GeneAnswers, Integrated Interpretation of Genes

Gang Feng[‡], Pan Du[‡], Warren A. Kibbe[‡], Simon Lin[‡] May 10, 2010

[‡]Northwestern University Biomedical Informatics Center Northwestern University, Chicago, IL, 60611, USA

Contents

1	Overview of GeneAnswers	1
2	Citation	2
3	Installation of GeneAnswers package	2
4	Object models of major classes	3
5	Data preprocessing5.1 Build a GeneAnswers instance5.2 Visulization5.3 Multigroup Genes Analysis5.4 Homologous Gene Mapping	3 4 6 26 26
6	Session Info	31
7	Acknowledgments	32
8	References	32

1 Overview of GeneAnswers

Microarray techniques have been widely employed in genomic scale studies for more than one decade. The standard analysis of microarray data is to filter out a group of genes from thousands of probes by certain statistical criteria. These genes are usually called significantly differentially expressed genes. Recently, next generation sequencing (NGS) is gradually adopted to explore gene transcription, methylation, etc. Also a gene list can be obtained by NGS preliminary data analysis. However, this type of information is not enough to understand

^{*}g-feng (at) northwestern.edu

[†]dupan (at) northwestern.edu

[‡]wakibbe (at) northwestern.edu

[§]s-lin2 (at) northwestern.edu

the potential linkage between identified genes and interested functions. The integrated functional and pathway analysis with gene expression data would be very helpful for researchers to interpret the relationship between the identified genes and proposed biological or medical functions and pathways.

The GeneAnswers package provides an integrated solution for a group of genes and specified categories (biological or medical functions, such as Gene Ontology, Disease Ontology, KEGG, etc) to reveal the potential relationship between them by means of statistical methods, and make user-friendly network visualization to interpret the results. Besides the package has a function to combine gene expression profile and category analysis together by outputting concept-gene cross tables, keywords query on NCBI Entrez Gene and application of human based Disease ontology analysis of given genes from other species can help people to understand or discover potential connection between genes and functions.

2 Citation

For the people using *GeneAnswers* package, please cite the following papers in your publications.

* For DOLite:

Du, P., Feng, G., Flatow, J., Song, J., Holko, M., Kibbe, W.A. and Lin, S.M., (2009) 'From disease ontology to disease-ontology lite: statistical methods to adapt a general-purpose ontology for the test of gene-ontology associations', Bioinformatics 25(12):i63-8

* For GeneAnswers:

Feng, G., Du, P., Krett, N.L., Tessel, M., Rosen, S., Kibbe, W.A., and Lin, S.M., (submitted) 'Bioconductor Methods to Visualize Gene-list Annotations', Thanks for your help!

3 Installation of GeneAnswers package

In order to install the *GeneAnswers* package, the user needs to first install R, some related Bioconductor packages. You can easily install them by the following codes.

source("http://bioconductor.org/biocLite.R")
biocLite("GeneAnswers")

For the users want to install the latest developing version of *GeneAnswers*, which can be downloaded from the developing section of Bioconductor website. Some additional packages might be required to be installed because of the update the Bioconductor. These packages can also be found from the developing section of Bioconductor website. You can also directly install the source packages from the Bioconductor website by specify the developing version number, which can be found at the Bioconductor website. Suppose the developing version is 2.5, to install the latest *GeneAnswers* pakcage in the Bioconductor developing version, you can use the following command:

install.packages("GeneAnswers",repos="http://www.bioconductor.org/packages/2.5/bioc",type=

4 Object models of major classes

The *GeneAnswers* package has one major class: **GeneAnswers**. It includes the following slots:

- 1. geneInput: a data frame containing gene Entrez IDs with or without any related values. The values could be foldChange, p value, or other values. These data can be used for concept-gene network. Genes with positive values will be represented as red nodes, while negative value genes are green nodes.
- 2. testType: statistical test method. Current version supports hypergeometric test to test relationship between genes and specified categories.
- 3. *pvalueT*: the cutoff of statistical test. Any categories will not be reported if the p value is more than the cutoff.
- 4. *genesInCategory*: a list containing genes belonging to categories. The names of the list are categories.
- 5. geneExpProfile: a data frame to store gene expression data. If not available, it could be NULL.
 - 6. annLib: annotation database used for statistical test.
 - 7. category Type: functional or medical category used for statistical test.
- 8. enrichmentInfo: a data frame containing filtered categories with statistical results by specified pvalueT.

The figure, 'Flow chart of GeneAnswers', shows how GeneAnswers package works. A group of genes are essential. We use unique Entrez gene IDs to represent genes. Any relative feature values of these genes can also be optional input information, like fold changes, p values, etc. If the gene expression profile of these genes are available, it can be considered as input, too. Since we want to find the potential connections between genes and categories, category type is also need to be specified. GeneAnswers currently supports Gene Ontology (GO), Pathway (KEGG) and developing Disease Ontology (DOLite) in our team. Furthermore, GeneAnswers supports Entrez eUtilis so that users can make customized annotation library based on interested keywords. If users have own annotation library, GeneAnswers can use it to build relationship between it and given genes.

Besides usual barplot and pie chart of top categories, *GeneAnswers* also provides four types of visualization. One is concepts-genes network, which show the concepts and genes on a network layout. The second one is concepts-genes cross table that integrated gene expression profile and corresponding categories together. The third one is a concepts-network shows connections between categories only. The last one is a table, which contains all of information of categories and genes. Combining all of these presentations can be helpful to find and explain the possible linkages between genes and categories.

5 Data preprocessing

First of all, load the GeneAnswers package.

> library(GeneAnswers)

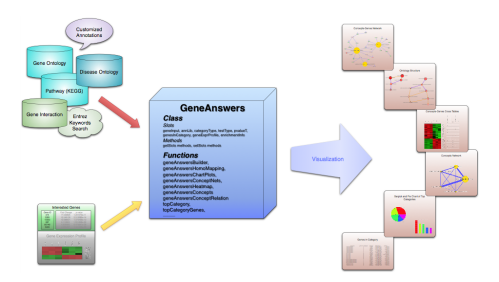


Figure 1: Flow chart of GeneAnswers

5.1 Build a GeneAnswers instance

The key point of GeneAnswers package is to build a GeneAnswers instance. The essential input for GeneAnswers is an Entrez gene IDs vector (a character vector). However, if users have any interested values associated with genes, these values can also be as optional inputs. In this case, the input, geneInput, could be a matrix or a dataframe. The first column is always for Entrez gene IDs. Other columns are used to store those interested values. Rownames for the matrix or dataframe are not necessary, but colnames are recommended for further usage. We use two internal datasets, one is from human and another is from mouse, as examples to show how to implement GeneAnswers package. The human and mouse datasets coming with the GeneAnswers package are from human and mouse Illumina beadarray experiments. Each dataset contains two dataframes. For example, humanGeneInput is a dataframe containing Entrez gene IDs with fold changes and p values, while the data frame, humanExpr, includes two types, control and treatment, of gene expression profile of the genes in humanGeneInput.

- > data('humanGeneInput')
- > data('humanExpr')
- > ## build a GeneAnswers instance with statistical test based on biological process of GO
 > x <- geneAnswersBuilder(humanGeneInput, 'org.Hs.eg.db', categoryType='GO.BP', testType='</pre>
- [1] "geneInput has built in ..."
- [1] "annLib and categoryType have built in ..."
- [1] "genesInCategory has built in ..."
- [1] "Enrichment test is only performed based on annotated genes"
- [1] "testType, pvalueT and enrichmentInfo have built in ..."
- [1] "geneExpressionProfile has been built in ..."
- [1] "GeneAnswers instance has been successfully generated!"
- > class(x)

```
[1] "GeneAnswers"
attr(,"package")
[1] "GeneAnswers"

> ## build a GeneAnswers instance with statistical test based on KEGG and saved example da
> y <- geneAnswersBuilder(humanGeneInput, 'org.Hs.eg.db', categoryType='KEGG', testType='h

[1] "GeneAnswers instance has been successfully generated!"

> ## build a GeneAnswers instance with statistical test based on DOLite and saved example
> z <- geneAnswersBuilder(humanGeneInput, 'org.Hs.eg.db', categoryType='DOLITE', testType=
[1] "GeneAnswers instance has been successfully generated!"

> w <- geneAnswersBuilder(humanGeneInput, 'org.Hs.eg.db', categoryType='GO.BP', testType='
[1] "GeneAnswers instance has been successfully generated!"

We have four GeneAnswers objects, x, y, z and w, containing the statistical</pre>
```

We have four GeneAnswers objects, x, y, z and w, containing the statistical test of biological process of GO, KEGG, DOLite and GO (The first two level nodes are removed), respectively. For Gene Ontology, sometimes, users think some nodes are too general and not very relative to their interests. So we provide parameter *level* to determine how many top levels of GO nodes are removed. The instances have included the relationship between given genes and specified categories.

GeneAnswers package also provides a function searchEntrez to retrieve Entrez genes for given keywords by Entrez XML query. National Center for Biotechnology Information (NCBI) provides many powerful online query systems. One of them is Entrez Programming Utilities (eUtils). Users can query NCBI databases by simple keywords logical operations based on XML protocol. This is very helpful to find potential or interested biological functions or pathways. Hence, the retrieved information can be considered as a customized annotation library to test whether the given genes are relative to interested keywords. Here is a case to build a customized GeneAnswers instance.

```
> keywordsList <- list(Apoptosis=c('apoptosis'), CellAdhesion=c('cell adhesion'))
> entrezIDList <- searchEntrez(keywordsList)

[1] "search link: http://eutils.ncbi.nlm.nih.gov/entrez/eutils/esearch.fcgi?db=gene&term=a
[1] "search link: http://eutils.ncbi.nlm.nih.gov/entrez/eutils/esearch.fcgi?db=gene&term=%
> q <- geneAnswersBuilder(humanGeneInput, entrezIDList, testType='hyperG', totalGeneNumber
[1] "GeneAnswers instance has been successfully generated!"
> class(q)

[1] "GeneAnswers"
attr(,"package")
[1] "GeneAnswers"
> getAnnLib(q)
```

NULL

```
> getCategoryType(q)
```

[1] "User defined"

Customized GeneAnswers instances have NULL at annLib slot and "User defiend" in categoryType slot.

5.2 Visulization

Besides barplot and pie chart, GeneAnswers package can generate a network (concept-gene network) show how genes are connected to specified categories as well as general barplot and piechart. Function GeneAnswersConceptNet can generate a common R canvas or tcl/tk interactive canvas to draw the network by calling igraph. Genes are presented as red nodes, if specified values are positive, and the gene nodes are green with negative values. The category nodes are yellow nodes, the sizes are relative to user-specified values. Currently, if function GeneAnswersBuilder successfully returns a GeneAnswers instance, the genes are represented as entrez IDs and categories are also category IDs. User can map them to gene symbols and categories terms by function GeneAnswersReadable. Function GeneAnswersReadable reads slot annLib to map Entrez IDs to gene symbols, so make sure slot annLib is correct before mapping.

```
> ## mapping gene IDs and category IDs to gene symbols and category terms
> xx <- geneAnswersReadable(x)

[1] "Mapping geneInput ..."
[1] "Mapping genesInCategory ..."
[1] "Mapping enrichmentInfo rownames ..."

[1] "Mapping geneExprProfile rownames ..."

> yy <- geneAnswersReadable(y, verbose=FALSE)
> zz <- geneAnswersReadable(z, verbose=FALSE)
> ww <- geneAnswersReadable(w, verbose=FALSE)
> q <- setAnnLib(q, 'org.Hs.eg.db')
> qq <- geneAnswersReadable(q, catTerm=FALSE)

[1] "Mapping geneInput ..."
[1] "Mapping genesInCategory ..."
[1] "Mapping geneExprProfile rownames ..."</pre>
```

Since function *geneAnswersReadable* implements mapping based on annotation database in slot annLib, we assign 'org.Hs.eg.db' to customized GeneAnswers instance annLib slot at first for make it readable.

```
> ## plot barplot and / or piechart
> geneAnswersChartPlots(xx, chartType='all')
```

The nodes could be represented by different colors for different specified values, like fold change. In this case, overexpressed genes are represented as red nodes while green dots stand for underexpressed genes. If the node is more red, which means its fold change is larger. It's same for green nodes, but different direction.

0.0.0 Quartz3

Top 5 Categories Distribution based on genes in Category

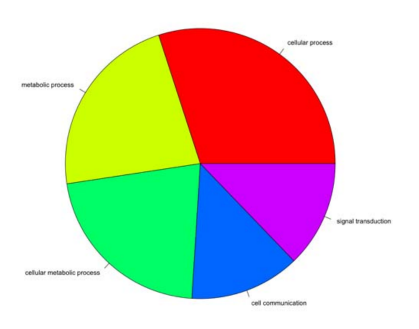


Figure 2: Screen shot of pie chart of top categories

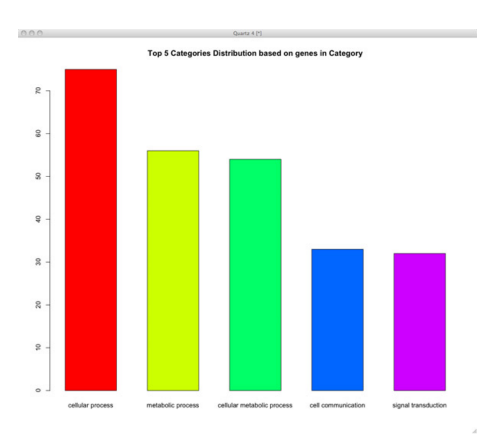


Figure 3: Screen shot of barplot of top categories

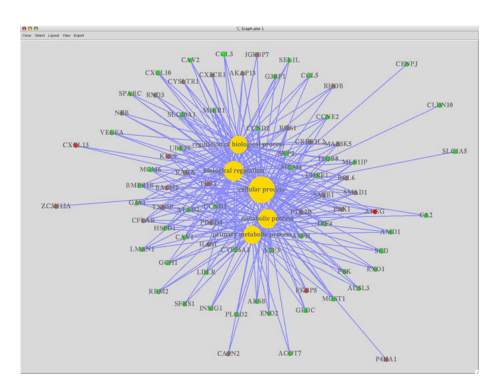


Figure 4: Screen shot of concept-gene network

- > ## plot interactive concept-gene network
- > geneAnswersConceptNet(xx, colorValueColumn='foldChange', centroidSize='pvalue', output='

The top 5 categories might not be very specific. Users might get a tree view to see relative category terms by calling function geneAnswersConceptRelationif the category has an ontology structure. The size of nodes is proportional to number of genes in these GO categories. The color of nodes stand for how relative the given genes are to the GO categories. More red, more relative. The given GO categories are yellow framed dots with dark purple edges connections.

- > ## plot interactive go structure network
- > geneAnswersConceptRelation(x, direction='both', netMode='connection', catTerm=TRUE, catI

Also the new version GeneAnswers integrates gene or protein interaction database from NCBI. The following case is a typical one to show interaction information could be included in basic concepts-genes network.

- > ## plot interactive concept-gene network
- > geneAnswersConceptNet(x, color='foldChange', geneLayer=5, output='interactive', showCats

This function can also be used to show how the given genes interact with each other. For example, for the given genes in the GeneAnswers instance, x, users can use the following command to show gene '444','3638', '5087' and '55835' interact with each other and other genes in the given geneInput of x.

- > ## plot the given gene interaction
- > buildNet(c('444','3638', '5087','55835'), idType='GeneInteraction', layers=2, filterGrap

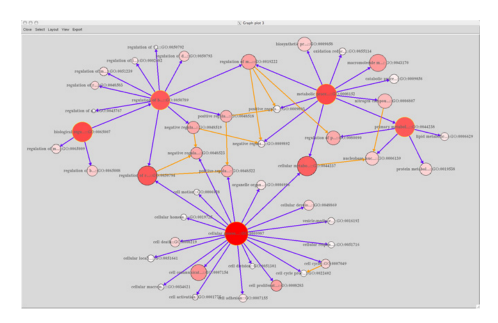


Figure 5: Screen shot of go structure network

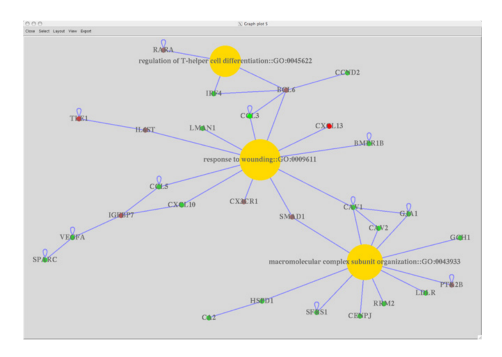


Figure 6: Screen shot of concept-gene network

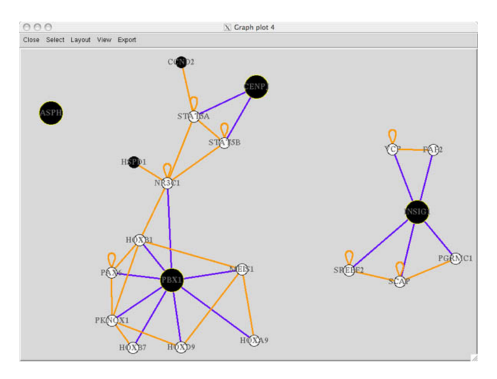


Figure 7: Screen shot of the given gene interaction network

In this case, large black dots with yellow frame stand for the 4 given genes. They also connect to other genes by dark-blue-purple edges. Small black dots represent the other genes from getGeneInput(x). Small white dots are genes that are not in the genes from getGeneInput(x), but interact with these genes.

If there are some certain values associate with genes in geneInput of x, like the example data, you can represent any one column by colors. For example, if users want to show how genes interaction with gene expression information, the following command can show this:

- > ## plot the given gene interaction
- > buildNet(c('444', '3638', '5087', '55835'), idType='GeneInteraction', layers=2, filterGraphy and solution', layers=2, filterGraphy and soluti

The following one is based on expression p-values, we often use - $\log 2$ to transform p-values.

- > ## plot the given gene interaction
- > buildNet(c('444','3638', '5087','55835'), idType='GeneInteraction', layers=2, filterGrap

Users can also define customized color scheme and transfer it into 'buildNet' function by parameter 'colorMap'. Details are available in 'buildNet' manual page. If users want to have a overview for the interaction of all of the given genes from getGeneInput(x), the following command could be used:

- > ## plot the given gene interaction
- > buildNet(getGeneInput(x)[,1], idType='GeneInteraction', layers=2, filterGraphIDs=getGene

If there are a lot of genes, the network could be very complicated. We strongly recommend to use 'interactive' mode to show the network since it's

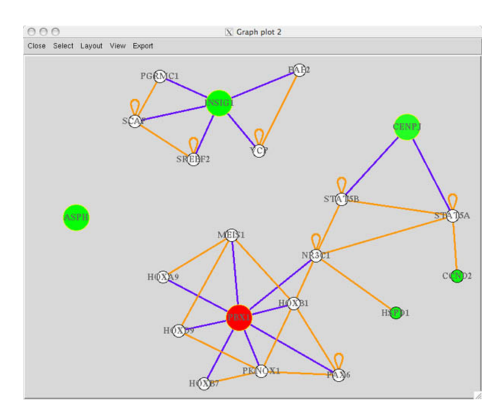


Figure 8: Screen shot of the given gene interaction network with expression information

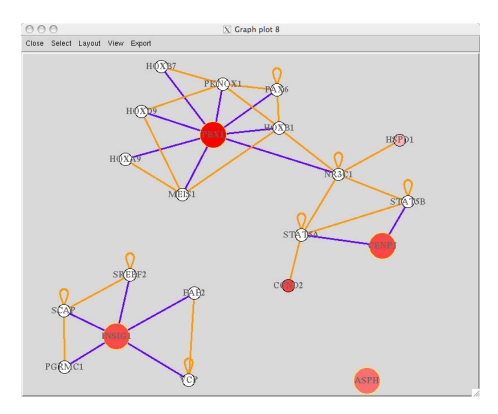


Figure 9: Screen shot of the given gene interaction network with p-value information ${\bf p}$

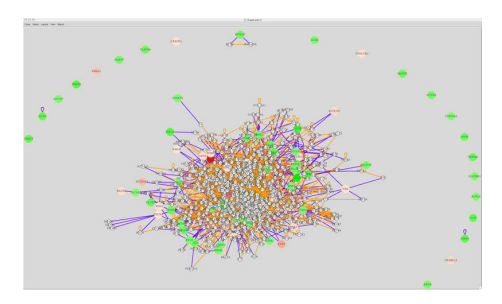


Figure 10: Screen shot of all of the given gene interaction network with expression information ${\bf r}$

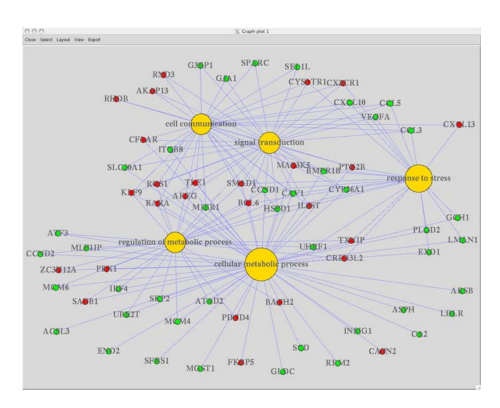


Figure 11: Screen shot of concept-gene network for top 2 GO level nodes removal

easy to manually change layout. The default setting is 'interactive', but users can change it to 'fixed' for a small or simple network.

The following example is to show top 5 GO-gene network for the first 2 level GO nodes removal.

- > ## plot Go-concept network for 2 level nodes removal
- > geneAnswersConceptNet(ww, colorValueColumn='foldChange', centroidSize='pvalue', output='

Also, users can sort enrichment test information and plot it.

- > ## sort enrichmentInfo dataframe by fdr adjusted p value
- > xxx <- geneAnswersSort(xx, sortBy='correctedPvalue')
- > yyy <- geneAnswersSort(yy, sortBy='pvalue')
- > zzz <- geneAnswersSort(zz, sortBy='geneNum')
- > geneAnswersConceptNet(yyy, colorValueColumn='foldChange', centroidSize='geneNum', output
- > geneAnswersConceptNet(zzz, colorValueColumn='foldChange', centroidSize='pvalue', output=

If users provide a gene expression profile, *GeneAnswers* package can generate a table or heatmap labeling relationship between genes and categories with a heatmap of these genes expression. We call this type of representation as concept-gene cross tabulation.

- > ## generate GO-gene cross tabulation
- > geneAnswersHeatmap(x, catTerm=TRUE, geneSymbol=TRUE)

0 0 0 Quartz 2 (*)

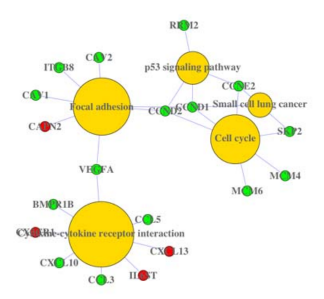


Figure 12: Screen shot of KEGG-gene network

0.0 0 Quart 2 [1]

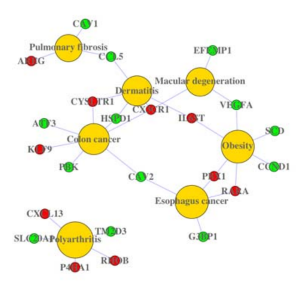


Figure 13: Screen shot of DOLite-gene network

- > ## generate GO-gene cross tabulation
- > geneAnswersHeatmap(x, catTerm=TRUE, geneSymbol=TRUE)

```
initial value 2.095268
       5 value 0.284632
iter
     10 value 0.180610
iter
iter
     15 value 0.104728
iter
     20 value 0.073371
iter 25 value 0.061866
     30 value 0.044067
     35 value 0.025023
iter
iter
     40 value 0.014669
final value 0.004753
converged
initial value 4.654638
iter
       5 value 2.542047
final value 0.000000
converged
```

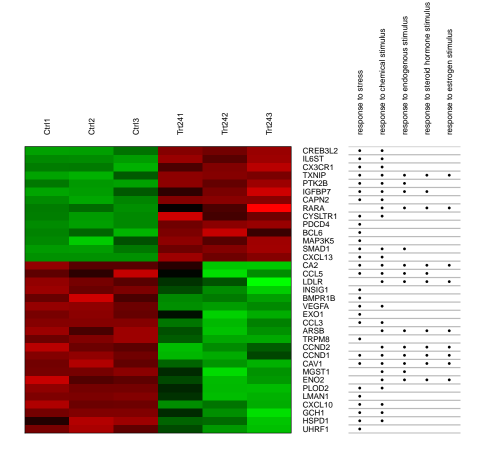


Figure 14: GO-gene cross tabulation

> geneAnswersHeatmap(yyy)

For cross table, there are two types of representations. One is a table, which is better for few genes, and another one is a two-color heatmap that is adopted for a lot of genes. In the latter, the default setting is that white bar stands for that a gene in that category.

Besides top categories, users can also show interested categories.

```
> GOBPIDs <- c("GO:0007049", "GO:0042592", "GO:0006259", "GO:0016265", "GO:0007243")
```

- > GOBPTerms <- c("cell cycle", "death", "protein kinase cascade", "homeostatic process", "
- > ## generate concept-gene cross tabulation
- > geneAnswersConceptNet(x, colorValueColumn='foldChange', centroidSize='pvalue', output='f
- > geneAnswersHeatmap(x, showCats=GOBPIDs, catTerm=TRUE, geneSymbol=TRUE)

Function gene Answers Concepts shows the linkages of specified categories. The width of edge stands for how overlapping between two categories.

- > ## generate concept-gene cross tabulation
- > geneAnswersConcepts(xxx, centroidSize='geneNum', output='fixed', showCats=GOBPTerms)

Users can also print top categories and genes on screen and save them in files by specification as well as these two types of visualization. The default file names are "topCategory.txt" and "topCategoryGenes.txt" for top categories with or without corresponding genes, respectively.

```
> ## print top GO categories sorted by hypergeometric test p value
> topGOGenes(x, orderby='pvalue')
```

```
[1] "****** response to estrogen stimulus p value: 4.89360705325764e-09 *******
   Symbol foldChange
                           pValue
411
     ARSB -2.122207 3.199329e-06
760
      CA2 -2.256972 8.992257e-04
857
     CAV1 -2.122246 3.842276e-06
595 CCND1 -2.257123 5.932920e-07
894 CCND2 -2.239806 1.208734e-06
[1] "****** response to chemical stimulus p value : 5.82089966265016e-09 *******
    Symbol foldChange
                           pValue
411
      ARSB -2.122207 3.199329e-06
760
       CA2 -2.256972 8.992257e-04
824
     CAPN2
            2.568772 1.890638e-08
      CAV1 -2.122246 3.842276e-06
857
      CCL3 -4.031781 1.147014e-07
                                                  p value : 8.4667158655129e-09 ******
[1] "***** response to steroid hormone stimulus
                            pValue
    Symbol foldChange
      ARSB -2.122207 3.199329e-06
411
760
       CA2 -2.256972 8.992257e-04
857
      CAV1 -2.122246 3.842276e-06
6352
     CCL5 -2.154827 1.092361e-03
```

CCND1 -2.257123 5.932920e-07

595

> geneAnswersHeatmap(yyy)

```
initial value 1.716305
      5 value 0.407656
iter
iter
     10 value 0.112360
iter
     15 value 0.072109
     20 value 0.052017
iter 25 value 0.029419
final value 0.006019
converged
initial value 4.466884
      5 value 1.279433
iter
iter 10 value 0.084821
final value 0.008157
converged
```

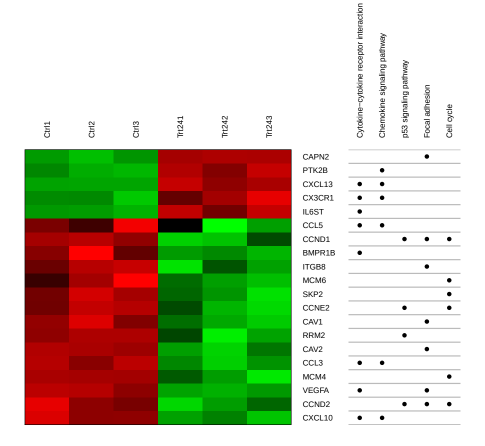


Figure 15: KEGG-gene cross tabulation

```
initial value 1.851350
iter
       5 value 0.390287
     10 value 0.146730
iter
     15 value 0.120260
iter
     20 value 0.092560
     25 value 0.046348
iter
     30 value 0.035369
iter
     35 value 0.028131
iter
iter
     40 value 0.021161
iter 45 value 0.011081
iter 45 value 0.008695
iter 45 value 0.005071
final value 0.005071
converged
initial value 19.020185
      5 value 2.491750
iter
iter
     10 value 0.333367
iter 15 value 0.027846
final value 0.002865
converged
```

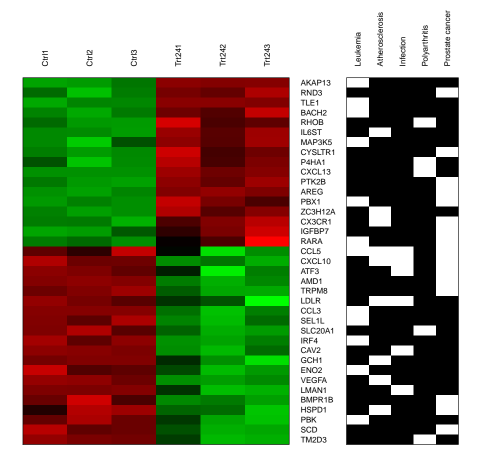


Figure 16: DOLite-gene cross tabulation

```
[1] "Some specified categories might not be statistical significant! Only show significant
initial value 1.955644
iter
       5 value 0.250440
     10 value 0.129463
iter
     15 value 0.075394
iter
      20 value 0.040824
iter
     25 value 0.016440
     30 value 0.013925
iter
final value 0.006865
converged
initial value 0.000000
final value 0.000000
```

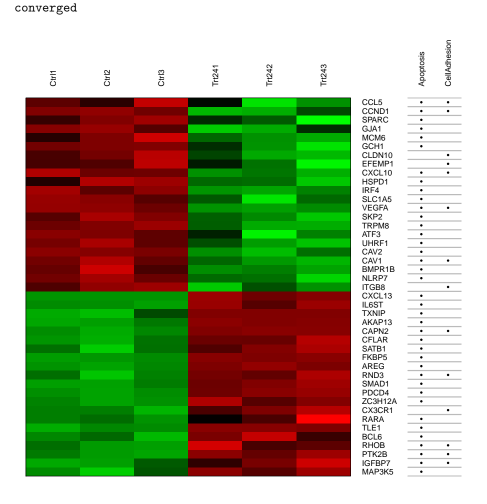


Figure 17: Customized GO-gene cross tabulation

O O O Quartz 2 [*]

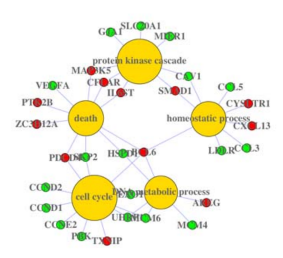


Figure 18: Screen shot of customized GO-gene network

[1] "***** response to stress

```
pValue
   Symbol foldChange
604
     BCL6
            2.380046 2.436277e-04
           -2.161504 1.308048e-04
658 BMPR1B
760
      CA2 -2.256972 8.992257e-04
824 CAPN2
           2.568772 1.890638e-08
     CAV1 -2.122246 3.842276e-06
[1] "***** response to endogenous stimulus  p value : 6.56782483074139e-08 *******
    Symbol foldChange
                            pValue
      ARSB -2.122207 3.199329e-06
411
760
       CA2 -2.256972 8.992257e-04
857
      CAV1
            -2.122246 3.842276e-06
6352
      CCL5 -2.154827 1.092361e-03
595
     CCND1 -2.257123 5.932920e-07
> ## print top KEGG categories sorted by gene numbers and sort genes by fold changes
> topPATHGenes(y, orderby='geneNum', top=4, topGenes=8, genesOrderBy='foldChange')
[1] "****** Cytokine-cytokine receptor interaction
                                                      genes in Category: 8 *******
      Symbol foldChange
                             pValue
6348
       CCL3 -4.031781 1.147014e-07
7422
      VEGFA -2.391115 7.147830e-09
     CXCL10 -2.376811 1.315856e-07
3627
658
     BMPR1B -2.161504 1.308048e-04
```

p value : 4.9255559399943e-08 *******

```
initial value 2.019969
       5 value 0.291962
iter
     10 value 0.188187
iter
iter
     15 value 0.144754
iter
     20 value 0.082005
iter
     25 value 0.065685
     30 value 0.023664
iter
     35 value 0.022465
iter
final value 0.006909
converged
initial value 9.435938
       5 value 2.999357
iter
iter
     10 value 0.139216
     15 value 0.011432
iter
iter
     15 value 0.002732
iter 15 value 0.002297
final value 0.002297
converged
```

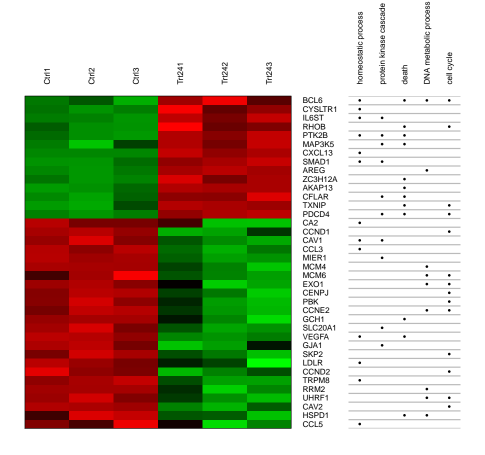


Figure 19: Customized concept-gene cross tabulation

0 0 0 Quartz 2 [*]

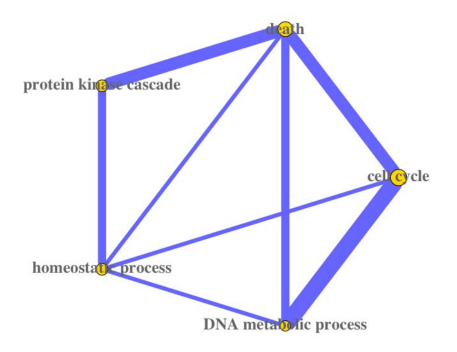


Figure 20: Screen shot of customized GO category linkage

```
1524 CX3CR1 2.127494 4.412558e-06
     IL6ST 2.213384 8.018338e-08
3572
10563 CXCL13 10.688601 1.073456e-10
[1] "***** Focal adhesion genes in Category: 7 *******
    Symbol foldChange pValue
3696 ITGB8 -3.097913 1.178489e-06
      CAV2 -2.643498 2.625770e-08
7422 VEGFA -2.391115 7.147830e-09
     CCND1 -2.257123 5.932920e-07
595
894
     CCND2 -2.239806 1.208734e-06
857
     CAV1 -2.122246 3.842276e-06
824
     CAPN2 2.568772 1.890638e-08
[1] "****** Cell cycle
                         genes in Category: 6 *******
    Symbol foldChange
                         pValue
9134 CCNE2 -2.379943 3.326011e-06
     MCM6 -2.356668 2.080808e-04
4175
     SKP2 -2.276824 1.013445e-05
6502
595 CCND1 -2.257123 5.932920e-07
894 CCND2 -2.239806 1.208734e-06
4173 MCM4 -2.082429 8.413734e-06
[1] "***** Chemokine signaling pathway
                                        genes in Category : 6 *******
     Symbol foldChange
                            pValue
6348
      CCL3 -4.031781 1.147014e-07
3627 CXCL10 -2.376811 1.315856e-07
     CCL5 -2.154827 1.092361e-03
6352
2185
      PTK2B
             2.036789 1.970066e-07
1524 CX3CR1
            2.127494 4.412558e-06
10563 CXCL13 10.688601 1.073456e-10
> ## print and save top 10 DOLites information
> topDOLiteGenes(z, orderby='pvalue', top=5, topGenes='ALL', genesOrderBy='pValue', file=T
[1] "****** Hyperlipidemia p value : 0.00198151640638039 *******
     Symbol foldChange
                           pValue
10628 TXNIP 3.899457 4.296784e-09
6319
        SCD -2.480571 6.976419e-07
2181 ACSL3 -2.049592 5.282896e-04
      CCL5 -2.154827 1.092361e-03
6352
[1] "****** Prostate cancer p value : 0.00396329360969883 *******
                         pValue
    Symbol foldChange
374
      AREG 17.553825 1.241073e-11
262
      AMD1 -2.538954 1.216657e-09
3490 IGFBP7 3.787547 2.688423e-08
            2.552036 1.120693e-07
5087
      PBX1
3627 CXCL10 -2.376811 1.315856e-07
[1] "****** Alveolar bone loss p value : 0.00872917145369052 *******
                           pValue
    Symbol foldChange
6348 CCL3 -4.031781 1.147014e-07
6678 SPARC -2.289622 7.751533e-04
```

6352

CCL5 -2.154827 1.092361e-03

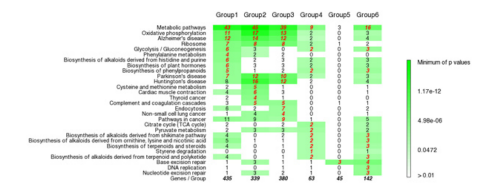


Figure 21: Screen shot of multigroup genes KEGG analysis

```
p value : 0.0114791957044022 *******
[1] "***** Leukodystrophy NOS
    Symbol foldChange
                            pValue
      ARSB -2.122207 3.199329e-06
411
3329 HSPD1 -2.144728 7.145728e-05
[1] "****** Bronchiolitis p value : 0.0114791957044022 *******
                            pValue
    Symbol foldChange
           -4.031781 1.147014e-07
6348
      CCL3
      CCL5 -2.154827 1.092361e-03
6352
[1] "File topCategoryGenes.txt is successfully generated!"
```

5.3 Multigroup Genes Analysis

Several groups of genes are used to dynamically study biological processes by different treatment or time points. In this case, *GeneAnswers* provide a solution to integrate enrichment test information of different groups of genes.

- > ##load multigroup genes sample data
- > data(sampleGroupsData)
- > ##Build a GeneAnswers List
- > gAKEGGL <- lapply(sampleGroupsData, geneAnswersBuilder, 'org.Hs.eg.db', categoryType='KE
- [1] "GeneAnswers instance has been successfully generated!"
- > ##Output integrated text table
- > output<- getConceptTable(gAKEGGL, items='geneNum')

Function groupReport can generate a html file including all information.

5.4 Homologous Gene Mapping

Since DOLite is developed for human, any gene from other species can not take advantage of this novel annotation database. Therefore, *GeneAnswers* package

provides two functions for this type of data interpretation. getHomoGeneIDs can map other species gene Entrez IDs to human homologous gene Entrez IDs at first. Then users can perform normal GeneAnswers functions. Finally, function geneAnswersHomoMapping maps back to original species gene Entrez IDs. Current version supports two types of homologous gene mapping. One is called "direct", which is simple and only works between mouse and human. Since all of human gene symbols are capitalized, while only first letter of mouse homologous gene symbols is uppercase, this method simply maps homologous genes by capitalized moues gene symbols. Another method adopts biomaRt to do mapping. biomaRt contacts its online server to mapping homologous genes. Its database include more accurate information, but it might take longer to do that, while 'direct' method can rapidly do conversation though it is possible to miss some information.

```
## load mouse example data
> data('mouseExpr')
> data('mouseGeneInput')
> mouseExpr[1:10,]
   GeneID
                S11
                          S12
                                    S13
                                             S21
                                                      S22
                                                               S23
1
   93695 11.140745 11.555394 11.199022 13.53989 13.68489 13.52166
   20750 10.378364 10.780340 10.280152 12.51370 12.77777 12.72755
3
   16854 10.576541 10.823445 10.539105 12.52568 12.94808 12.75282
4
   20210 10.417790 10.503403 10.603501 12.38010 12.64376 12.45370
5
   14282 9.392208 9.574147 9.456061 11.47399 11.24749 11.42666
   17105 10.599174 11.078450 10.565310 12.47790 12.79757 12.50897
7
   17110 12.674773 13.153840 12.672851 14.56094 14.89131 14.57835
8
   16002 11.766943 12.268368 11.557304 13.42105 13.62164 13.60838
   21924 8.874513 9.096380 8.860733 10.46360 10.66965 10.62615
10 269994 10.913894 10.330857 10.853911 9.07294 9.07630 9.04366
```

> mouseGeneInput[1:10,]

```
Symbol foldChange
                                pValue
93695
        93695
                4.869452 1.864011e-08
        20750
                4.573777 1.224957e-07
20750
16854
        16854
                4.274721 6.526113e-08
20210
        20210
                3.956676 4.098411e-09
14282
        14282
                3.754383 3.190981e-09
17105
        17105
                3.597932 1.088294e-06
        17110
                3.587662 1.035619e-06
17110
16002
        16002
                3.217968 5.465650e-06
21924
        21924
                3.122260 2.337725e-08
269994 269994
               -3.106423 1.962161e-06
```

- > ## only keep first one for one to more mapping
- > pickHomo <- function(element, inputV) {return(names(inputV[inputV == element])[1])}
- > ## mapping geneInput to homo entrez IDs.
- > homoLL <- getHomoGeneIDs(mouseGeneInput[,1], species='mouse', speciesL='human', mappingM
- [1] "Warning: homogenes of some input genes can not be found and are removed!!!"

```
> newGeneInput <- mouseGeneInput[mouseGeneInput[,1] %in% unlist(lapply(unique(homoLL), pic
> dim(mouseGeneInput)
[1] 71 3
> dim(newGeneInput)
[1] 67 3
> newGeneInput[,1] <- homoLL[newGeneInput[,1]]</pre>
> ## mapping geneExpr to homo entrez IDs.
> homoLLExpr <- getHomoGeneIDs(as.character(mouseExpr[,1]), species='mouse', speciesL='hum</pre>
[1] "Warning: homogenes of some input genes can not be found and are removed!!!"
> newExpr <- mouseExpr[as.character(mouseExpr[,1]) %in% unlist(lapply(unique(homoLLExpr) ,
> newExpr[,1] <- homoLLExpr[as.character(newExpr[,1])]</pre>
> dim(mouseExpr)
[1] 71 7
> dim(newExpr)
[1] 67 7
> ## build a GeneAnswers instance based on mapped data
> v <- geneAnswersBuilder(newGeneInput, 'org.Hs.eg.db', categoryType='DOLITE', testType='h
[1] "geneInput has built in ..."
[1] "annLib and categoryType have built in ..."
[1] "genesInCategory has built in ..."
[1] "Enrichment test is only performed based on annotated genes"
[1] "testType, pvalueT and enrichmentInfo have built in ..."
[1] "geneExpressionProfile has been built in ..."
[1] "GeneAnswers instance has been successfully generated!"
> ## make the GeneAnswers instance readable, only map DOLite IDs to terms
> vv <- geneAnswersReadable(v, geneSymbol=F)
[1] "Mapping genesInCategory ..."
[1] "Mapping enrichmentInfo rownames ..."
> getAnnLib(vv)
[1] "org.Hs.eg.db"
> ## mapping back to mouse genes
> uu <- geneAnswersHomoMapping(vv, species='human', speciesL='mouse', mappingMethod='direc</p>
[1] "Change annLib ..."
[1] "Mapping geneInput ..."
[1] "Mapping genesInCategory ..."
```

[1] "Mapping geneExprProfile ..."

1 0 0 Quartz 2 (*)

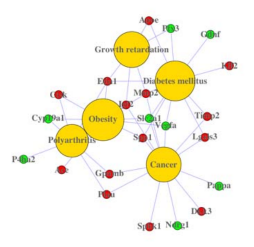


Figure 22: Screen shot of homogene DOLite-gene network

```
> getAnnLib(uu)
[1] "org.Mm.eg.db"
> ## make mapped genes readable, DOLite terms are not mapped
> u <- geneAnswersReadable(uu, catTerm=FALSE)
[1] "Mapping geneInput ..."
[1] "Mapping genesInCategory ..."
[1] "Mapping geneExprProfile rownames ..."
> ## sort new GeneAnswers instance
> u1 <- geneAnswersSort(u, sortBy='pvalue')
> ## plot concept-gene network
> geneAnswersConceptNet(u, colorValueColumn='foldChange', centroidSize='pvalue', output='f
> ## plot homogene DOLite-gene cross tabulation
> geneAnswersHeatmap(u1)
> ## output top information
> topDOLiteGenes(u, geneSymbol=FALSE, catTerm=FALSE, orderby='pvalue', top=6, topGenes='AL
[1] "****** Growth retardation p value : 1.86018082726357e-06 *******
```

pValue

Symbol foldChange

initial value 0.561792
iter 5 value 0.185166
iter 10 value 0.055120
iter 15 value 0.051535
iter 20 value 0.035624
final value 0.009132
converged
initial value 16.218142
iter 5 value 9.824398
final value 9.383898
converged

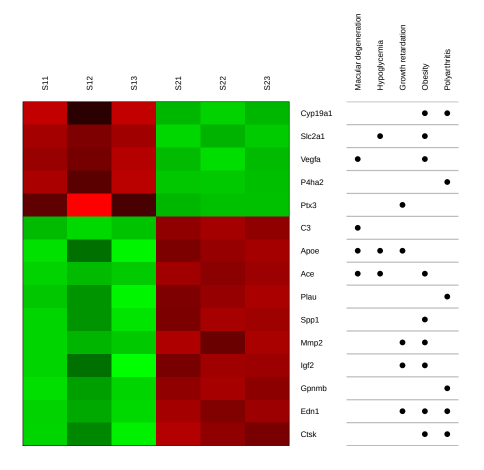


Figure 23: homogene DOLite-gene cross tabulation

```
Edn1
              2.504625 1.536084e-08
13614
17390
       Mmp2
              2.932919 5.771626e-08
11816
       Apoe
              2.465045 4.135459e-06
16002
       Igf2
              3.217968 5.465650e-06
       Ptx3 -2.100947 4.364358e-05
19288
                       p value : 5.28156537991024e-06 *******
[1] "****** Obesity
     Symbol foldChange
                             pValue
              2.897884 4.811431e-10
11421
        Ace
20525 Slc2a1 -2.590159 6.290580e-09
       Edn1
              2.504625 1.536084e-08
13614
22339
      Vegfa -2.535157 3.651889e-08
17390
       Mmp2
              2.932919 5.771626e-08
[1] "***** Hypoglycemia
                            p value : 9.69236642456048e-06 *******
     Symbol foldChange
                             pValue
        Ace
              2.897884 4.811431e-10
11421
20525 Slc2a1 -2.590159 6.290580e-09
             2.465045 4.135459e-06
11816
      Apoe
[1] "****** Polyarthritis p value : 2.85339797023958e-05 *******
     Symbol foldChange
                             pValue
13614
       Edn1
              2.504625 1.536084e-08
93695 Gpnmb
              4.869452 1.864011e-08
18452 P4ha2 -2.584996 3.363470e-07
             2.456354 4.029683e-07
18792
       Plau
13038
       Ctsk
             2.220556 1.394015e-06
[1] "****** Macular degeneration
                                    p value : 3.4593344673306e-05 *******
     Symbol foldChange
                             pValue
              2.897884 4.811431e-10
11421
        Ace
12266
         СЗ
              2.580510 1.185513e-09
22339 Vegfa -2.535157 3.651889e-08
11816
       Apoe
              2.465045 4.135459e-06
                                   p value : 5.68729958212342e-05 *******
[1] "***** Capillaries disease
     Symbol foldChange
                             pValue
              2.504625 1.536084e-08
13614
       Edn1
22339 Vegfa -2.535157 3.651889e-08
17390
       Mmp2
              2.932919 5.771626e-08
[1] "File topCategoryGenes.txt is successfully generated!"
```

6 Session Info

- > toLatex(sessionInfo())
 - R version 2.11.0 (2010-04-22), i386-apple-darwin9.8.0
 - Locale: en_US.UTF-8/en_US.UTF-8/C/C/en_US.UTF-8/en_US.UTF-8
 - Base packages: base, datasets, graphics, grDevices, methods, stats, tools, utils
 - Other packages: annotate 1.26.0, AnnotationDbi 1.9.8, Biobase 2.8.0, bitops 1.0-4.1, DBI 0.2-5, GeneAnswers 1.5.3, GO.db 2.4.1,

Heatplus 1.17.0, igraph 0.5.3, KEGG.db 2.4.1, MASS 7.3-5, org.Hs.eg.db 2.4.1, org.Mm.eg.db 2.4.1, RColorBrewer 1.0-2, RCurl 1.3-1, rgl 0.90, RSQLite 0.8-4, XML 2.8-1

• Loaded via a namespace (and not attached): xtable 1.5-6

7 Acknowledgments

We would like to thank the users and researchers around the world contribute to the *GeneAnswers* package, provide great comments and suggestions and report bugs

8 References

Du, P., Feng, G., Flatow, J., Song, J., Holko, M., Kibbe, W.A. and Lin, S.M., (2009) 'From disease ontology to disease-ontology lite: statistical methods to adapt a general-purpose ontology for the test of gene-ontology associations', Bioinformatics 25(12):i63-8

Feng, G., Du, P., Krett, N.L., Tessel, M., Rosen, S., Kibbe, W.A., and Lin, S.M., (submitted) 'Bioconductor Methods to Visualize Gene-list Annotations',