RDAVIDWebService: a versatile R interface to DAVID

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ABSTRACT

Summary: The RDAVIDWebService package provides a class-based interface from R programs/scripts to fully access/control the database for annotation, visualization and integrated discovery, without the need for human interaction on its Web site (http://david.abcc.ncifcrf.gov). The library enhances the database for annotation, visualization and integrated discovery capabilities for Gene Ontology analysis by means of GOstats-based direct acyclic graph conversion methods, in addition to the usual many-genes-to-many-terms visualization.

Availability and implementation: RDAVIDWebService is available as an R package from the Bioconductor project (www.bioconductor.org) and on the authors’ Web site (www.bdmg.com.ar) under GPL-2 license, subjected to the terms of use of DAVID (http://david.abcc. ncifcrf.gov/content.jsp?file=WS.html).

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1 INTRODUCTION

One of the most accessed systems for functional genomics/proteomics analysis is the database for annotation, visualization and integrated discovery (DAVID), a web-based online bioinformatics resource (http://david.abcc.ncifcrf.gov) that aims to provide tools for functional interpretation of large lists of genes/proteins (Huang et al., 2009). Its access is carried mainly through a Web site. There is also a uniform resource locator (URL)-based application programming interface, to query DAVID programmatically, accessible through DAVIDQuery R package (Day and Lisovich, 2010). However, the URL-application programming interface has limited capabilities, such as URL length, and only works with the default settings. In 2012, a web service interface was made available allowing full access and control over all its functionalities except visualization (Jiao et al., 2012). Although it is possible to handle DAVID web services (DWS) through R, it requires high programming skills. In addition, query results are difficult to manage, as they are XML (SOAP package, Temple Lang, 2012) or Java objects (rJava package, Urbanek, 2013).

Here we provide a versatile class-based R interface to access DAVID. It is an R wrapper to all DWS functionalities, with several new features such as offline processing (allows using previously queried saved reports) and native R classes data types. Additionally, it overcomes DWS visualization constraints, providing the usual many-genes-to-many-terms feature, and enhances DAVID capabilities for Gene Ontology (GO, Ashburner et al., 2000) analysis by means of GOstats-based (Falcon and Gentleman, 2007) direct acyclic graph (DAG) conversion methods. Therefore, it expands DAVID features by allowing new developments through one of the most used computer languages in Bioinformatics, R (R Core Team, 2012).

2 IMPLEMENTATION

The package implements a reference class object, by means of R5 paradigm, for DWS communication through a Java client. This allows the establishment of a unique user access point.

To reduce Java-to-R handshaking due to parsing data structures (a time-consuming computational task), the provided Java-based file report client was enhanced to allow formatting of all the DAVID outputs into appropriate RDAVIDWebService S4 R classes. This speeds up the bottleneck data importation process. In addition, it allows locally saving DAVID query to file for further analysis, as well as using Web site-generated reports. Thus, permits using web services and Web site query results interchangeably.

This implementation carries all the limitations of DWS as stated at DAVID’s Web site (http://david.abcc.ncifcrf.gov/content.jsp?file=WS.html): (i) gene or term cluster report will handle up to 3000 genes, (ii) a user or computer can compute up to 200 jobs in a day, and (ii) DAVID Team reserves the right to suspend any improper uses of DWS without notice.

3 FEATURES

Ease of use: It provides a uniform framework to access DAVID analysis straight from R without the need of ad hoc parsing queried reports.

Data import/export: Results from DAVID can be accessed through R or also generated on the Web site. In both cases, they are stored in the same format for later use. This lets on-/offline processing capabilities within R. Thus, queried reports generated anywhere and anywhere can be used for processing, without the need to redo the uploading to DAVID.

Visualization: Customizable many-genes-to-many-terms 2D relationship views are also available with ggplot2 package (Wickham, 2009).

GO structure: DAVID set enrichment analysis or modular enrichment analysis results can be mapped into GOstats-based DAGs. This enables visualization of Expression Analysis Systematic Explorer (EASE) score-based enriched biological process (BP), molecular function and cellular component GO terms in the DAGs. Thus, the exploration and analysis of blurred pattern presence is facilitated, compared with the usual tabular format.
4 EXAMPLE APPLICATION

DAVIDWebService requires a registered DAVID user (this is a prerequisite to use DWS). By means of the registered institutional e-mail, the user can build a DAVIDWebService object and establish a connection. Then, a gene list should be uploaded providing a name and type of list. Here, the one provided in the DAVID Web site is used (demoList1 with Affymetrix® identifiers of 155 genes). Then, the status of the connection is saved in david object.

```r
> david<-DAVIDWebService$new("user@inst.org")
> data(demoList1,
+ idType="AFFYMETRIX_3PRIME_IVT_ID",
+ listName="demoList1", listType="Gene")
> david
DAVIDWebService object to access DAVID’s website.
User email: user@inst.org
Available Gene List/s:
  Name Using
1 demoList1 *
Available Specie/s:
  Name Using
1 Homo sapiens (155) *
Available Background List/s:
  Name Using
1 Homo sapiens *
```

In this example, the complete genome is selected as the default background. If required, the user can select which annotation category to use, e.g. GOTERM_BP_ALL.

A user can obtain the functional annotation cluster report and inspect the results using the following code:

```r
> setAnnotationCategories(david,
+ "GOTERM_BP_ALL")
> termCluster<-+
+ getClusterReport(david,type="Term")
> head(summary(termCluster))
Cluster   Enrichment    Members
1       1 2.9042961     14
2       2 1.9768081     14
3       3 1.3473934     4
4       4 1.3095547     8
5       5 1.2982877     6
6       6 0.8279339     4
```

Then, visually explore the 2D view of a particular cluster (e.g. the third).

```r
> clustNumber<-3
> plot2D(termCluster, clustNumber)
```

However, in Figure 1A, the four term/category members of this cluster share all the ids at ‘system process’ (bottom row).

![Image](https://academic.oup.com/bioinformatics/article-abstract/29/21/2810/196568)

**Fig. 1.** Functional annotation cluster exploration of the third cluster. (A) A 2D view of the evidence of the four term/category members present in the cluster. (B) GO DAG induced by cluster three in (A). Terms with an EASE score <0.1 are shown in gray. In addition, the ratio between genes on the list versus background reference is displayed. Because no information is available regarding the other terms into the mapped GO structure from the cluster in (A), NAs (not available) are introduced when required.

However, as we go up toward the first row (regulation of blood pressure), only four ids have evidence related to it. In this view, neither the hierarchical structure of GO is considered nor the members that are enriched or not (default option). Therefore, the user can extend DAVID’s features obtaining the associated induced DAG structure of the cluster (DAVIDGODag) and contextualize it using GOstats functionalities (plotGOTermGraph, see Fig. 1B).

```r
> davidGODag<-+
+ DAVIDGODag(members(termCluster),
+ 
+ (clustNumber)), pvalueCutoff=0.1, "BP")
> plotGOTermGraph(g=goDag(davidGODag),
+ r=davidGODag, max.nchar=30,
+ node.shape="ellipse")
```

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### REFERENCES


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