# Package 'gProfileR'

October 13, 2022

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License GPL (>= 2)			
Description This package has been deprecated and will not be updated.  New users should use the package 'gprofiler2' ( <a href="https://cran.r-project.org/package=gprofiler2">https://cran.r-project.org/package=gprofiler2</a> )  for up-to-date data and improved functionality.  Functional enrichment analysis, gene identifier conversion and mapping homologous genes across related organisms via the 'g:Profiler' toolkit ( <a href="https://biit.cs.ut.ee/gprofiler/">https://biit.cs.ut.ee/gprofiler/</a> ).			
Title Interface to the 'g:Profiler' Toolkit			
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BugReports http://biit.cs.ut.ee/gprofiler/contact.cgi  Depends R (>= 2.10)  Imports RCurl, plyr, utils  Collate 'gProfileR.R'  RoxygenNote 6.1.1  NeedsCompilation no  Repository CRAN  Date/Publication 2019-11-04 16:30:17 UTC  R topics documented:			
r topics documented:			
gconvert get_base_url get_tls_version get_user_agent gorth gprofiler set_base_url			

2 gconvert

```
      set_tls_version
      7

      set_user_agent
      7

      Index
      9
```

gconvert Convert gene IDs.

#### **Description**

Interface to the g:Convert tool. Organism names are constructed by concatenating the first letter of the name and the family name. Example: human - 'hsapiens', mouse - 'mmusculus'.

## Usage

```
gconvert(query, organism = "hsapiens", target = "ENSG",
  region_query = F, numeric_ns = "", mthreshold = Inf,
  filter_na = T, df = T)
```

#### **Arguments**

list of gene IDs. query organism organism name. target target namespace. region\_query interpret query as chromosomal ranges. namespace to use for fully numeric IDs. numeric\_ns mthreshold maximum number of results per initial alias to show. filter\_na logical indicating whether to filter out results without a corresponding target. df logical indicating whether the output will be a data.frame or list.

## Value

The output can be either a list or a data.frame. The list has an entry for every input gene. The data frame is a table closely corresponding to the web interface output.

#### Author(s)

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## References

J. Reimand, M. Kull, H. Peterson, J. Hansen, J. Vilo: g:Profiler - a web-based toolset for functional profiling of gene lists from large-scale experiments (2007) NAR 35 W193-W200

get\_base\_url 3

## Examples

```
## Not run:
gconvert(c("POU5F1", "SOX2", "NANOG"), organism = "hsapiens", target="AFFY_HG_U133_PLUS_2")
## End(Not run)
```

get\_base\_url

Get the base URL.

## Description

Get the base URL.

## Usage

```
get_base_url()
```

get\_tls\_version

Get the TLS version for SSL

## Description

Get the TLS version for SSL

## Usage

```
get_tls_version()
```

get\_user\_agent

Get current user agent string.

## Description

Get the HTTP User-Agent string.

## Usage

```
get_user_agent()
```

4 gorth

gorth	Find orthologs.	

#### **Description**

Interface to the g:Orth tool. Organism names are constructed by concatenating the first letter of the name and the family name. Example: human - 'hsapiens', mouse - 'mmusculus'.

#### Usage

```
gorth(query, source_organism = "hsapiens",
  target_organism = "mmusculus", region_query = F, numeric_ns = "",
  mthreshold = Inf, filter_na = T, df = T)
```

#### **Arguments**

query list of gene IDs to be translated.

source\_organism

name of the source organism.

target\_organism

name of the target organism.

region\_query interpret query as chromosomal ranges.

numeric\_ns namespace to use for fully numeric IDs.

mthreshold maximum number of ortholog names per gene to show.

filter\_na logical indicating whether to filter out results without a corresponding target name.

#### **Details**

df

To alleviate the problem of having many orthologs per gene (most of them uninformative) one can set a threshold for the number of results. The program tries to find the most informative by selecting the most popular ones.

logical indicating whether the output will be a data.frame or list.

#### Value

The output can be either a list or a data.frame. The list has an entry for every input gene. The data frame is a table closely corresponding to the web interface output.

#### Author(s)

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#### References

J. Reimand, M. Kull, H. Peterson, J. Hansen, J. Vilo: g:Profiler – a web-based toolset for functional profiling of gene lists from large-scale experiments (2007) NAR 35 W193-W200

gprofiler 5

#### **Examples**

```
## Not run:
gorth(c("Klf4","Pax5","Sox2","Nanog"), source_organism="mmusculus", target_organism="hsapiens")
## End(Not run)
```

gprofiler

Annotate gene list functionally.

#### **Description**

Interface to the g:Profiler tool for finding enrichments in gene lists. Organism names are constructed by concatenating the first letter of the name and the family name. Example: human - 'hsapiens', mouse - 'mmusculus'. If requesting PNG output, the request is directed to the g:GOSt tool in case 'query' is a vector and the g:Cocoa (compact view of multiple queries) tool in case 'query' is a list. PNG output can fail (return FALSE) in case the input query is too large. In such case, it is advisable to fall back to a non-image request.

#### **Usage**

```
gprofiler(query, organism = "hsapiens", sort_by_structure = T,
  ordered_query = F, significant = T, exclude_iea = F,
  underrep = F, evcodes = F, region_query = F, max_p_value = 1,
  min_set_size = 0, max_set_size = 0, min_isect_size = 0,
  correction_method = "analytical", hier_filtering = "none",
  domain_size = "annotated", custom_bg = "", numeric_ns = "",
  png_fn = NULL, include_graph = F, src_filter = NULL)
```

#### **Arguments**

query vector of gene IDs or a list of such vectors. In the latter case, the query is

directed to g:Cocoa, which yields a different graphical output if requested with

the png\_fn parameter.

organism organism name.

sort\_by\_structure

whether hierarchical sorting is enabled or disabled.

ordered\_query in case output gene lists are ranked this option may be used to get GSEA style

p-values.

significant whether all or only statistically significant results should be returned.

exclude\_iea exclude electronic annotations (IEA).

underrep measure underrepresentation.

evcodes include GO evidence codes as the final column of output. Note that this can

decrease performance and make the query slower.

region\_query interpret query as chromosomal ranges.

6 gprofiler

	max_p_value	custom p-value threshold, results with a larger p-value are excluded.		
	min_set_size	minimum size of functional category, smaller categories are excluded.		
	max_set_size	maximum size of functional category, larger categories are excluded.		
	min_isect_size	minimum size of the overlap (intersection) between query and functional category, smaller intersections are excluded.		
correction_method				
		the algorithm used for determining the significance threshold, one of "gSCS", "fdr", "bonferroni".		
	hier_filtering	hierarchical filtering strength, one of "none", "moderate", "strong".		
	domain_size	statistical domain size, one of "annotated", "known".		
	custom_bg	vector of gene names to use as a statistical background.		
	numeric_ns	namespace to use for fully numeric IDs.		
	png_fn	request the result as PNG image and write it to png_fn.		
	include_graph	request inclusion of network data with the result.		
	<pre>src_filter</pre>	a vector of data sources to use. Currently, these include GO (GO:BP, GO:MF, GO:CC to select a particular GO branch), KEGG, REAC, TF, MI, CORUM, HP, HPA, OMIM. Please see the g:GOSt web tool for the comprehensive list and details on incorporated data sources.		

#### Value

A data frame with the enrichment analysis results. If the input consisted of several lists the corresponding list is indicated with a variable 'query number'. When requesting a PNG image, either TRUE or FALSE, depending on whether a non-empty result was received and a file written or not, respectively. If 'include\_graph' is set, the return value may include the attribute 'networks', containing a list of all network sources, each in turn containing a list of graph edges. The edge structure is a list containing the two interacting symbols and two boolean values (in that order), indicating whether the first or second interactor is part of the input query (core nodes).

#### Author(s)

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#### References

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## Examples

```
## Not run:
gprofiler(c("Klf4", "Pax5", "Sox2", "Nanog"), organism = "mmusculus")
## End(Not run)
```

set\_base\_url 7

set\_base\_url

Set the base URL.

#### **Description**

Set the base URL. Useful for overriding the default URL (http://biit.cs.ut.ee/gprofiler) with the bleeding-edge beta or an archived version.

## Usage

```
set_base_url(url)
```

#### **Arguments**

url

the base URL.

 $set_tls_version$ 

Set the TLS version to use for SSL

## Description

Set the TLS version. Could be useful at environments where SSL was built without TLS 1.2 support

#### Usage

```
set_tls_version(v)
```

#### **Arguments**

٧

version: "1.2" (default), "1.1" (fallback)

set\_user\_agent

Set custom user agent string.

## Description

Set the HTTP User-Agent string. Useful for overriding the default user agent for packages that depend on gProfileR functionality.

## Usage

```
set_user_agent(ua, append = F)
```

8 set\_user\_agent

## Arguments

ua the user agent string.

append logical indicating whether to append the passed string to the default user agent

string.

## **Index**

```
gconvert, 2
get_base_url, 3
get_tls_version, 3
get_user_agent, 3
gorth, 4
gprofiler, 5
set_base_url, 7
set_tls_version, 7
set_user_agent, 7
```