

# Disease Ontology Semantic and Enrichment analysis

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## Abstract

Disease Ontology (DO) aims to provide an open source ontology for the integration of biomedical data that is associated with human disease. We developed *DOSE* package to promote the investigation of diseases. *DOSE* provides five methods including Resnik, Lin, Jiang, Rel and Wang for measuring semantic similarities among DO terms and gene products; Hypergeometric model and gene set enrichment analysis were also implemented for extracting disease association insight from genome wide expression profiles.

**DOSE version:** 2.6.6

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## 1 Introduction

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Public health is an important driving force behind biological and medical research. A major challenge of the post-genomic era is bridging the gap between fundamental biological research and its clinical applications. Recent research has increasingly demonstrated that many seemingly dissimilar diseases have common molecular mechanisms. Understanding similarities among disease aids in early diagnosis and new drug development.

Formal knowledge representation of gene-disease association is demanded for this purpose. Ontologies, such as Gene Ontology, have been successfully applied to represent biological knowledge, and many related techniques have been adopted to extract information. Disease Ontology (DO) [1] was developed to create a consistent description of gene products with disease perspectives, and is essential for supporting functional genomics in disease context. Accurate disease descriptions can discover new relationships between genes and disease, and new functions for previous uncharacterized genes and alleles.

Unlike other clinical vocabularies that defined disease related concepts disparately, DO is organized as a directed acyclic graph, laying the foundation for quantitative computation of disease knowledge. The application of disease ontology is in its infancy, lacking programs for mining DO knowledge automatically.

Here, we present an R package *DOSE* [2] for analyzing semantic similarities among DO terms and gene products annotated with DO terms, and extracting disease association insight from genome wide expression profiles.

Four information content (IC)-based methods and one graph structure-based method were implemented for measuring semantic similarity. Hypergeometric test and Gene Set Enrichment Analysis were implemented for extracting biological insight.

To start with *DOSE* package, type following code below:

```
library(DOSE)
help(DOSE)
```

## 2 DO term semantic similarity measurement

---

Four methods determine the semantic similarity of two terms based on the Information Content of their common ancestor term were proposed by Resnik [3], Jiang [4], Lin [5] and Schlicker [6]. Wang [7] presented a method to measure the similarity based on the graph structure. Each of these methods has its own advantage and weakness. *DOSE* implemented all these methods to compute semantic similarity among DO terms and gene products. We have developed another package *GOSemSim* [8] to explore the functional similarity at GO perspective, including molecular function (MF), biological process (BP) and cellular component (CC).

## 2.1 Information content-based method

Information content (IC) is defined as the negative logarithm of the frequency of each term occurs in the corpus of DO annotation.

The frequency of a term  $t$  is defined as:

$$p(t) = \frac{n_{t'}}{N} | t' \in \{t, \text{children of } t\}$$

where  $n_{t'}$  is the number of term  $t'$ , and  $N$  is the total number of terms in DO corpus.

Thus the information content is defined as:

$$IC(t) = -\log(p(t))$$

IC-based methods calculate similarity of two DO terms based on the information content of their closest common ancestor term, which was also called most informative information ancestor (MICA).

### 2.1.1 Resnik method

The Resnik method is defined as:

$$sim_{Resnik}(t_1, t_2) = IC(MICA)$$

### 2.1.2 Lin method

The Lin method is defined as:

$$sim_{Lin}(t_1, t_2) = \frac{2IC(MICA)}{IC(t_1) + IC(t_2)}$$

### 2.1.3 Rel method

The Relevance method, which was proposed by Schlicker, combine Resnik's and Lin's method and is defined as:

$$sim_{Rel}(t_1, t_2) = \frac{2IC(MICA)(1 - p(MICA))}{IC(t_1) + IC(t_2)}$$

### 2.1.4 Jiang method

The Jiang and Conrath's method is defined as:

$$sim_{Jiang}(t_1, t_2) = 1 - \min(1, IC(t_1) + IC(t_2) - 2IC(MICA))$$

## 2.2 Graph-based method

Graph-based methods using the topology of DO graph structure to compute semantic similarity. Formally, a DO term A can be represented as  $DAG_A = (A, T_A, E_A)$  where  $T_A$  is the set of DO terms in  $DAG_A$ , including term A and all of its ancestor terms in the DO graph, and  $E_A$  is the set of edges connecting the DO terms in  $DAG_A$ .

### 2.2.1 Wang method

To encode the semantic of a DO term in a measurable format to enable a quantitative comparison, Wang firstly defined the semantic value of term A as the aggregate contribution of all terms in  $DAG_A$  to the semantics of term A, terms closer to term A in  $DAG_A$  contribute more to its semantics. Thus, defined the contribution of a DO term  $t$  to the semantic of DO term A as the S-value of DO term  $t$  related to term A. For any of term  $t$  in  $DAG_A$ , its S-value related to term A,  $S_A(t)$  is defined as:

$$\begin{cases} S_A(A) = 1 \\ S_A(t) = \max\{w_e \times S_A(t') | t' \in \text{children of}(t)\} \text{ if } t \neq A \end{cases}$$

where  $w_e$  is the semantic contribution factor for edge  $e \in E_A$  linking term  $t$  with its child term  $t'$ . Term A contributes to its own is defined as one. After obtaining the S-values for all terms in  $DAG_A$ , the semantic value of DO term A,  $SV(A)$ , is calculated as:

$$SV(A) = \sum_{t \in T_A} S_A(t)$$

Thus given two DO terms A and B, the semantic similarity between these two terms is defined as:

$$\text{sim}_{Wang}(A, B) = \frac{\sum_{t \in T_A \cap T_B} S_A(t) + S_B(t)}{SV(A) + SV(B)}$$

where  $S_A(t)$  is the S-value of DO term  $t$  related to term A and  $S_B(t)$  is the S-value of DO term  $t$  related to term B.

## 2.3 doSim function

In *DOSE*, we implemented all these IC-based and graph-based methods. `doSim` can calculate semantic similarity between two DO terms and two set of DO terms.

```
data(DO2EG)
set.seed(123)
a <- sample(names(DO2EG), 10)
a

## [1] "DOID:1407" "DOID:5844" "DOID:2034" "DOID:8432" "DOID:9146"
## [6] "DOID:10584" "DOID:3209" "DOID:848" "DOID:3341" "DOID:2512"
```

```

b <- sample(names(DO2EG), 5)
b
## [1] "DOID:9409" "DOID:2481" "DOID:4465" "DOID:3498" "DOID:11252"
doSim(a[1], b[1], measure="Wang")
## [1] 0.113
doSim(a[1], b[1], measure="Resnik")
## [1] 0.0809
doSim(a[1], b[1], measure="Lin")
## [1] 0.0941
s <- doSim(a, b, measure="Wang")
s
##           DOID:9409 DOID:2481 DOID:4465 DOID:3498 DOID:11252
## DOID:1407      0.1133    0.0860    0.0152    0.0152    0.0819
## DOID:5844      0.1490    0.0783    0.0280    0.0280    0.1156
## DOID:2034      0.1735    0.4755    0.0368    0.0368    0.1388
## DOID:8432      0.1735    0.1000    0.0368    0.0368    0.4220
## DOID:9146      0.0714    0.0412    0.0368    0.0368    0.0571
## DOID:10584     0.1211    0.0987    0.0180    0.0180    0.0893
## DOID:3209      0.1490    0.0783    0.0280    0.0280    0.1156
## DOID:848       0.1490    0.0783    0.0280    0.0280    0.1156
## DOID:3341      0.1324    0.0637    0.0221    0.0221    0.1000
## DOID:2512      0.0714    0.0412    0.0368    0.0368    0.0571

```

doSim requires three parameter DOID1, DOID2 and measure. DOID1 and DOID2 should be a vector of DO terms, while measure should be one of Resnik, Jiang, Lin, Rel, and Wang.

We also implement a plot function simplot to visualize the similarity result.

```

simplot(s,
  color.low="white", color.high="red",
  labs=TRUE, digits=2, labs.size=5,
  font.size=14, xlab="", ylab="")

```

Parameter color.low and color.high are used to setting the color gradient; labs is a logical parameter indicating whether to show the similarity values or not, digits to indicate the number of decimal places to be used and labs.size setting the size of similarity values; font.size setting the font size of axis and label of the coordinate system.

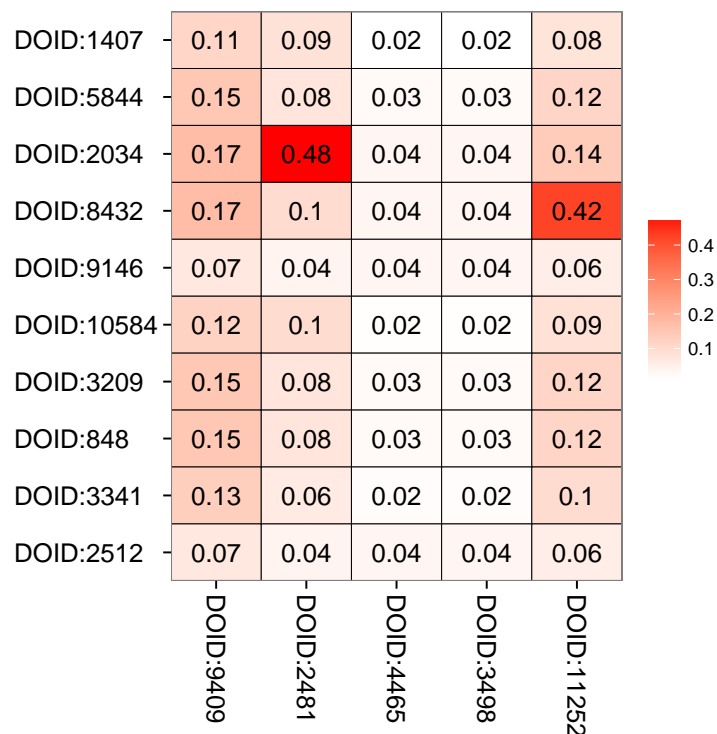


Figure 1: Visualizing similarity matrix.

### 3 Gene semantic similarity measurement

On the basis of semantic similarity between DO terms, *DOSE* can also compute semantic similarity among gene products.

Suppose we have gene  $g_1$  annotated by DO term set  $DO_1 = \{do_{11}, do_{12} \cdots do_{1m}\}$  and  $g_2$  annotated by  $DO_2 = \{do_{21}, do_{22} \cdots do_{2n}\}$ , *DOSE* implemented four methods which called max, avg, rcmax and BMA to combine semantic similarity scores of multiple DO terms.

#### 3.1 Combine method

##### 3.1.1 max

The max method calculates the maximum semantic similarity score over all pairs of DO terms between these two DO term sets.

$$sim_{max}(g_1, g_2) = \max_{1 \leq i \leq m, 1 \leq j \leq n} sim(do_{1i}, do_{2j})$$

### 3.1.2 avg

The avg calculates the average semantic similarity score over all pairs of DO terms.

$$sim_{avg}(g_1, g_2) = \frac{\sum_{i=1}^m \sum_{j=1}^n sim(do_{1i}, do_{2j})}{m \times n}$$

### 3.1.3 rcmax

Similarities among two sets of DO terms form a matrix, the rcmax method uses the maximum of RowScore and ColumnScore as the similarity, where RowScore (or ColumnScore) is the average of maximum similarity on each row (or column).

$$sim_{rcmax}(g_1, g_2) = \max\left(\frac{\sum_{i=1}^m \max_{1 \leq j \leq n} sim(do_{1i}, do_{2j})}{m}, \frac{\sum_{j=1}^n \max_{1 \leq i \leq m} sim(do_{1i}, do_{2j})}{n}\right)$$

### 3.1.4 BMA

The BMA method, used the best-match average strategy, calculates the average of all maximum similarities on each row and column, and is defined as:

$$sim_{BMA}(g_1, g_2) = \frac{\sum_{i=1}^m \max_{1 \leq j \leq n} sim(do_{1i}, do_{2j}) + \sum_{j=1}^n \max_{1 \leq i \leq m} sim(do_{1i}, do_{2j})}{m + n}$$

## 3.2 geneSim function

In *DOSE*, we implemented geneSim to measure semantic similarities among genes.

```
data(EG2D0)
g1 <- sample(names(EG2D0), 5)
g1
## [1] "84842" "2521" "10592" "3069" "91746"

g2 <- sample(names(EG2D0), 4)
g2
## [1] "84289" "6045" "56999" "9869"

geneSim(g1[1], g2[1], measure="Wang", combine="BMA")
## [1] 0.057
```



```
gs <- geneSim(g1, g2, measure="Wang", combine="BMA")
gs
##          84289   6045 56999   9869
## 84842 0.057 0.135 0.355 0.098
## 2521  0.573 0.253 0.511 0.482
## 10592 0.057 0.187 0.296 0.128
## 3069  0.573 0.517 1.000 1.000
## 91746 0.573 0.308 0.527 0.501
```

geneSim requires four parameter geneID1, geneID2, measure and combine. geneID1 and geneID2 should be a vector of entrez gene IDs, measure should be one of Resnik, Jiang, Lin, Rel, and Wang, while combine should be one of max, avg, rcmax and BMA as described previously.

The simplot works well with both the output of doSim and geneSim.

## 4 DO term enrichment analysis

---

### 4.1 Hypergeometric model

Enrichment analysis [9] is a widely used approach to identify biological themes. Here we implement hypergeometric model to assess whether the number of selected genes associated with disease is larger than expected.

To determine whether any DO terms annotate a specified list of genes at frequency greater than that would be expected by chance, *DOSE* calculates a p-value using the hypergeometric distribution:

$$p = 1 - \sum_{i=0}^{k-1} \frac{\binom{M}{i} \binom{N-M}{n-i}}{\binom{N}{n}}$$

In this equation,  $N$  is the total number of genes in the background distribution,  $M$  is the number of genes within that distribution that are annotated (either directly or indirectly) to the node of interest,  $n$  is the size of the list of genes of interest and  $k$  is the number of genes within that list which are annotated to the node. The background distribution by default is all the genes that have DO annotation.

P-values were adjusted for multiple comparison, and q-values were also calculated for FDR control.

### 4.2 enrichDO function

*DOSE* provides an example dataset *geneList* which was derived from R package *breastCancerMAINZ* that contained 200 samples, including 29 samples in grade I, 136 samples in grade II and 35 samples in grade III. We computed the ratios of geometric means of grade III samples versus geometric means of grade I samples. Logarithm of these ratios (base 2) were stored in *geneList* dataset.

In the following example, we selected fold change above 1 as the differential genes and analyzing their disease association.

```
data(geneList)
gene <- names(geneList)[abs(geneList) > 1.5]
head(gene)

## [1] "4312" "8318" "10874" "55143" "55388" "991"

x <- enrichDO(gene      = gene,
               ont       = "DO",
               pvalueCutoff = 0.05,
               pAdjustMethod = "BH",
               universe    = names(geneList),
               minGSSize   = 5,
               readable    = FALSE)
head(summary(x))

##              ID              Description GeneRatio  BgRatio
## DOID:162      DOID:162      cancer      266/331 4259/6274
## DOID:14566    DOID:14566 disease of cellular proliferation 267/331 4307/6274
## DOID:0050686 DOID:0050686 organ system cancer      187/331 2756/6274
## DOID:2994     DOID:2994     germ cell cancer      47/331  483/6274
## DOID:193      DOID:193      reproductive organ cancer 61/331  691/6274
## DOID:10283    DOID:10283    prostate cancer    40/331  394/6274
##              pvalue p.adjust  qvalue
## DOID:162      1.41e-07 0.000123 9.48e-05
## DOID:14566    3.27e-07 0.000143 1.10e-04
## DOID:0050686 1.61e-06 0.000469 3.61e-04
## DOID:2994     2.27e-05 0.004538 3.50e-03
## DOID:193      2.60e-05 0.004538 3.50e-03
## DOID:10283    3.78e-05 0.005496 4.23e-03
##
## DOID:162      4312/8318/10874/55143/991/6280/2305/1062/4605/9833/9133/6279/10403/597/7153
## DOID:14566    4312/8318/10874/55143/991/6280/2305/1062/4605/9833/9133/6279/10403/597/7153
## DOID:0050686
## DOID:2994
## DOID:193
## DOID:10283
##              Count
## DOID:162      266
## DOID:14566    267
## DOID:0050686  187
## DOID:2994      47
## DOID:193       61
## DOID:10283     40
```

The `enrichDO` requires an entrezgene ID vector as input, mostly is the differential gene list of gene expression profile studies. The `ont` parameter can be "DO" or "DOLite", DOLite [10] was constructed to aggregate the redundant DO terms. The DOLite data is not updated, we recommend user use `ont="DO"`. `pvalueCutoff` setting the cutoff value of p value and p value adjust; `pAdjustMethod` setting the p value correction methods, include the Bonferroni correction ("bonferroni"), Holm ("holm"), Hochberg ("hochberg"), Hommel ("hommel"), Benjamini & Hochberg ("BH") and Benjamini & Yekutieli ("BY").

The `universe` setting the background gene universe for testing. If user do not explicitly setting this parameter, `enrichDO` will set the universe to all human genes that have DO annotation.

The `minGSSize` indicates that only those DO terms that have more than `minGSSize` genes annotated will be tested.

The `readable` is a logical parameter, indicates whether the entrezgene IDs will mapping to gene symbols or not.

We also implement `setReadable` function that helps the user to convert entrezgene IDs to gene symbols.

```
x <- setReadable(x)
head(summary(x))
```

##	ID	Description	GeneRatio	BgRatio
## DOID:162	DOID:162	cancer	266/331	4259/6274
## DOID:14566	DOID:14566	disease of cellular proliferation	267/331	4307/6274
## DOID:0050686	DOID:0050686	organ system cancer	187/331	2756/6274
## DOID:2994	DOID:2994	germ cell cancer	47/331	483/6274
## DOID:193	DOID:193	reproductive organ cancer	61/331	691/6274
## DOID:10283	DOID:10283	prostate cancer	40/331	394/6274
##	pvalue	p.adjust	qvalue	
## DOID:162	1.41e-07	0.000123	9.48e-05	
## DOID:14566	3.27e-07	0.000143	1.10e-04	
## DOID:0050686	1.61e-06	0.000469	3.61e-04	
## DOID:2994	2.27e-05	0.004538	3.50e-03	
## DOID:193	2.60e-05	0.004538	3.50e-03	
## DOID:10283	3.78e-05	0.005496	4.23e-03	
##				
## DOID:162	MMP1/CDC45/NMU/CDCA8/CDC20/S100A9/FOXN1/CENPE/MYBL2/MELK/CCNB2/S100A8/NDC80			
## DOID:14566	MMP1/CDC45/NMU/CDCA8/CDC20/S100A9/FOXN1/CENPE/MYBL2/MELK/CCNB2/S100A8/NDC80			
## DOID:0050686				
## DOID:2994				
## DOID:193				
## DOID:10283				
##	Count			
## DOID:162	266			
## DOID:14566	267			
## DOID:0050686	187			

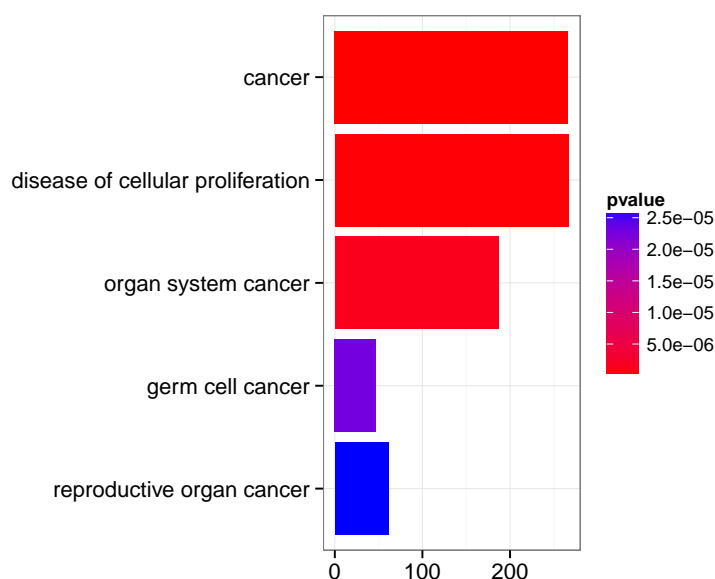


Figure 2: barplot of DO enrichment result.

```
## DOID:2994      47
## DOID:193       61
## DOID:10283    40
```

### 4.3 Visualize enrichment result

We also implement a bar plot and category-gene-network for visualization. It is very common to visualize the enrichment result in bar or pie chart. We believe the pie chart is misleading and only provide bar chart.

```
barplot(x)
```

In order to consider the potentially biological complexities in which a gene may belong to multiple annotation categories, we developed `cnetplot` function to extract the complex association between genes and diseases.

```
cnetplot(x, categorySize="pvalue", foldChange=geneList)
```

### 4.4 Disease association comparison

We have developed an R package *clusterProfiler* [11] for comparing biological themes among gene clusters. *DOSE* works fine with *clusterProfiler* and can compare biological themes at disease perspective.



```
require(clusterProfiler)
data(gcSample)
cdo <- compareCluster(gcSample, fun="enrichD0")
plot(cdo)
```

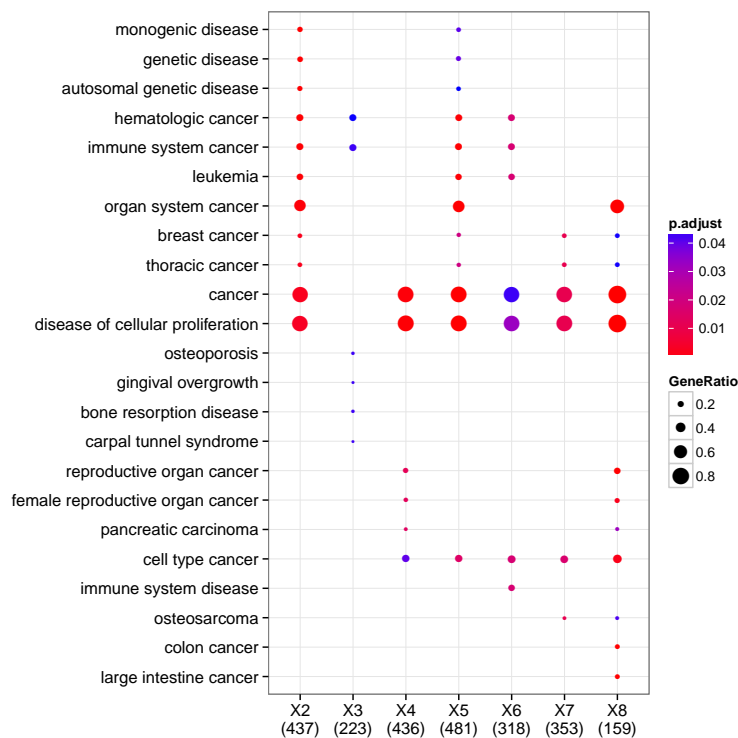


Figure 4: DOSE with clusterProfiler.

## 5 Gene set enrichment analysis

### 5.1 GSEA algorithm

A common approach in analyzing gene expression profiles was identifying differential expressed genes that are deemed interesting. The DO term enrichment analysis we demonstrated previous were based on these differential expressed genes. This approach will find genes where the difference is large, but it will not detect a situation where the difference is small, but evidenced in coordinated way in a set of related genes. Gene Set Enrichment Analysis (GSEA) [12] directly addresses this limitation. All genes can be used in GSEA; GSEA aggregates the per gene statistics across genes within a gene set, therefore making it possible to detect situations where all genes in a predefined set change in a small but coordinated way. Since it is likely that many relevant phenotypic differences are manifested by small but consistent changes in a set of genes.

Genes are ranked based on their phenotypes. Given a priori defined set of genes  $S$  (e.g., genes sharing the same *DO* or *DOLite* category), the goal of GSEA is to determine whether the members of  $S$  are randomly distributed throughout the ranked gene list ( $L$ ) or primarily found at the top or bottom.

There are three key elements of the GSEA method:

- Calculation of an Enrichment Score.

The enrichment score ( $ES$ ) represent the degree to which a set  $S$  is over-represented at the top or bottom of the ranked list  $L$ . The score is calculated by walking down the list  $L$ , increasing a running-sum statistic when we encounter a gene in  $S$  and decreasing when it is not. The magnitude of the increment depends on the gene statistics (e.g., correlation of the gene with phenotype). The  $ES$  is the maximum deviation from zero encountered in the random walk; it corresponds to a weighted Kolmogorov-Smirnov-like statistic [12].

- Estimation of Significance Level of  $ES$ .

The  $p$ -value of the  $ES$  is calculated using permutation test. Specifically, we permute the gene labels of the gene list  $L$  and recompute the  $ES$  of the gene set for the permuted data, which generate a null distribution for the  $ES$ . The  $p$ -value of the observed  $ES$  is then calculated relative to this null distribution.

- Adjustment for Multiple Hypothesis Testing.

When the entire  $DO$  or  $DOLite$  gene sets is evaluated,  $DOSE$  adjust the estimated significance level to account for multiple hypothesis testing and also  $q$ -values were calculated for FDR control.

## 5.2 gseAnalyzer fuction

In *DOSE*, we implemented GSEA algorithm proposed by Subramanian [12] in *gseAnalyzer* function.

In the following example, in order to speedup the compilation of this document, only gene sets with size above 120 were tested and only 100 permutations were performed.

```
y <- gseAnalyzer(geneList,
                 setType      = "DO",
                 nPerm        = 100,
                 minGSSize    = 120,
                 pvalueCutoff  = 0.05,
                 pAdjustMethod = "BH",
                 verbose       = FALSE)
res <- summary(y)
head(res)
```

##	ID	Description	setSize		
##	DOID:0050117 DOID:0050117	disease by infectious agent	851		
##	DOID:0050161 DOID:0050161	lower respiratory tract disease	455		
##	DOID:0050177 DOID:0050177	monogenic disease	575		
##	DOID:0050338 DOID:0050338	primary bacterial infectious disease	214		
##	DOID:0050615 DOID:0050615	respiratory system cancer	558		
##	DOID:0050686 DOID:0050686	organ system cancer	2756		
##	enrichmentScore	pvalue	p.adjust	qvalues	
##	DOID:0050117	0.311	0.0099	0.0198	0.00643
##	DOID:0050161	0.300	0.0099	0.0198	0.00643
##	DOID:0050177	0.230	0.0099	0.0198	0.00643
##	DOID:0050338	0.457	0.0099	0.0198	0.00643
##	DOID:0050615	0.335	0.0099	0.0198	0.00643

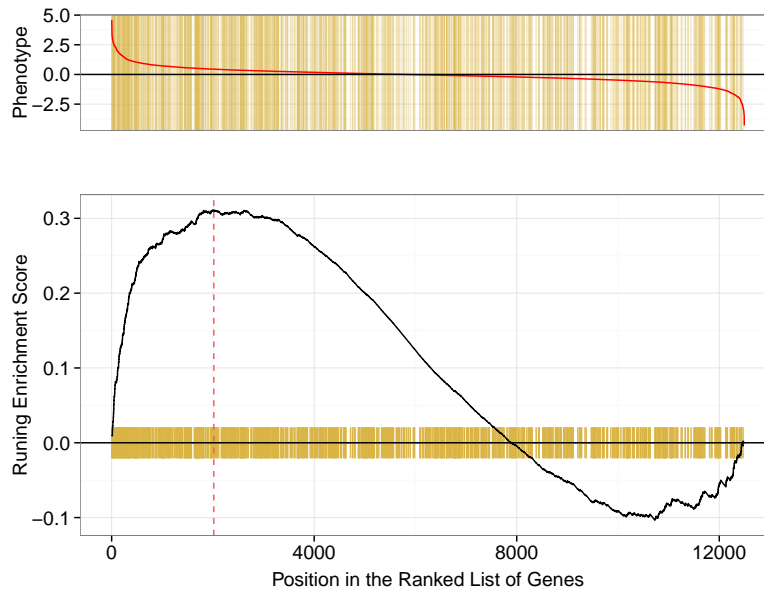


Figure 5: gseaplot example.

```
## D0ID:0050686          0.251 0.0099    0.0198 0.00643
```

The `setType` should be one of "DO" or "DOLite" and was required for `gseaAnalyzer` to prepare the corresponding gene sets.

```
topID <- res[1,1]
topID
## [1] "D0ID:0050117"
plot(y, geneSetID = topID)
```

Parameter `geneSetID` can be numeric, the following command will generate the same figure as illustrated above.

```
plot(y, geneSetID = 1)
```

## 6 enrichMap

Enrichment Map can be visualized by `enrichMap` function. It supports both enrichment result and GSEA result.

```
enrichMap(x)
```



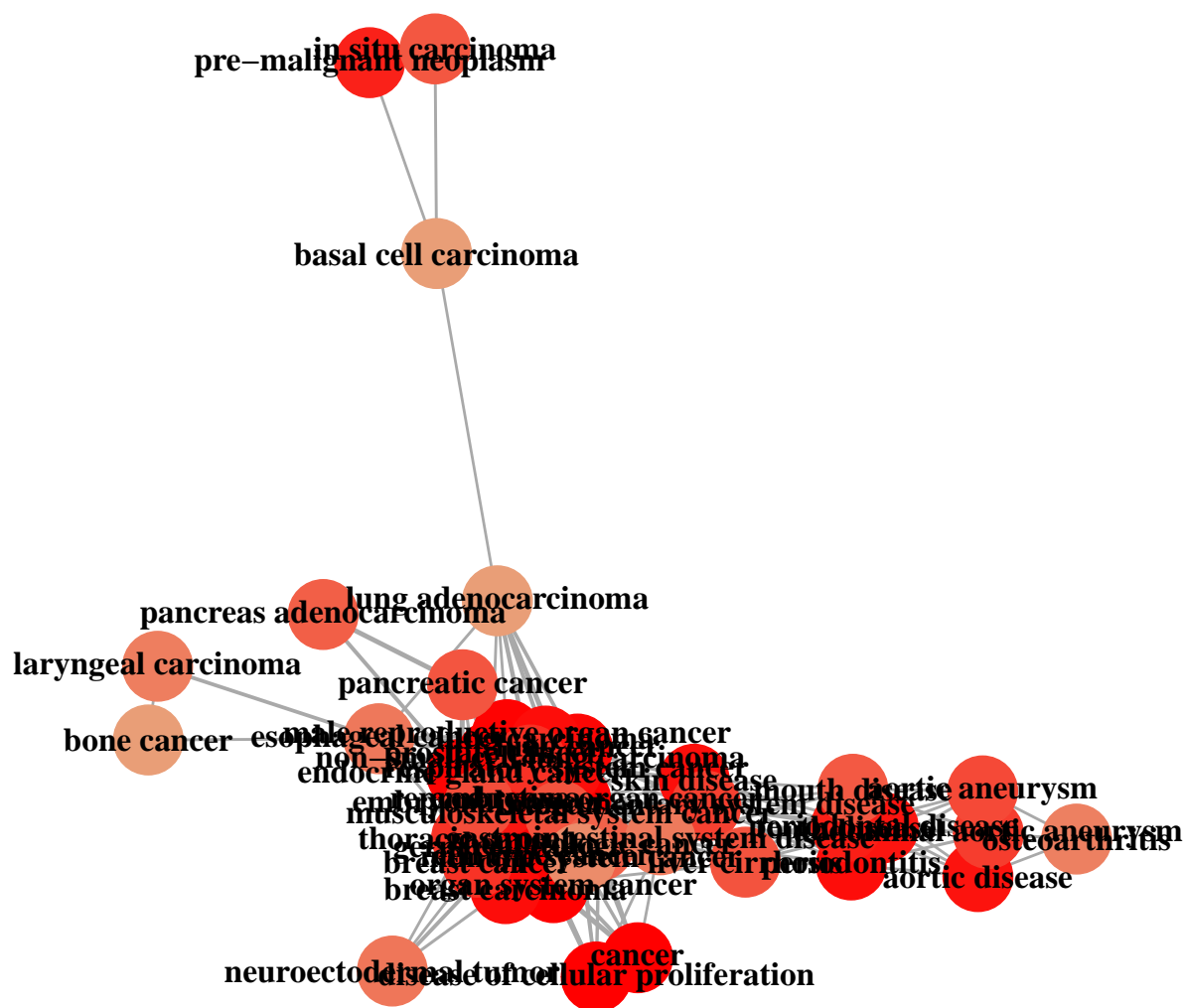


Figure 6: enrichMap of DO enrichment result.

## 7 GO semantic similarity calculation

GO Semantic similarity can be calculated by *GOSemSim* [8].

## 8 Other enrichment analysis tools

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We provides GO & KEGG enrichment analysis in [clusterProfiler](#) [11] and Reactome pathway enrichment analysis in [ReactomePA](#) package. Both hypergeometric test and GSEA are supported.

## 9 External document

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- [Why clusterProfiler fails](#)
- [NCG enrichment implemented in DOSE <- bioinfoblog.it](#)
- [Enrichment map](#)

## 10 Bugs/Feature Requests

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If you have any, [let me know](#).

## 11 Session Information

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Here is the output of `sessionInfo()` on the system on which this document was compiled:

- R version 3.2.2 (2015-08-14), x86\_64-apple-darwin13.4.0
- Locale: C/en\_US.UTF-8/en\_US.UTF-8/C/en\_US.UTF-8/en\_US.UTF-8
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: AnnotationDbi 1.30.1, Biobase 2.28.0, BiocGenerics 0.14.0, DBI 0.3.1, DO.db 2.9, DOSE 2.6.6, GenomInfoDb 1.4.2, IRanges 2.2.7, RSQLite 1.0.0, S4Vectors 0.6.3, clusterProfiler 2.2.5, org.Hs.eg.db 3.1.2
- Loaded via a namespace (and not attached): BiocStyle 1.6.0, Biostrings 2.36.4, GO.db 3.1.2, GOSemSim 1.26.0, KEGGREST 1.8.0, MASS 7.3-43, R6 2.1.1, Rcpp 0.12.0, XVector 0.8.0, colorspace 1.2-6, digest 0.6.8, evaluate 0.7.2, formatR 1.2, ggplot2 1.0.1, grid 3.2.2, gtable 0.1.2, highr 0.5, httr 1.0.0, igraph 1.0.1, knitr 1.11, labeling 0.3, magrittr 1.5, munsell 0.4.2, plyr 1.8.3, png 0.1-7, proto 0.3-10, qvalue 2.0.0, reshape2 1.4.1, scales 0.3.0, splines 3.2.2, stringi 0.5-5, stringr 1.0.0, tools 3.2.2, zlibbioc 1.14.0

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