How To Use DOSim

Jiang Li

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1 Overview

This vignette demonstrates how to easily use the DOSim package. DOSim is used to calculate DO terms similarity and genes similarity based on terms similarity, and meanwhile it provides information for disease ontology and can do DO Enrichment analysis. We take GOSim[1] as a reference to organize our code.

To start with DOSim package, type following code below.

- > library(DOSim)
- > help(DOSim)

2 Calculate DO Terms Similarity

Terms in disease ontology(DO) are organized in Directed Acyclic Graph (DAG). Previous studies have developed many methods to calculate their similarities, information content(IC) based is the most popular one. In our package, we provide 13 different methods to get DO terms similarity, including IC based and graph based. The function getTerm-Sim is the interface for user to calculate DO terms similarity.

An example of how to calculate DO Terms similarity is shown below:

Detailed information for each method implemented in DOSim is shown below.

2.1 Resnik

Method Resnik [2] is IC based, the similarity between term1 and term2 is the maximum IC of their common ancestors. Defined as

$$IC(term1, term2) = \max_{t \in S(term1, term2)} [IC(t)] = IC_{ms}$$

where S(term1, term2) is the set of terms that subsume both term1 and term2.

2.2 JiangConrath

In 1997, Jay J. Jiang and David W. Conrath[3] proposed a new method and the formular is below:

$$IC(term1, term2) = 1 - \min(1, IC(term1) - 2IC_{ms} + IC(term2))$$

where IC_{ms} is the similarity defiend by Resnik.

2.3 Lin

The formula for Lin[4] is below:

$$IC(term1, term2) = \frac{2IC_{ms}}{IC(term1) + IC(term2)}$$

where IC_{ms} is the similarity defiend by Resnik.

2.4 CoutoEnriched

These method is proposed by Couto in 2003[5], please see the original paper for detail.

2.5 CoutoResnik

It is similar to Resnik, but instead of using common ancestor, the similarity of term1 and term2 is the maximum IC of all the common **disjunctive ancestors** of term1 and term2[6]. It is defined as:

$$IC(term1, term2) = \max_{t \in CommonDisjAnc(term1, term2)} \left[IC(t)\right] = IC_{share}$$

where CommonDisjAnc(term1, term2) is the set of common disjunctive ancestors of term1 and term2.

2.6 CoutoJiangConrath

Similar to JiangConrath, use the Couto's [6] concept and defined as:

$$IC(term1, term2) = 1 - \min(1, IC(term1) - 2IC_{share} + IC(term2))$$

where IC_{share} is the similarity defiend by CoutoResnik.

2.7 CoutoLin

Similar to Lin, use the Couto's [6] concept and defined as:

$$IC(term1, term2) = \frac{2IC_{share}}{IC(term1) + IC(term2)}$$

where IC_{share} is the similarity defiend by CoutoResnik.

2.8 relevance

Proposed by Schlicker[7] in 2006.

$$IC(term1, term2) = Sim_{Lin} * (1 - e^{-IC_{ms}})$$

where Sim_{Lin} is the similarity defined by Lin and IC_{ms} for Resnik.

2.9 GIC

Proposed by Pesquita[8] in 2007.

$$IC(term1, term2) = \frac{\sum\limits_{t \in (Ancestor(term1) \cap Ancestor(term2))} IC(t)}{\sum\limits_{t \in (Ancestor(term1) \cup Ancestor(term2))} IC(t)}$$

where Ancestor(t) is the set of all ancestor terms of term t

$2.10 \quad simIC$

Proposed by Li[9] in 2009.

$$IC(term1, term2) = Sim_{Lin} * \left(1 - \frac{1}{1 + IC_{ms}}\right)$$

where Sim_{Lin} is the similarity defined by Lin and IC_{ms} for Resnik.

2.11 path

This method is not IC based and first proposed by Wu Z[10] in 1994 and mentiond in Pedersen's[11] article in 2007.

$$IC(term1, term2) = \frac{1}{p}$$

where p is the number of nodes on the shortest path between term1 and term2.

2.12 lch

This method is also not IC based and first proposed by Leacock C[12] in 1998 and mentiond in Pedersen's[11] article in 2007.

$$IC(term1, term2) = -\log(\frac{p}{2*depth})$$

where p is the number of nodes on the shortest path between term1 and term2 and depth is the maximum depth of the hierarchy.

2.13 Wang

Proposed by Wang[13] in 2007 and see the original paper for detail.

$$Sim(term1, term2) = \frac{\sum\limits_{t \in T_{term1} \cap T_{term2}} (S_{term1}(t) + S_{term2}(t))}{SV(term1) + SV(term2)}$$

where $S_{term1}(t)$ is the S-value of term t related to term term1. In DO, term term1 can be represented as $DAG_{term1} = (term1, T_{term1}, E_{term1})$ where T_{term1} is the set of DO terms in DAG_{term1} , including term term1 and all of its ancestor terms in the DO graph, and E_{term1} is the set of edges connecting the DO terms in DAG_{term1} . And for any term t in $DAG_{term1} = (term1, T_{term1}, E_{term1})$, its S-value is defined as:

$$\begin{cases} S_{term1}(term1) = 1 \\ S_{term1}(t) = \max\{w_e * S_{term1}(t') | t' \in childrenof(t)\} & if \ t \neq A \end{cases}$$

where w_e is the semantic contribution factor for edge $e \in E_{term1}$ linking term t with its child term t. After obtaining the S-values for all terms in DAG_{term1} , we calculate the semantic value of DO term term1,SV(term1),as:

$$SV(term1) = \sum_{t \in T_{term1}} S_{term1}(t)$$

3 Calculate Genes Similarity

Genes similarity is calculate based on their annotated DO terms similarity. DOSim provides users a function named getGeneSim to calculate genes similarity. It provides 8 methods to calculate genes similarity. A basic example is shown below:

```
> genelist <- c("10003", "10008", "10015", "10042", "10036")
> gsim <- getGeneSim(genelist, similarity = "funSimMax", similarityTerm = "Lin")</pre>
```

> gsim

		10003	10008	10015	10042	10036
10	2003	1.000000000	0	0.003439812	0.002969545	0.281067587
10	8000	0.00000000	1	0.000000000	0.00000000	0.000000000
10	0015	0.003439812	0	1.000000000	0.001945925	0.002137409
10	0042	0.002969545	0	0.001945925	1.000000000	0.001945925
10	0036	0.281067587	0	0.002137409	0.001945925	1.000000000

Here we define some formula and detail information for each method is described below. Assume gene1 have m DO annoated($DO_1 = \{do_{11}, do_{12}, \ldots, do_{1m}\}$) and gene2 have n DO annotated($DO_2 = \{do_{21}, do_{22}, \ldots, do_{2n}\}$). We define Sim_{matrix} is an $m \times n$ matrix of any pairwise DO terms similarity from DO_1 to DO_2 .

$$Sim_{matrix} = \left\{ \begin{array}{cccc} sim_{11} & sim_{12} & \cdots & sim_{1n} \\ sim_{21} & sim_{22} & \cdots & sim_{2n} \\ \vdots & \vdots & \vdots & \vdots \\ sim_{m1} & sim_{m2} & \cdots & sim_{mn} \end{array} \right\}$$

3.1 max

The maximum similarity between any two DO terms.

$$Sim(gene1, gene2) = \max(Sim_{matrix})$$

3.2 mean

The average similarity between any two DO terms

$$Sim(gene1, gene2) = mean(Sim_{matrix})$$

3.3 funSimMax

The average of best matching DO term similarities. Take the maximum of the scores achieved by assignments of DO terms from gene 1 to gene 2 and vice versa.[14]

$$Sim(qene1, qene2) = \max(rowMax, colMax)$$

where rowMax is the average score of each row's maximum score, and same for colMax.

3.4 funSimAvg

The average of best matching DO term similarities. Take the average of the scores achieved by assignments of DO terms from gene 1 to gene 2 and vice versa. [14]

$$Sim(gene1, gene2) = \frac{rowMax + colMax}{2}$$

where rowMax is the average score of each row's maximum score, and same for colMax.

3.5 OA

The optimal assignment (maximally weighted bipartite matching) of DO terms associated to the gene having fewer annotation to the DO terms of the other gene.[15]. See the original paper for details.

3.6 hausdorff

The translation of the Hausdorff distance between two sets:[16] Let A and B be the two sets of DO terms associated to two genes(gene1 and gene2). Then

$$Sim(gene1, gene2) = \min \left(\min \left(\max_{x \in A} (x, y) \right), \min \left(\max_{y \in B} (x, y) \right) \right)$$

3.7 dot

The dot product between feature vectors describing the absence/presence of each DO term. The absence of each DO term is weighted by its information content. Depending on the type of later normalization one can arrive at the cosine similarity (method="sqrt") or at the Tanimoto coefficient (method="Tanimoto").[17].See the original paper for details.

3.8 Wang

Propose by Wang in 2007.[13]. Give two genes gene1 and gene2 annotated by DO term sets $DO_1 = \{do_{11}, do_{12}, \ldots, do_{1m}\}$ and $DO_2 = \{do_{21}, do_{22}, \ldots, do_{2n}\}$ respectively, we define their similarity as:

$$Sim(gene1, gene2) = \frac{\sum\limits_{1 \le i \le m} Sim(do_{1i}, DO_2) + \sum\limits_{1 \le j \le n} Sim(do_{2j}, DO_1)}{m+n}$$

where
$$Sim(do_{1i}, DO_2) = \max_{do_j \in DO_2} (sim(do_{1i}, do_j))$$

4 Get Information of Disease Ontology

The Disease Ontology is a community driven, open source ontology that is designed to link disparate datasets through disease concepts. Terms in DO are organized in Directed Acyclic Graph (DAG). With the work of John D.Osborne in 2009[18], human genes are annotated to DO terms. In DOSim, we provide 7 functions to fetch information of DO terms. They are:

- getParents
- getAncestors
- qetOffsprings
- \bullet getChildren
- getDoTerm
- $\bullet \ getDoAnno$
- getDOGraph

Basic example of each of the 7 functions are show in the following sections below.

4.1 getParents

Returns a list of all direct parents associated to each DO term.

```
> terms <- c("DOID:934", "DOID:1579")
> getParents(terms)

[1] "Start to fetch the parents"
$`DOID:934`
[1] "DOID:95"

$`DOID:1579`
[1] "DOID:13"
```

4.2 getAncestors

Returns the list of all ancestors associated to each DO term.

```
> terms <- c("DOID:934", "DOID:1579")
> getAncestors(terms)
```

```
[1] "Start to fetch the ancestors"
$`DOID:934`
[1] "DOID:0050117" "DOID:2040"
                                  "DOID:4"
                                               "DOID:95"
$`DOID:1579`
[1] "DOID:8" "DOID:2" "DOID:5" "DOID:4" "DOID:13" "DOID:7"
     getOffsprings
4.3
Returns the list of all offspring associated to each DO term.
> terms <- c("DOID:10533", "DOID:550")
> getOffsprings(terms)
[1] "Start to fetch the offsprings"
$`DOID:10533`
[1] "DOID:5460" "DOID:874" "DOID:12017" "DOID:14474" "DOID:10531"
 [6] "DOID:13277" "DOID:10510" "DOID:13275" "DOID:13815" "DOID:12607"
[11] "DOID:5461" "DOID:12888" "DOID:14473" "DOID:10508" "DOID:10509"
[16] "DOID:14338" "DOID:14475" "DOID:13272" "DOID:12608" "DOID:14319"
[21] "DOID:13278" "DOID:13273" "DOID:10457" "DOID:13164" "DOID:12019"
[26] "DOID:11742" "DOID:14472" "DOID:14477" "DOID:11741" "DOID:10532"
[31] "DOID:13167" "DOID:10527" "DOID:14476" "DOID:13276" "DOID:13274"
[36] "DOID:13165" "DOID:873" "DOID:12375"
$`DOID:550`
```

4.4 getChildren

Returns the list of all direct children associated to each DO term.

[1] "DOID:549" "DOID:551" "DOID:554" "DOID:553"

```
> terms <- c("DOID:934", "DOID:1579")
> getChildren(terms)
```

[1] "Start to fetch the children" \$`DOID:934`

```
[1] "DOID:623"
                    "DOID:1329"
                                   "DOID:5064"
                                                   "DOID:1301"
                                                                   "DOID:2950"
[6] "DOID:937"
                    "DOID:2295"
                                   "DOID:0050079" "DOID:13801"
                                                                   "DOID:1274"
[11] "DOID:3294"
                    "DOID:2940"
                                   "DOID:2941"
                                                   "DOID:1304"
                                                                   "DOID:4121"
                                                   "DOID:2931"
                                                                   "DOID:2932"
[16] "DOID:6297"
                    "DOID:1310"
                                   "DOID:4146"
[21] "DOID:2947"
                    "DOID:1885"
                                   "DOID:1331"
                                                   "DOID:2937"
                                                                  "DOID:1385"
[26] "DOID:10533"
```

```
[1] "DOID:11023" "DOID:1585" "DOID:12118" "DOID:12117" "DOID:3224"
                  "DOID:11091" "DOID:13016" "DOID:6144" "DOID:766"
 [6] "DOID:974"
[11] "DOID:550"
4.5
      getDoTerm
Returns the list of DO term's name associated to each DO ID.
> terms <- c("DOID:934", "DOID:1579")
> getDoTerm(terms)
$`DOID:934`
[1] "Virus diseases"
$`DOID:1579`
[1] "respiratory system disease"
4.6
      getDoAnno
Get gene list associated to each DO term
> terms <- c("DOID:934", "DOID:1579")
> getDoAnno(terms)
$`DOID:934`
 [1] "3596"
              "943"
                        "3802"
                                 "941"
                                           "2159"
                                                    "7098"
                                                              "5806"
                                                                       "3837"
 [9] "348"
              "3659"
                        "3665"
                                 "3566"
                                           "29110"
                                                    "60489"
                                                              "939"
                                                                       "282618"
[17] "3105"
              "10859"
                        "4599"
                                 "5133"
                                           "3439"
                                                    "3824"
                                                              "8797"
                                                                       "3491"
[25] "1231"
              "3821"
                        "5322"
                                 "57062"
                                           "3661"
                                                    "1487"
                                                              "3567"
                                                                       "796"
[33] "708"
                        "103"
                                 "3565"
                                                    "3576"
                                                              "4277"
              "2022"
                                           "4000"
                                                                       "5058"
[41] "3553"
              "6504"
                        "325"
                                 "942"
                                           "3627"
                                                    "64135"
                                                              "3554"
                                                                       "3456"
                                                    "282616" "929"
[49] "332"
              "3998"
                        "3586"
                                 "3106"
                                           "3265"
                                                                       "59067"
[57] "5932"
              "3676"
                        "3620"
                                 "5371"
                                           "10010"
                                                    "842"
                                                                       "1616"
                                                              "4153"
[65] "5366"
              "3438"
                        "1234"
                                 "10344"
                                           "4001"
                                                    "3609"
                                                              "1147"
                                                                       "57506"
[73] "10219"
              "3838"
                        "7293"
                                 "6041"
                                           "10673"
$`DOID:1579`
```

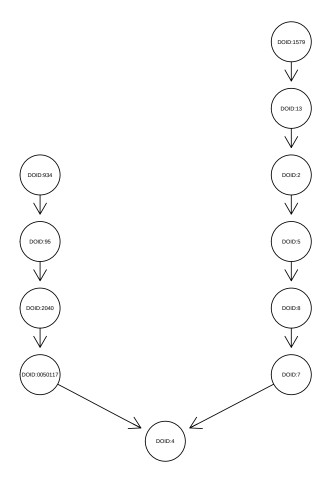
4.7 getDOGraph

[1] "1636"

\$`DOID:1579`

Get DO graph with specified DO terms at its leave.

```
> terms <- c("DOID:934", "DOID:1579")
> if (require(graph)) {
+          g <- getDOGraph(terms)
+          if (require(Rgraphviz)) {
+              plot(g)
+          }
+ }</pre>
```



5 DO Enrichment Analysis

DOSim can do DO enrichment analysis for a list of Entrez gene ids by using **hyper geometric test** or **fisher test**. To do it, you can simply invoke the function *DOEnrichment*. Here is an example.

```
> genelist = as.character(1:100)
> DOEnrichment(genelist, method = "hypertest", filter = 50, cutoff = 0.001)
```

	DOID	pvalue	odds	genenum1	genenum2
DOID:14330	DOID:14330	3.732400e-07	18.068317	101	5
DOID:10652	DOID:10652	2.854039e-05	8.527570	214	5
DOID:759	DOID:759	3.416859e-05	8.257466	221	5
DOID:10591	DOID:10591	2.537661e-04	9.864324	111	3
DOID:12603	DOID:12603	3.777357e-04	14.312941	51	2
DOID:3683	DOID:3683	4.000677e-04	14.037692	52	2
DOID:722	DOID:722	4.232310e-04	13.772830	53	2
DOID:9074	DOID:9074	4.761334e-04	8.358321	131	3
DOID:10825	DOID:10825	5.519650e-04	12.585517	58	2
DOID:10283	DOID:10283	5.594589e-04	4.243953	516	6
DOID:3300	DOID:3300	8.417936e-04	10.894925	67	2
DOID:2370	DOID:2370	8.417936e-04	10.894925	67	2
DOID:12849	DOID:12849	8.789028e-04	10.734706	68	2

References

- [1] Frohlich H, Speer N, Poustka A, BeiSZbarth T: GOSim an R-package for computation of information theoretic GO similarities between terms and gene products. *BMC Bioinformatics* 2007, 8:166.
- [2] Resnik P: Using Information Content to Evaluate Semantic Similarity in a Taxonomy. Proceedings of the 14th International Joint Conference on Artificial Intelligence, Montreal 1995, 1:448–453.
- [3] Jiang J, Conrath D: Semantic similarity based on corpus statistics and lexical taxonomy. Proceedings of the International Conference on Research in Computational Linguistics, Taiwan 1998.
- [4] Jiang J, Conrath D: Semantic similarity based on corpus statistics and lexical taxonomy. Proceedings of the International Conference on Research in Computational Linguistics, Taiwan 1998.
- [5] Couto F, Silva M, Coutinho P: Implementation of a Functional Semantic Similarity Measure between Gene-Products. Tech Rep DI/FCUL TR 03-29 2003.
- [6] Couto F, Silva M, Coutinho P: Semantic Similarity over the Gene Ontology: Family Correlation and Selecting Disjunctive Ancestors. Conference in Information and Knowledge Management 2005.
- [7] A Schlicker FD: A new measure for functional similarity of gene products based on Gene Ontology. BMC Bioinformatics 2006.

- [8] C Pesquita DF: Evaluating GO-based Semantic Similarity Measures. *In:* Proc. 10th Annual Bio-Ontologies Meeting 2007, :37–40.
- [9] B Li AF J Wang: Effectively Integrating Information Content and Structural Relationship to Improve the GO-based Similarity Measure Between Proteins. *BMC Bioinformatics* 2009.
- [10] Wu Z PM: Verb semantics and lexical selection. In:Proceedings of the 32nd annual meeting of the association for computational linguistics 1994, :133–8.
- [11] Pedersen T, Pakhomov SV, Patwardhan S, Chute CG: Measures of semantic similarity and relatedness in the biomedical domain. *Journal of Biomedical Informatics* 2007, **40**(3):288 299.
- [12] Leacock C CM: Combining local context and WordNet similarity for word sense identification. In: Fellbaum C, editor. WordNet: An electronic lexical database. Cambridge, MA: MIT Press 1998, :265–83.
- [13] James ZWang ZD: A new method to measure the semantic similarity of GO terms. *Bioinformatics* 2007, :1274–1281.
- [14] A Schlicker FD: A new measure for functional similarity of gene products based on Gene Ontology. *BMC Bioinformatics* 2006.
- [15] H Froehlich NS: **Kernel Based Functional Gene Grouping**. Proc. Int. Joint Conf. on Neural Networks (IJCNN) 2006, :6886 6891.
- [16] A del Pozo AV F Pazos: **Defining functional distances over Gene Ontology**. *BMC Bioinformatics* 2008, :9:50.
- [17] M Mistry PP: Gene Ontology term overlap as a measure of gene functional similarity. *BMC Bioinformatics* 2008, :9:327.
- [18] Osborne J, Flatow J, Holko M, Lin S, Kibbe W, Zhu L, Danila M, Feng G, Chisholm R: **Annotating the human genome with Disease Ontology**. *BMC Genomics* 2009, **10**(Suppl 1):S6.