# Getting started with go Tools package

Agnes Paquet<sup>1</sup> and (Jean) Yee Hwa Yang<sup>2</sup>
April 21, 2009

### 1. paquetagnes@yahoo.com

2. Department of Medicine, University of California, San Francisco, http://www.biostat.ucsf.edu/jean

### Contents

| 1        | Getting started                                 |
|----------|---|
| <b>2</b> | Graphical comparisons of two sets of genes      |
|          | 2.1 How to use goTools                          |
|          | 2.1.1 Methods for computing percentages         |
|          | 2.2 Plotting the results                        |
| 3        | How to set up the "end nodes"  3.1 Default list |

## 1 Getting started

This document provides a tutorial for the goTools package, which allows graphical comparisons of functional groups between two sets of genes.

Installing the package: To install the *goTools* package, go to the Bioconductor installation web site http://www.bioconductor.org/faq.html for more detailed instructions.

**Help files:** As with any R package, detailed information on functions, classes and methods can be obtained in the help files. For instance, to view the help file for the function ontoCompare in a browser, use help.start() followed by ?ontoCompare.

We demonstrate the functionality with a randomly selected set of probe IDs from both Affymetrix hgu133a chip (affylist). To load the probeID dataset, use data(probeID), and to view a description of the experiments and data, type ?probeID.

**Sweave:** This document was generated using the **Sweave** function from the R *tools* package. The source file is in the /inst.doc directory of the package *goTools*.

To begin, let's load the package and the probeID datasets into your R session.

```
> library("goTools", verbose = FALSE)
> data(probeID)
```

As shown below, affylist is a vector of lists containing 3 list of vectors of probe ids from Affymetrix hgu133a chip.

```
> class(affylist)
[1] "list"
> length(affylist)
[1] 3
> affylist[[1]][1:5]
[1] "215828_at" "201849_at" "219719_at" "213690_s_at" "203172_at"
```

## 2 Graphical comparisons of two sets of genes

Gene Ontology is a Direct Acyclic Graph (DAG) that provides three structured networks of defined terms to describe gene product attributes: Molecular Function (MF), Biological Process (BP) and Cellular Component (CC). A gene product has one or more molecular functions and is used in one or more biological processes; it might be associated with one or more cellular components. To learn more about GO and DAG, please refer to Gene Ontology web site http://www.geneontology.org/.

We have created a set of R functions that use GO structure to describe and compare the composition of sets of genes (or probes). We use the following algorithm:

- 1. Read in a **list** of sets of probe id you want to compare.
- 2. Map each probe id to corresponding ontologies in the GO tree, if any.
- 3. Create the set of GO ids of interest used to compare your datasets (endnode). The function EndNodeList() will create a set of nodes of the DAG located one level under MF, BP or CC, but you can use any sets of GO ids.
- 4. For each GO id, go up the GO tree until reaching the nodes in endnode. Search may be limited to MF, BP or CC if specified in goType.
- 5. Compute the percentage of direct children found under each node in endnode.
- 6. Return the results. Plot them if plot=TRUE.

#### 2.1 How to use goTools

The main function that we provide is ontoCompare. It takes as argument a list of probe ids. Their type must be specified in the argument probeType. For more details about it,you can refer to the corresponding help file by typing: ?ontoCompare.

```
> library(GO.db)
> subset = c(L1 = list(affylist[[1]][1:5]), L2 = list(affylist[[2]][1:5]))
> res <- ontoCompare(subset, probeType = "hgu133a")</pre>
```

#### 2.1.1 Methods for computing percentages

ontoCompare allows you to choose from 3 different methods to estimate the percentage of probes under each element of endnode. The default method is TGenes.

- 1. TGenes: for each end node, return the number of direct children found / total number of probe ids.
  - This includes oligos which do not have GO annotations.
- 2. TIDS: for each end node, return the number of direct children found / total number of GO ids describing the list.
- 3. none: for each end node, return the number of direct children found.

### 2.2 Plotting the results

The plots are produced using the function ontoPlot. It is called by ontoCompare when you set plot=TRUE. You can also call it directly, passing as argument ontoCompare results. If only one set of genes is passed to ontoCompare, ontoPlot will return a pie chart. In other cases, it will return a bargraph. You can modify ontoPlot layout parameters using usual R graphics layout parameters. For more details, type ?par.

```
> library(GO.db)
> subset = c(L1 = list(affylist[[1]][1:5]), L2 = list(affylist[[2]][1:5]))
> res <- ontoCompare(subset, probeType = "hgu133a", plot = TRUE)</pre>
```

## 3 How to set up the "end nodes"

#### 3.1 Default list

The default end nodes list is defined by a call to the function EndNodeList. It contains all children of MF(GO:0003674), BP(GO:0008150) and CC (GO:0005575).

#### > EndNodeList()

| "GD:0008150" | "GO:0005575" | "GD:0003674" |
|--------------|--------------|--------------|
| isa          | isa          | isa          |
| "GO:0045735" | "GD:0003824" | "GD:0009055" |

| isa                  | isa                  | isa          |
|----------------------|----------------------|--------------|
| "GD:0005198"         | "GO:0005215"         | "GD:0005488" |
| isa                  | isa                  | isa          |
| "GO:0031386"         | "GO:0010860"         | "GO:0015457" |
| isa                  | isa                  | isa          |
| "GD:0016209"         | "GO:0016530"         | "GD:0030234" |
| isa                  | isa                  | isa          |
| "GO:0030528"         | "GO:0042056"         | "GD:0045182" |
| isa                  | isa                  | isa          |
| "GO:0045499"         | "GD:0060089"         | "GD:0005576" |
| isa                  | isa                  | isa          |
| "GD:0005623"         | "GO:0019012"         | "GO:0031974" |
| isa                  | isa                  | isa          |
| "GO:0031975"         | "GO:0032991"         | "GD:0043226" |
| isa                  | isa                  | isa          |
| "GO:0044421"         | "GO:0044422"         | "GD:0044423" |
| isa                  | isa                  | isa          |
| "GD:0044456"         | "GD:0044464"         | "GO:0045202" |
| isa                  | isa                  | isa          |
| "GO:0055044"         | "GD:0000003"         | "GO:0008152" |
| isa                  | isa                  | isa          |
| "GD:0001906"         | "GD:0002376"         | "GD:0040007" |
| isa                  | isa                  | isa          |
| "GD:0016032"         | "GD:0009987"         | "GO:0022414" |
| isa                  | isa                  | isa          |
| "GO:0022610"         | "GO:0032501"         | "GO:0032502" |
| isa                  | isa                  | isa          |
| "GD:0040011"         | "GD:0043473"         | "GO:0048511" |
| positively_regulates | negatively_regulates | regulates    |
| "GO:0048518"         | "GO:0048519"         | "GD:0050789" |
| isa                  | isa                  | isa          |
| "GD:0050896"         | "GO:0051179"         | "GO:0051234" |
| isa                  | isa                  |              |
| "GO:0051704"         | "GD:0065007"         |              |
|                      |                      |              |

#### 3.2 Customized end node list

If you want to use more ontologies to describe your set of genes, you can use the function CustomEndNodeList(id,rank) to create a bigger set of end nodes. It returns all GO ids children of id up to rank levels below id.

```
> MFendnode <- CustomEndNodeList("GO:0003674", rank = 2)</pre>
```

Finally, the code below shows you how to use a custom end node list, and also how to modify the goType argument to select only Molecular Function (MF) ontologies.

```
> res <- ontoCompare(subset, probeType = "hgu133a", endnode = MFendnode,
+ goType = "MF")</pre>
```

You can also create a list of GO ids of nodes of interest and pass it directly to the endnode argument in ontoCompare. GO ids must be in the following format: "GO:XXXXXXXX."