  
  replica = NULL, project = NULL, user_key = deepblue_options("user_key"))
```

Arguments

expression_type	- A string (expression type (supported: 'gene'))
sample_id	- A string or a vector of string (sample ID(s))
replica	- A int or a vector of int (replica(s))
project	- A string or a vector of string (project(s) name)
user_key	- A string (users token key)

Value

expressions - A array (expressions names and IDS)

See Also

Other Expression data: [deepblue_select_expressions](#)

Examples

```
deepblue_list_expressions(expression_type='gene')
```

deepblue_list_genes *list_genes*

Description

List the Genes currently available in DeepBlue.

Usage

```
deepblue_list_genes(genes = NULL, go_terms = NULL, chromosome = NULL,  
  start = NULL, end = NULL, gene_model = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

genes	- A string or a vector of string (Name(s) or ENSEMBL ID (ENSGXXXXXXXXXXXXX.X) of the gene(s).)
go_terms	- A string or a vector of string (gene ontology terms - ID or label)
chromosome	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
gene_model	- A string (the gene model)
user_key	- A string (users token key)

Value

genes - A array (genes names and its content)

See Also

Other Gene models and genes identifiers: [deepblue_count_gene_ontology_terms](#), [deepblue_list_gene_models](#), [deepblue_select_genes](#)

Examples

```
deepblue_list_genes(  
  chromosome="chr20",  
  start=10000000,  
  end=21696620,  
  gene_model='Gencode v22')
```

deepblue_list_gene_models
list_gene_models

Description

List all the Gene Models currently available in DeepBlue. A gene model is a set of genes usually imported from GENCODE. For example Gencode v22.

Usage

```
deepblue_list_gene_models(user_key = deepblue_options("user_key"))
```

Arguments

user_key - A string (users token key)

Value

gene_models - A array (gene models names and IDS)

See Also

Other Gene models and genes identifiers: [deepblue_count_gene_ontology_terms](#), [deepblue_list_genes](#), [deepblue_select_genes](#)

Examples

```
deepblue_list_gene_models()
```

deepblue_list_genomes *list_genomes*

Description

List Genomes assemblies that are registered in DeepBlue.

Usage

```
deepblue_list_genomes(user_key = deepblue_options("user_key"))
```

Arguments

user_key - A string (users token key)

Value

genomes - A array (genome names)

See Also

Other Inserting and listing genomes: [deepblue_chromosomes](#), [deepblue_list_similar_genomes](#)

Examples

```
deepblue_list_genomes()
```

deepblue_list_in_use *list_in_use*

Description

List all terms used by the Experiments mandatory metadata that have at least one Experiment or Annotation using them.

Usage

```
deepblue_list_in_use(controlled_vocabulary = NULL,  
                    user_key = deepblue_options("user_key"))
```

Arguments

controlled_vocabulary
 - A string (controlled vocabulary name)

user_key - A string (users token key)

Value

terms - A array (controlled_vocabulary terms with count)

See Also

Other Commands for all types of data: [deepblue_cancel_request](#), [deepblue_info](#), [deepblue_is_biosource](#), [deepblue_name_to_id](#), [deepblue_search](#)

Examples

```
deepblue_list_in_use(controlled_vocabulary = "biosources")
```

```
deepblue_list_projects
    list_projects
```

Description

List Projects included in DeepBlue.

Usage

```
deepblue_list_projects(user_key = deepblue_options("user_key"))
```

Arguments

`user_key` - A string (users token key)

Value

`projects` - A array (project names)

See Also

Other Inserting and listing projects: [deepblue_list_similar_projects](#)

Examples

```
deepblue_list_projects()
```

```
deepblue_list_recent_experiments
    list_recent_experiments
```

Description

List the latest Experiments included in DeepBlue that match criteria defined in the parameters. The returned experiments are sorted by insertion date.

Usage

```
deepblue_list_recent_experiments(days = NULL, genome = NULL,
    epigenetic_mark = NULL, sample = NULL, technique = NULL,
    project = NULL, user_key = deepblue_options("user_key"))
```

Arguments

days	- A double (maximum days ago the experiments were added)
genome	- A string or a vector of string (the target genome)
epigenetic_mark	- A string or a vector of string (name(s) of selected epigenetic mark(s))
sample	- A string or a vector of string (id(s) of selected sample(s))
technique	- A string or a vector of string (name(s) of selected technique(es))
project	- A string or a vector of string (name(s) of selected projects)
user_key	- A string (users token key)

Value

experiments - A array (names of recent experiments)

See Also

Other Inserting and listing experiments: [deepblue_collection_experiments_count](#), [deepblue_faceting_experiments](#), [deepblue_list_experiments](#), [deepblue_list_similar_experiments](#), [deepblue_preview_experiment](#)

Examples

```
deepblue_list_recent_experiments(days = 2, genome = "hg19")
```

```
deepblue_list_requests
      list_requests
```

Description

List the Requests made by the user. It is possible to obtain only the requests of a given state.

Usage

```
deepblue_list_requests(request_state = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

request_state	- A string (Name of the state to get requests for. The valid states are: new, running, done, and failed.)
user_key	- A string (users token key)

Value

data_state - A array (Request-IDs and their state)

See Also

Other Requests status information and results: [deepblue_get_request_data](#)

Examples

```
deepblue_list_requests(request_state = 'running')
```

deepblue_list_samples *list_samples*

Description

List Samples included in DeepBlue. It is possible to filter by the BioSource and by extra_metadata fields content.

Usage

```
deepblue_list_samples(biosource = NULL, extra_metadata = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

biosource - A string or a vector of string (name(s) of selected biosource(s))
extra_metadata - A struct (Metadata that must be matched)
user_key - A string (users token key)

Value

samples - A array (samples id with their content)

Examples

```
deepblue_list_samples(biosource = "Blood")
```

deepblue_list_similar_biosources
list_similar_biosources

Description

List all BioSources that have a similar name compared to the provided name. A BioSource refers to a term describing the origin of a given sample, such as a tissue or cell line. The similarity is calculated using the Levenshtein method.

Usage

```
deepblue_list_similar_biosources(name = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

name - A string (biosource name)
user_key - A string (users token key)

Value

biosource - A string (biosource name)

See Also

Other Inserting and listing biosources: [deepblue_list_biosources](#)

Examples

```
deepblue_list_similar_biosources(name = "blood")
```

deepblue_list_similar_epigenetic_marks
list_similar_epigenetic_marks

Description

List all Epigenetic Marks that have a similar name compared to the provided name. The similarity is calculated using the Levenshtein method.

Usage

```
deepblue_list_similar_epigenetic_marks(name = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

name - A string (epigenetic mark name)
user_key - A string (users token key)

Value

epigenetic_marks - A array (similar epigenetic mark names)

See Also

Other Inserting and listing epigenetic marks: [deepblue_list_epigenetic_marks](#)

Examples

```
deepblue_list_similar_epigenetic_marks(name = "H3k27ac")
```

deepblue_list_similar_experiments
list_similar_experiments

Description

List all Experiments that have a similar name compared to the provided name. The similarity is calculated using the Levenshtein method.

Usage

```
deepblue_list_similar_experiments(name = NULL, genome = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

name - A string (experiment name)
genome - A string or a vector of string (the target genome)
user_key - A string (users token key)

Value

experiments - A array (similar experiment names)

See Also

Other Inserting and listing experiments: [deepblue_collection_experiments_count](#), [deepblue_faceting_experiments](#), [deepblue_list_experiments](#), [deepblue_list_recent_experiments](#), [deepblue_preview_experiment](#)

Examples

```
deepblue_list_similar_experiments(name = "blood", genome = "hg19")
```

```
deepblue_list_similar_genomes  
    list_similar_genomes
```

Description

Lists all Genomes that have a similar name compared to the provided name. The similarity is calculated using the Levenshtein method.

Usage

```
deepblue_list_similar_genomes(name = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

name	- A string (genome name)
user_key	- A string (users token key)

Value

genomes - A array (similar genome names)

See Also

Other Inserting and listing genomes: [deepblue_chromosomes](#), [deepblue_list_genomes](#)

Examples

```
deepblue_list_similar_genomes(name = "grc")
```

deepblue_list_similar_projects
list_similar_projects

Description

List Projects that have a similar name compared to the provided name. The similarity is calculated using the Levenshtein method.

Usage

```
deepblue_list_similar_projects(name = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

name - A string (project name)
user_key - A string (users token key)

Value

projects - A array (similar project names)

See Also

Other Inserting and listing projects: [deepblue_list_projects](#)

Examples

```
deepblue_list_similar_projects(name = "BLUEPRINT")
```

deepblue_list_similar_techniques
list_similar_techniques

Description

List Techniques that have a similar name compared to the provided name. The similarity is calculated using the Levenshtein method.

Usage

```
deepblue_list_similar_techniques(name = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

name - A string (technique name)
user_key - A string (users token key)

Value

techniques - A array (similar techniques)

See Also

Other Inserting and listing techniques: [deepblue_list_techniques](#)

Examples

```
deepblue_list_similar_techniques(name = "chip seq")
```

deepblue_list_techniques
list_techniques

Description

List the Techniques included in DeepBlue.

Usage

```
deepblue_list_techniques(user_key = deepblue_options("user_key"))
```

Arguments

user_key - A string (users token key)

Value

techniques - A array (techniques)

See Also

Other Inserting and listing techniques: [deepblue_list_similar_techniques](#)

Examples

```
deepblue_list_techniques()
```

deepblue_merge_queries
merge_queries

Description

Merge regions from two queries in a new query.

Usage

```
deepblue_merge_queries(query_a_id = NULL, query_b_id = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

query_a_id - A string (id of the first query)
query_b_id - A string or a vector of string (id of the second query (or use an array to include multiple queries))
user_key - A string (users token key)

Value

id - A string (new query id)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(  
  annotation_name="CpG Islands",  
  genome="hg19", chromosome="chr1")  
data_id = deepblue_select_experiments(  
  experiment_name="E002-H3K9ac.narrowPeak.bed")  
deepblue_merge_queries(  
  query_a_id = annotation_id,  
  query_b_id = data_id)
```

deepblue_meta_data_to_table

Convert XML structured meta data to table format

Description

Convert XML structured meta data to table format

Usage

```
deepblue_meta_data_to_table(ids, user_key = deepblue_options("user_key"))
```

Arguments

ids	an id or a list of ids
user_key	a DeepBlue user key (optional for public data)

Value

a data frame with meta data

Examples

```
#works for sample ids
deepblue_meta_data_to_table(list("s2694", "s2695"))

#or experiment ids
deepblue_meta_data_to_table(list("e30035", "e30036"))
```

deepblue_name_to_id *name_to_id*

Description

Obtain the data ID(s) from the informed data name(s).

Usage

```
deepblue_name_to_id(name = NULL, collection = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

name	- A string or a vector of string (ID or an array of IDs)
collection	- A string (Collection where the data name is in)
user_key	- A string (users token key)

Value

information - A array or a vector of array (List of IDs.)

See Also

Other Commands for all types of data: [deepblue_cancel_request](#), [deepblue_info](#), [deepblue_is_biosource](#), [deepblue_list_in_use](#), [deepblue_search](#)

Examples

```
deepblue_name_to_id("E002-H3K9ac.narrowPeak.bed", "experiments")
deepblue_name_to_id("prostate duct", "biosources")
deepblue_name_to_id("DNA Methylation", "Epigenetic_marks")
```

deepblue_options	<i>options</i>
------------------	----------------

Description

options manager from the settings package

Usage

```
deepblue_options(..., .__defaults = FALSE, .__reset = FALSE)
```

Arguments

...	list of new options
.__defaults	disallowed option
.__reset	disallowed option

Value

default options

deepblue_overlap *overlap*

Description

Select genomic regions that overlap or not overlap with with the specified number of regions of the second query. Important: This command is still experimental and changes may occur.

Usage

```
deepblue_overlap(query_data_id = NULL, query_filter_id = NULL,
  overlap = NULL, amount = NULL, amount_type = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

query_data_id - A string (query data that will be filtered.)

query_filter_id
- A string (query containing the regions that the regions of the query_data_id must overlap.)

overlap - A boolean (True if must overlap, or false if must not overlap.)

amount - A int (Amount of regions that must overlap. Use the parameter 'amount_type' ('bp' or '%') to specify the unit. For example, use the value '10' with the amount_type '%' to specify that 10% of the bases in both regions must overlap, or use '10' with the amount_type 'bp' to specify that at least 10 bases must or must not overlap.)

amount_type - A string (Type of the amount: 'bp' for base pairs and '%' for percentage.)

user_key - A string (users token key)

Value

id - A string (id of the new query)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(  
    annotation_name="CpG Islands",  
    genome="hg19", chromosome="chr1")  
experiment_id = deepblue_select_experiments(  
    experiment_name="S00XDKH1.ERX712765.H3K27ac.bwa.GRCh38.20150527.bed")  
deepblue_overlap(query_data_id = experiment_id, query_filter_id = annotation_id,  
    overlap = TRUE, amount=10, amount_type="%")
```

deepblue_preview_experiment

preview_experiment

Description

List the DeepBlue Experiments that matches the search criteria defined by this command parameters.

Usage

```
deepblue_preview_experiment(experiment_name = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

experiment_name - A string (name(s) of selected experiment(s))

user_key - A string (users token key)

Value

experiment - A string (experiment's regions)

See Also

Other Inserting and listing experiments: [deepblue_collection_experiments_count](#), [deepblue_faceting_experiments](#), [deepblue_list_experiments](#), [deepblue_list_recent_experiments](#), [deepblue_list_similar_experiments](#)

Examples

```
deepblue_preview_experiment('S00JJRH1.ERX683143.H3K4me3.bwa.GRCh38.20150527.bed')
```

deepblue_query_cache *query_cache*

Description

Cache a query result in DeepBlue memory. This command is useful when the same query ID is used multiple times in different requests. The command is an advice for DeepBlue to cache the query result and there is no guarantee that this query data access will be faster.

Usage

```
deepblue_query_cache(query_id = NULL, cache = NULL,  
user_key = deepblue_options("user_key"))
```

Arguments

query_id - A string (Query ID)
cache - A boolean (set or unset this query caching)
user_key - A string (users token key)

Value

information - A string (New query ID.)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
annotation_id = deepblue_select_annotations(  
  annotation_name="CpG Islands",  
  genome="hg19", chromosome="chr1")  
data_id = deepblue_select_experiments(  
  experiment_name="E002-H3K9ac.narrowPeak.bed")  
merged_regions = deepblue_merge_queries(  
  query_a_id = annotation_id,  
  query_b_id = data_id)  
deepblue_query_cache(  
  query_id = merged_regions, cache = TRUE)
```

deepblue_query_experiment_type
query_experiment_type

Description

Filter the query ID for regions associated with experiments of a given type. For example, it is possible to select only peaks using this command with the 'peaks' parameter.

Usage

```
deepblue_query_experiment_type(query_id = NULL, type = NULL,  
    user_key = deepblue_options("user_key"))
```

Arguments

query_id	- A string (Query ID)
type	- A string (experiment type (peaks or signal))
user_key	- A string (users token key)

Value

information - A string (New query ID.)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
h3k27ac_regions = deepblue_select_regions(  
    genome = 'GRCh38',  
    epigenetic_mark = 'H3k27ac',  
    project = 'BLUEPRINT Epigenome',  
    chromosome = 'chr1')  
deepblue_query_experiment_type(  
    query_id = h3k27ac_regions,  
    type = "peaks")
```

deepblue_reset_options

Reset DeepBlueR options

Description

Reset DeepBlueR options

Usage

```
deepblue_reset_options(new_options = NULL)
```

Arguments

new_options list of new options that should be used. default options if NULL

Value

new (default) options

Examples

```
deepblue_reset_options()
```

deepblue_score_matrix *score_matrix*

Description

Build a matrix containing the aggregation result of the the experiments data by the aggregation boundaries.

Usage

```
deepblue_score_matrix(experiments_columns = NULL,
  aggregation_function = NULL, aggregation_regions_id = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

experiments_columns

- A struct (map with experiments names and columns to be processed. Example : 'wgEncodeBroadHistoneDnd41H3k27acSig.wig': 'VALUE', 'wgEncodeBroadHistoneCd20ro01794H3k27acSig.wig': 'VALUE')

aggregation_function
 - A string (aggregation function name: min, max, sum, mean, var, sd, median, count, boolean)

aggregation_regions_id
 - A string (query ID of the regions that will be used as the aggregation boundaries)

user_key
 - A string (users token key)

Value

score_matrix - A string (the score matrix containing the summarized data)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
tiling_regions = deepblue_tiling_regions(
  size=100000, genome="mm10", chromosome="chr1")
deepblue_score_matrix(
  experiments_columns =
    list(ENCF721EKA="VALUE", ENCF781VH="VALUE"),
  aggregation_function = "mean",
  aggregation_regions_id = tiling_regions)
```

deepblue_search	<i>search</i>
-----------------	---------------

Description

Search all data of all types for the given keyword. A minus (-) character in front of a keyword searches for data without the given keyword. The search can be restricted to the following data types are: Annotations, Biosources, Column_types, Epigenetic_marks, Experiments, Genomes, Gene_models, Gene_expressions, Genes, Gene_ontology, Projects, Samples, Techniques, Tilings.

Usage

```
deepblue_search(keyword = NULL, type = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

keyword	- A string (keyword to search by)
type	- A string or a vector of string (type of data to search for - Annotations, Biosources, Column_types, Epigenetic_marks, Experiments, Genomes, Gene_models, Gene_expressions, Genes, Gene_ontology, Projects, Samples, Techniques, Tilings)
user_key	- A string (users token key)

Value

results - A array (search results as [id, name, type])

See Also

Other Commands for all types of data: [deepblue_cancel_request](#), [deepblue_info](#), [deepblue_is_biosource](#), [deepblue_list_in_use](#), [deepblue_name_to_id](#)

Examples

```
deepblue_search(keyword = "DNA Methylation BLUEPRINT",
  type = "experiments")
```

```
deepblue_select_annotations
  select_annotations
```

Description

Select regions from the Annotations that match the selection criteria.

Usage

```
deepblue_select_annotations(annotation_name = NULL, genome = NULL,
  chromosome = NULL, start = NULL, end = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

annotation_name	- A string or a vector of string (name(s) of selected annotation(s))
genome	- A string (the target genome)
chromosome	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
user_key	- A string (users token key)

Value

id - A string (query id)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_experiments](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
deepblue_select_annotations(  
  annotation_name = "Cpg Islands",  
  genome = "hg19",  
  chromosome = "chr1",  
  start = 0,  
  end = 2000000)
```

```
deepblue_select_column  
      select column
```

Description

A utility command that creates a list of experiments in which a specific column is selected. Such a list is needed as input for `deepblue_score_matrix`.

Usage

```
deepblue_select_column(experiments, column,  
  user_key = deepblue_options("user_key"))
```

Arguments

`experiments` - A data frame with experiments obtained from `deepblue_list_experiments`
`column` - The name of the column that is extracted from each experiment file
`user_key` - A string (users token key)

Value

A list of experiments with the selected column

See Also

[deepblue_score_matrix](#)

[deepblue_list_experiments](#)

Other Utilities for information processing: [deepblue_diff](#)

Examples

```
blueprint_DNA_meth <- deepblue_list_experiments(  
  genome = "GRCh38",  
  epigenetic_mark = "DNA Methylation",  
  technique = "Bisulfite-Seq",  
  project = "BLUEPRINT EPIGENOME")  
  
blueprint_DNA_meth <- blueprint_DNA_meth[grepl("bs_call",  
  deepblue_extract_names(blueprint_DNA_meth)),]  
  
exp_columns <- deepblue_select_column(blueprint_DNA_meth, "VALUE")
```

```
deepblue_select_experiments  
  select_experiments
```

Description

Selects regions from Experiments by the experiments names.

Usage

```
deepblue_select_experiments(experiment_name = NULL, chromosome = NULL,  
  start = NULL, end = NULL, user_key = deepblue_options("user_key"))
```

Arguments

experiment_name	- A string or a vector of string (name(s) of selected experiment(s))
chromosome	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
user_key	- A string (users token key)

Value

id - A string (query id)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_regions](#), [deepblue_tiling_regions](#)

Examples

```
deepblue_select_experiments(
  experiment_name = c("E002-H3K9ac.narrowPeak.bed",
    "E001-H3K4me3.gappedPeak.bed")
)
```

```
deepblue_select_expressions
      select_expressions
```

Description

Select expressions (by their name or ID) as genomic regions from the specified model.

Usage

```
deepblue_select_expressions(expression_type = NULL, sample_ids = NULL,
  replicas = NULL, identifiers = NULL, projects = NULL,
  gene_model = NULL, user_key = deepblue_options("user_key"))
```

Arguments

<code>expression_type</code>	- A string (expression type (supported: 'gene'))
<code>sample_ids</code>	- A string or a vector of string (id(s) of selected sample(s))
<code>replicas</code>	- A int or a vector of int (replica(s))
<code>identifiers</code>	- A string or a vector of string (identifier(s) (for genes: ensembl ID or ENSB name).)
<code>projects</code>	- A string or a vector of string (projects(s))
<code>gene_model</code>	- A string (gene model name)
<code>user_key</code>	- A string (users token key)

Value

`id` - A string (query id)

See Also

Other Expression data: [deepblue_list_expressions](#)

Examples

```
genes_names =
  c('CCR1', 'CD164', 'CD1D', 'CD2', 'CD34', 'CD3G', 'CD44')
deepblue_select_expressions(
  expression_type="gene",
  sample_ids="s10205",
  identifiers = genes_names,
  gene_model = "gencode v23")
```

deepblue_select_genes *select_genes*

Description

Select genes (by their name or ID) as genomic regions from the specified gene model.

Usage

```
deepblue_select_genes(genes = NULL, go_terms = NULL, gene_model = NULL,
  chromosome = NULL, start = NULL, end = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

genes	- A string or a vector of string (Name(s) or ENSEMBL ID (ENSGXXXXXXXXXXXXX.X) of the gene(s).)
go_terms	- A string or a vector of string (gene ontology terms - ID or label)
gene_model	- A string (the gene model)
chromosome	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
user_key	- A string (users token key)

Value

id - A string (query id)

See Also

Other Gene models and genes identifiers: [deepblue_count_gene_ontology_terms](#), [deepblue_list_gene_models](#), [deepblue_list_genes](#)

Examples

```
genes_names =
  c('CCR1', 'CD164', 'CD1D', 'CD2', 'CD34', 'CD3G', 'CD44')
deepblue_select_genes(
  genes = genes_names,
  gene_model = "gencode v23")
```

```
deepblue_select_regions
      select_regions
```

Description

Selects Experiment regions that matches the criteria informed by the operation parameters.

Usage

```
deepblue_select_regions(experiment_name = NULL, genome = NULL,
  epigenetic_mark = NULL, sample_id = NULL, technique = NULL,
  project = NULL, chromosomes = NULL, start = NULL, end = NULL,
  user_key = deepblue_options("user_key"))
```

Arguments

experiment_name	- A string or a vector of string (name(s) of selected experiment(s))
genome	- A string or a vector of string (the target genome)
epigenetic_mark	- A string or a vector of string (name(s) of selected epigenetic mark(s))
sample_id	- A string or a vector of string (id(s) of selected sample(s))
technique	- A string or a vector of string (name(s) of selected technique(es))
project	- A string or a vector of string (name(s) of selected projects)
chromosomes	- A string or a vector of string (chromosome name(s))
start	- A int (minimum start region)
end	- A int (maximum end region)
user_key	- A string (users token key)

Value

id - A string (query id)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_tiling_regions](#)

Examples

```
deepblue_select_regions(  
  genome="hg19",  
  epigenetic_mark = "H3K27ac",  
  project = " BLUEPRINT Epigenome")
```

```
deepblue_tiling_regions  
    tiling_regions
```

Description

Generate tiling regions across the genome chromosomes. The idea is to "bin" genomic regions systematically in order to obtain discrete regions over which one can aggregate. Using the 'score_matrix' command, these bins (tiles) can be compared directly across experiments.

Usage

```
deepblue_tiling_regions(size = NULL, genome = NULL, chromosome = NULL,  
  user_key = deepblue_options("user_key"))
```

Arguments

size	- A int (tiling size)
genome	- A string (the target genome)
chromosome	- A string or a vector of string (chromosome name(s))
user_key	- A string (users token key)

Value

id - A string (query id)

See Also

Other Operating on the data regions: [deepblue_aggregate](#), [deepblue_binning](#), [deepblue_count_regions](#), [deepblue_coverage](#), [deepblue_distinct_column_values](#), [deepblue_extend](#), [deepblue_filter_regions](#), [deepblue_flank](#), [deepblue_get_experiments_by_query](#), [deepblue_get_regions](#), [deepblue_input_regions](#), [deepblue_intersection](#), [deepblue_merge_queries](#), [deepblue_overlap](#), [deepblue_query_cache](#), [deepblue_query_experiment_type](#), [deepblue_score_matrix](#), [deepblue_select_annotations](#), [deepblue_select_experiments](#), [deepblue_select_regions](#)

Examples

```
deepblue_tiling_regions(  
    size = 10000,  
    genome = "hg19",  
    chromosome = "chr1")
```

`deepblue_wait_request` *deepblue_wait_request*

Description

Process the user request. Takes in three parameters; requested regions, sleep time, and user key.

Usage

```
deepblue_wait_request(request_id, sleep_time = 1,  
    user_key = deepblue_options("user_key"))
```

Arguments

<code>request_id</code>	A string with the <code>request_id</code>
<code>sleep_time</code>	An integer with default value 1s
<code>user_key</code>	A string

Value

`request_id` info

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