guildifyR

February 2, 2018

get.species.info

Get info on available species and tissues

Description

Get info on available species and tissues

Usage

```
get.species.info()
```

Value

result List containing available species tax ids and names for tissue-specific networks

Examples

```
result = get.species.info()
```

query

Get protein/gene info associated with the query keywords

Description

Get protein/gene info associated with the query keywords

Usage

```
query(keywords, species = "9606", tissue = "All", quote.keywords = T)
```

Arguments

keywords Text containing description of a phenotype or list of genes (seperated by ";") species

Species tax identifier (9606: human, 10090: mouse, etc., see get.species.info

method)

Quotes the keywords to treat whitespaces as ANDs (e.g., "Muscular Dystrophy") quote.keywords

tissues Tissue identifier (All, brain, liver, etc., see get.species.info method) 2 retrieve.job

Value

result.table Data frame containing list of matching proteins/genes and their description

Examples

```
result.table = query("alzheimer", species="10090", tissue="All")
```

retrieve.job

Retrieve results using job id

Description

Retrieve results using job id

Usage

```
retrieve.job(job.id, n.top = NULL, fetch.files = F, output.dir = NULL)
```

Arguments

job.id	Job id
n.top	Number of top proteins to retrieve. If NULL top functionally enriched proteins are provided (upto 500 proteins)
fetch.files	Flag to fetch result files from server and save them locally in output.dir
output.dir	Directory to save the ranking, function, subnetwork and drug info files fetched from the server (if NULL, a folder named same as job.id in the current working directory)

Value

result List containing scores of top-ranking proteins, functions enriched among top-ranking proteins, drugs targeting top-ranking proteins, top-ranking subnetwork cutoff

Examples

```
result = retrieve.job(job.id)
getSlots(class(result))
head(scores(result))
```

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retrieve.overlap Retrieve	overlap between two results using two job ids
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Description

Retrieve overlap between two results using two job ids

Usage

```
retrieve.overlap(job.id1, job.id2, fetch.files = F, output.dir = "./")
```

Arguments

```
job.id1 Job id 1

job.id2 Job id 2

fetch.files Flag to fetch result files from server and save them locally in output.dir

output.dir Directory to save the ranking, function, subnetwork and drug info files fetched
```

from the server

Value

result List containing scores of common top-ranking proteins, common functions enriched among top-ranking proteins, drugs targeting common top-ranking proteins (Note that the number of top-ranking proteins and common functions are limited to 500)

Examples

```
result = retrieve.overlap(job.id1, job.id2)
getSlots(class(result))
head(scores(result))
```

submit.job

Submit job using the guildifyR query result table

Description

Submit job using the guildifyR query result table

Usage

```
submit.job(result.table, species, tissue, scoring.options = list(netcombo =
T))
```

Arguments

```
scoring.options
```

Prioritization method options (default NetCombo)

 $query.\, table \qquad Data\,\, frame\,\, containing\,\, information\,\, from\,\, guildify R\,\, query\,\, (user\,\, entitiy\,\, info,\, net-$

work info, source etc...)

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Value

job id

Examples

```
species="10090"
tissue="All"
result.table = query("alzheimer", species)
#Diamond
scoring.options = list(diamond=T)
#NetScore
scoring.options = list(netscore=T, repetitionSelector=3, iterationSelector=2)
#NetZcore
scoring.options = list(netzcore=T, repetitionZelector=3)
#NetShort
scoring.options = list(netshort=T)
#NetCombo
scoring.options = list(netcombo=T)
job.id = submit.job(result.table, species, tissue, scoring.options)
```

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