DAVIDQuery
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AffyProbesetList  Retrieve Affy probeset IDs from DAVID.

Description
For a given Affymetrix microarray chip, retrieve Affy probeset IDs from DAVID. Optionally, a
menu is used to pick the chip name.

Usage
AffyProbesetList(chipname = NULL, menu = TRUE, verbose=FALSE)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>chipname</td>
<td>Full name or regular expression.</td>
</tr>
<tr>
<td>menu</td>
<td>Select chipname from a menu (default=TRUE)</td>
</tr>
<tr>
<td>verbose</td>
<td>Print a bit of tracing information along the way (default=FALSE)</td>
</tr>
</tbody>
</table>
Details

First, DAVID’s table of chip names is retrieved. When the user selects or specifies one of the names, the associated file of probeset names is retrieved. If chipname is a regular expression, then the menu (if requested) is subsetted accordingly.

Value

Character vector of probeset names.

Note

Use with caution. The returned file is not guaranteed to be correct. In the example above, with the chip "Human Genome U133 Plus 2.0", the list returned includes 40907 probeset IDs on the chip (and no others), but appears to be missing 13768 others.

Author(s)

Roger Day

Examples

```r
head(AffyProbesetList("Human Genome U133 Plus 2.0", menu=FALSE, verbose=TRUE))
## Not run:
length(AffyProbesetList("133|95"))
## End(Not run)
```

---

**affyToUniprot**

**Obtain Affymetrix probeset IDs for given Uniprot IDs.**

Description

Obtain Affymetrix probeset IDs for given Uniprot IDs, using DAVIDQuery.

Usage

```r
affyToUniprot(ids = "88736_AT", ...)
```

Arguments

- `ids` Affymetrix probeset IDs.
- `...` Args to be passed to DAVIDQuery().

Value

The output of DAVIDQuery. If only the DAVIDQueryResult component is desired, include the arg details=FALSE.

Note

There is currently no provision for using DAVIDQueryLoop.
bracketedStrings

Author(s)
Roger Day

See Also
DAVIDQuery

bracketedStrings Extract bracketed substrings.

Description
Extract substrings that are bracketed by specified strings before and after.

Usage
bracketedStrings(s, before, after, verbose=FALSE, addNames=FALSE, drop.na=TRUE, warn.if.gt.1=TRUE)

Arguments
s Vector of strings to search.
before String to the left of the desired substring within s.
after String to the right of the desired substring within s.
verbose If TRUE, print the starting and ending index (or indices) of the desired substring(s).
addNames If TRUE, and if s is a vector, set the names attribute of the return value to s.
drop.na If TRUE, remove empty strings from the return value.
warn.if.gt.1 If TRUE, warn if a string has more than one pair of bracketed target strings.

Value
For a single input string s, the return value is the desired substring sandwiched between before and after. For a vector of inputs, list of outputs.

Author(s)
Roger Day

See Also
DAVIDQuery

Examples
bracketedStrings("quickbrownfox", "quick", "fox")
bracketedStrings(c("quickbrownfox", "quickredfox"), "quick", "fox", addNames=TRUE)
bracketedStrings("quickbrownfoxANDquickredfox", "quick", "fox")
bracketedStrings("quickbrownfoxANDquickredfox", "quick", "fox", warn.if.gt.1=FALSE)
catalogDAVIDResultsByTool

Create a catalog of types of DAVID results.

Description

Loops through values of tool, for the specified value of annot. Runs each DAVID query, and saves the result to create a catalog of types of DAVID results.

Usage

catalogDAVIDResultsByTool(annot = NULL, sleepSeconds = 10, ...)

Arguments

annot See DAVIDQuery.
sleepSeconds DAVIDQueryLoop.
... Extra args passed to DAVIDQuery.

Details

The purpose is to check comprehensively whether there are results that could be better formatted than the default output or the reformatting provided by formatDAVIDResult.

Value

A list of outputs from DAVIDQuery. Automatically assigned to the name catalogOfDAVIDResultsByTool.ANNOT where ANNOT is replaced by the annot argument.

Author(s)

Roger Day

DAVIDQueryLoop

Access DAVID multiple times.

Description

Make a query larger than DAVID allows in one go, by looping, respecting the limitations imposed by DAVID policies.
DAVIDQueryLoop

Usage

DAVIDQueryLoop(
  idList = unlist(strsplit(strsplit("P31946 P62258 P29360 P42655 Q63631
  " "\n")[[1]], "\n")),
  idLimit = 100,
  sleepSeconds = 10,
  hitsPerDayLimit = 200,
  verbose = FALSE,
  testMe = FALSE,
  type,
  annot,
  tool,
  graphicMenu = FALSE,
  formatEach = FALSE,
  formatAll = FALSE,
  ...
)

Arguments

idList  IDs of interest for query.
idLimit Published limit of number of ID’s to process in one call.
sleepSeconds Published minimum time between iterations
hitsPerDayLimit Published maximum URL calls to the API per day from one address.
verbose Print out tracking information as the queries are sent.
testMe Runs DAVIDQueryLoop with arguments set as follows: annot=NULL, tool="geneReportFull",
type="UNIPROT_ACCESSION", verbose=TRUE
annot See DAVIDQuery.
tool See DAVIDQuery.
graphicMenu See DAVIDQuery.
formatEach Passed to DAVIDQuery as the formatIt argument.
formatAll Assembled results are sent to formatDAVIDResult.
...

Value

The results of DAVIDQuery bound together with rbind. Not printed (returned invisibly).

Note

For some choice of the tool argument, the result returned may differ if idLimit is changed.

Author(s)

Roger Day

See Also

DAVIDQuery
Description

Tools to retrieve data from DAVID, the Database for Annotation, Visualization and Integrated Discovery.

Details

Package: DAVIDQuery
Type: Package
Version: 1.0
Date: 2008-10-31
License: GPL-2
LazyLoad: yes
Vignette and overview: vignette("DAVIDQuery")

Author(s)

Roger Day <day@upci.pitt.edu>

References

Home base for DAVID (in this package as DAVIDURBase): http://david.abcc.ncifcrf.gov

Description

Launch a query against DAVID, the Database for Annotation, Visualization and Integrated Discovery. Return the results into an R object.

Usage

DAVIDQuery(ids = "O00161,O75396", type = "UNIPROT_ACCESSION", annot, tool, URLlengthLimit = 2048, details = TRUE, verbose = FALSE, writeHTML = FALSE, testMe = FALSE, graphicMenu = TRUE)
DAVIDQuery

Arguments

ids
IDs for desired objects, as a character vector or as a single string with ids separated by ",". To be passed to the DAVID website, the format has to be the latter.

type
Type of input ids. If missing, a menu is constructed from the R object DAVID-TypeChoices. See http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html#input_list for up-to-date choices.

annot
Type of annotation requested. If missing, a menu is constructed from the R object DAVIDAnnotChoices. See http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html#input_list for up-to-date choices.

tool
Type of gene tool to use. If missing, a menu is constructed from the R object DAVIDToolChoices. See http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html#input_list for up-to-date choices. As of this writing, the tool choices corresponding to the Functional Annotation tools cannot be handled by this package.

URLlengthLimit
Published maximum length of the constructed URL.

details
If TRUE (default), a list of intermediate results is returned; otherwise, just the final query result.

verbose
If TRUE (default is FALSE), more debugging information is printed.

writeHTML
If TRUE (default is FALSE), write the received intermediate HTML to files.

testMe
If TRUE (default is FALSE), assign default values and run.

graphicMenu
If TRUE (default is FALSE), use a GUI window for the pick menus.

formatIt
If TRUE (default), try to interpret the returned character table and structure the result. If false, the unadorned character table returned by DAVID.

Details

The API described at http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html is used. The return is screen-scraped, a new URL is formulated and transmitted, again the return is screen-scraped to find the name of the results file, and finally that file is retrieved into a string matrix.

Obviously this approach is brittle, but it has survived the 2008 DAVID update. A real API would be better, of course.

The return value DAVIDQueryResult is just a character matrix. Its content structure depends on the choices of tool and annotation arguments, so there has been no attempt to manipulate it into, say, a data frame with nice column names.

Value

If detail==FALSE, only DAVIDQueryResult is returned. This a character matrix holding the results of the tab-delimited file returned by DAVID.

If detail==TRUE, a list with contents useful for trouble-shooting:

firstURL
firstStageResult

DAVIDaction
DAVIDToolChoices

secondURL
secondStageResult
hasSessionEnded
downloadFileName
downloadURL
DAVIDQueryResult

Author(s)
Roger Day

References
http://david.abcc.ncifcrf.gov

See Also
DAVIDQueryLoop, formatDAVIDResult

Examples
result = DAVIDQuery(testMe=TRUE)$DAVIDQueryResult
print(length(result))
print(result$firstURL)
print(result$secondURL)
print(result$000161$REFSEQ_MRNA) ## Uses UNIPROT ID's for input.

Description

DAVIDToolChoices, DAVIDTypeChoices, and DAVIDAnnotChoices are string vectors used to construct pick menus, when the corresponding arguments to DAVIDQuery are not provided.

Details

The source of these lists can be found at http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html#input_list The lists posted there may change in the future. No mechanism is included in this package to retrieve and parse the page.

Source

http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html#input_list
DAVIDURLBase

**Description**

Base URL for the DAVID database.

**Source**

http://david.abcc.ncifcrf.gov

**See Also**

DAVIDQuery

DAVIDToolChoices

---

formatDAVIDResult  

*Format the character table returned by DAVID.*

**Description**

These functions attempt to format the character table `result` returned by DAVID.

**Usage**

```
formatDAVIDResult(result, verbose=FALSE)
formatAnnotationReport(result)
formatGeneReport(result)
formatGeneReportFull(result)
formatList(result)
formatGene2Gene(result)
```

**Arguments**

- `result`  
  Character table returned by DAVID.
- `verbose`  
  If TRUE, print tool. Warn if tool="geneReportFull" that the result will be returned invisibly due to its size.
Details

`formatDAVIDResult` switches out to one of `formatGeneReport`, `formatGeneReportFull`, `formatGene2Gene`, or `formatList`, depending on the `tool` argument of `DAVIDQuery()` used to specify what query report to do. The `tool` argument is passed as an attribute attached to `result`.

WARNINGS: Not all values of `tool` have an associated format.

These format utilities are not guaranteed to work correctly for all combinations of inputs into `DAVIDQuery()`, or to continue to work correctly if or when the DAVID API changes. If results appear incorrect, one can use the option `DAVIDQuery(formatIt=FALSE)` to see the unformatted output, and/or paste `DAVIDQuery(details=TRUE)[firstURL]` into a browser.

In the case of `formatGene2Gene`, the `gene` column of the `details` component might not always contain a single identifier.

Value

For `tool == "geneAnnotationReport"`, a list, one component for each element in the `ids` arg. Each component has subcomponents:

- **Gene Name** Self-explanatory.
- **Species** Self-explanatory.
- `<id>` The identifier(s) in the query. The name is whatever the id type was.
- `<other items>` Items produced for the input, specified by the `annot` arg of the query.
  - `<gene name>` Using the same ID type as the `type` argument of `DAVIDQuery()`.
  - **Gene Name** Self-explanatory.
  - **Species** Self-explanatory.
  - **Gene Name** Self-explanatory.
  - **Species** Self-explanatory.
- `<other items>` The union of items produced for the input identifiers (generically called "genes" in DAVID). (The set of attributes is not fixed.)
  - **median** See DAVID documentation.
  - **geo** See DAVID documentation.
  - **diagram** An attempt to parse the fourth column of the Functional Group line of the input. See DAVID documentation and consult the DAVID team.
  - **details** A data frame with columns
    - **gene** "gene" as identified in the `ids` arg to `DAVIDQuery`
    - **geneName** gene name
    - **url** The URL for the Gene Report page at NIAID.

As of this writing, the tool choices corresponding to most Functional Annotation tools cannot be handled by this package.
### testGene2Gene

This function tests and demonstrates the use of DAVIDQuery to access the gene2gene tool.

#### Usage

```r
testGene2Gene(ids = "33246_AT,32469_AT,1786_AT,32680_AT,1355_G_AT,37968_AT,33530_G_AT", type="ENTREZ_GENE_ID")
```

#### Arguments

- `ids` Arg passed to `DAVIDQuery`.
- `type` Arg passed to `DAVIDQuery`.
- `...` Other args passed to `DAVIDQuery`.

#### Details

Input Affy IDS are taken from the example on the DAVID web site.

#### Value

The value returned by DAVIDQuery using tool=gene2gene.

#### Author(s)

Roger Day

#### See Also

- `DAVIDQuery`

#### Examples

```r
testGene2Gene(details=FALSE)
### Run example from http://david.abcc.ncifcrf.gov/gene2gene.jsp
testGene2Gene(ids=idExampleList, type="ENTREZ_GENE_ID", details=FALSE)
### Run example from http://david.abcc.ncifcrf.gov/content.jsp?file=DAVID_API.html
```
uniprotToAffy

Obtain Affymetrix probeset IDs for given Uniprot IDs.

Description

Obtain Affymetrix probeset IDs for given Uniprot IDs, using DAVIDQuery.

Usage

uniprotToAffy(uid = "O00161", ...)

Arguments

uid

Uniprot IDs, either a string with the IDs separated by commas, or else a character vector.

...

Args to be passed to DAVIDQuery().

Value

The output of DAVIDQuery. If only the DAVIDQueryResult component is desired, include the arg details=FALSE. If probesets from a specific chip are desired, then you can intersect these results with the results of AffyProbesetList.

Note

There is currently no provision for using DAVIDQueryLoop.

Author(s)

Roger Day

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