

# Package ‘goTools’

September 3, 2025

**Version** 1.82.0

**Date** 2009-10-23

**Title** Functions for Gene Ontology database

**Depends** GO.db

**Imports** AnnotationDbi, GO.db, graphics, grDevices

**Suggests** hgu133a.db

**biocViews** Microarray,GO,Visualization

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**Description** Wrapper functions for description/comparison of oligo ID  
list using Gene Ontology database

**License** GPL-2

**LazyLoad** yes

**git\_url** <https://git.bioconductor.org/packages/goTools>

**git\_branch** RELEASE\_3\_21

**git\_last\_commit** 7c93e7d

**git\_last\_commit\_date** 2025-04-15

**Repository** Bioconductor 3.21

**Date/Publication** 2025-09-03

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EndNodeList

*Reference GO nodes list.*


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## Description

The function `EndNodeList` builds the default end node list used in [ontoCompare](#). `CustomEndNodeList` helps you build a list of children nodes starting from a GO id of interest.

## Usage

```
EndNodeList()
CustomEndNodeList(id,rank=1)
```

## Arguments

|                   |                                                                                         |
|-------------------|-----------------------------------------------------------------------------------------|
| <code>id</code>   | Valid GO id: "GO:XXXXXXX".                                                              |
| <code>rank</code> | Number of levels of children of the GO DAG under id you want to add to your nodes list. |

## Value

`EndNodeList` returns a vector of all GO ids 1 level below MF("GO:0003674"), BP("GO:0008150") and CC("GO:0005575"). MF, BP and CC nodes are included. `CustomEndNodeList` returns a vector of all GO ids children of id, rank levels below it.

## Author(s)

Yee Hwa (Jean) Yang, Agnes Paquet

## See Also

[ontoCompare](#)

## Examples

```
## Examples use the probeID dataset. For description type ? probeID.
## library(GO.db)
## EndNodeList()
## MFendnode <- CustomEndNodeList("GO:0003674", rank=2)

## Example (not run)
## data(probeID)
## ontoCompare(affylist, probeType="hgu133a", endnode=MFendnode, goType="MF")
```

## Description

This functions will allow you to describe and compare sets of oligo ids using Gene Ontology database

## Usage

```
ontoCompare(genelist,probeType=c("GO","hgu133a"),
goType="All", endnode, method=c("TGenes", "TIDS", "none"), plot=FALSE,
...)

ontoPlot(objM, names.arg=NULL,beside=TRUE, las=2,legend.text=TRUE, ...)
```

## Arguments

|             |                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|-------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| genelist    | list of list of valid probe ids.                                                                                                                                                                                                                                                                                                                                                                                                         |
| method      | method used to evaluate the percentage of oligos for each end-node. 'TGenes' = for each end node, return the number of direct children found / total number of probe ids. (default). This includes oligos which do not have GO annotations. 'TIDS' = for each end node, return the number of direct children found / total number of GO ids describing the list. 'none' = for each end node, return the number of direct children found. |
| probeType   | type of input given to the function.Valid probe types include GO ids and any probes ids for which a BioC annotation package providing a mapping to GO is available. ontoCompare is expecting valid probe ids.                                                                                                                                                                                                                            |
| goType      | help sort the data by type. If 'All' (default), all oligos are taken into account. 'BP' restricts information to Biological Process, 'CC' to Cellular Component, and 'MF' to Molecular Function.                                                                                                                                                                                                                                         |
| plot        | logical: if 'TRUE', results are output as a graph.                                                                                                                                                                                                                                                                                                                                                                                       |
| endnode     | list of GO ids corresponding to end-nodes of interest.                                                                                                                                                                                                                                                                                                                                                                                   |
| beside      | Logical. If 'TRUE', the bars of the barplot are portrayed as juxtaposed bars. See ?barplot for more details.                                                                                                                                                                                                                                                                                                                             |
| las         | numeric: if las=2, the axis labels are displayed perpendicular to the axis. See ?par for more details.                                                                                                                                                                                                                                                                                                                                   |
| legend.text | vector of text used to construct a legend for the plot. See ?barplot for more details.                                                                                                                                                                                                                                                                                                                                                   |
| objM        | results from ontoCompare.                                                                                                                                                                                                                                                                                                                                                                                                                |
| names.arg   | Labels to use in ontoPlot.                                                                                                                                                                                                                                                                                                                                                                                                               |
| ...         | extra layout parameters to be passed to ontoPlot.                                                                                                                                                                                                                                                                                                                                                                                        |

**Value**

Returns the percentage of probes children of nodes contained in endnode. If 'plot' = TRUE, results are plotted as a pie chart or a bargraph.

**Author(s)**

Yee Hwa (Jean) Yang, Agnes Paquet

**Examples**

```
# Examples use the probeID dataset. For description type ?probeID.
# Not run

#library(GO.db)
#data(probeID)
#ontoCompare(affylist, probeType="hgu133a", plot=TRUE)
#res <- ontoCompare(operonlist["L1"], probeType="operon", method="TIDS")
#ontoPlot(res, cex=0.7)
```

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|                    |                                   |
|--------------------|-----------------------------------|
| Internal functions | <i>Internal goTools functions</i> |
|--------------------|-----------------------------------|

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**Description**

Internal goTools functions

**Details**

These are not to be called by the user.

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|         |                                                                                         |
|---------|-----------------------------------------------------------------------------------------|
| probeID | <i>List of probe ids from Affymetrix hgu133a chip and Operon Version 2 Human oligos</i> |
|---------|-----------------------------------------------------------------------------------------|

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**Description**

The probeID dataset consists of two lists of randomly chosen probe Ids. affylist contains 3 sets of ids from Affymetrix hgu133a. operonlist contains 2 sets of ids from Operon Version 2.

**Usage**

```
data(probeID)
```

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