

# Biological Theme Comparison

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## 1 Introduction

In recently years, high-throughput experimental techniques such as microarray, RNA-Seq and mass spectrometry can detect cellular moleculars at systems-level. These kinds of analysis generate huge quantities of data, which need to be given a biological interpretation. A commonly used approach is via clustering in the gene dimension for grouping different genes based on their similarities(Yu et al., 2010).

To search for shared functions among genes, a common way is to incorporate the biological knowledge, such as Gene Ontology (GO) and Kyoto Encyclopedia of genes and Genomes (KEGG), for identifying predominant biological themes of a collection of genes.

After clustering analysis, researchers not only want to determine whether there is a common theme of a particular gene cluster, but also to compare the biological themes among gene clusters. The manual step to choose interesting clusters followed by enrichment analysis on each selected cluster is slow and tedious. To bridge this gap, we designed *clusterProfiler*, for comparing and visulizing functional profiles among gene clusters.

## 2 Citation

Please cite the following articles when using *clusterProfiler*.

G Yu, LG Wang, Y Han, QY He. *clusterProfiler*: an R package for comparing biological themes among gene clusters. *OMICS: A Journal of Integrative Biology*. 2012, 16(5), in press.

## 3 Functional Profiles

In *clusterProfiler*, we implemented three functions to explore the functional profiles of a collection of genes.

- `groupGO` for gene classification based on GO distribution at a specific level

```
> data(gcSample)
> x <- groupGO(gene=gcSample[[1]],
+             organism="human",
+             ont="CC",
+             level=2,
+             readable=TRUE)
> head(summary(x))
```

ID	Description	Count
GO:0005576	GO:0005576	1
GO:0005623	GO:0005623	13
GO:0019012	GO:0019012	0
GO:0030054	GO:0030054	1
GO:0031974	GO:0031974	7
GO:0032991	GO:0032991	6

GO:0005576  
GO:0005623  
GO:0019012  
GO:0030054  
GO:0031974  
GO:0032991

SDF2L1/PA2G4/RAD50/RUVBL2/LONP1/RPL4/  
PA2G4/RAD50/RUVBL2/RPS23/RPL4/

• **enrichGO for GO enrichment analysis**

```
> y <- enrichGO(gene=gcSample[[2]],
+               organism="human",
+               ont="MF",
+               pvalueCutoff=0.01,
+               qvalueCutoff=0.05,
+               readable=TRUE)
> head(summary(y))
```

ID	Description	GeneRatio	BgRatio
GO:0003924	GO:0003924	4/18	230/17870
GO:0008135	GO:0008135	3/18	87/17870
GO:0000166	GO:0000166	9/18	2270/17870
GO:0003746	GO:0003746	2/18	20/17870
GO:0005525	GO:0005525	4/18	375/17870
GO:0019001	GO:0019001	4/18	388/17870

	pvalue	qvalue
GO:0003924	7.099129e-05	0.002724231
GO:0008135	8.626731e-05	0.002724231
GO:0000166	1.398773e-04	0.002844141
GO:0003746	1.801289e-04	0.002844141
GO:0005525	4.625035e-04	0.004745854
GO:0019001	5.259988e-04	0.004745854

GO ID	geneID
GO:0003924	EEF1A2/EEF2/RAB5A/EFTUD2
GO:0008135	EEF1A2/EEF2/EIF4A1
GO:0000166	CCT2/EEF1A2/EEF2/EIF4A1/NDUFA10/RAB5A/SNRPB2/SPR/EFTUD2
GO:0003746	EEF1A2/EEF2
GO:0005525	EEF1A2/EEF2/RAB5A/EFTUD2
GO:0019001	EEF1A2/EEF2/RAB5A/EFTUD2

GO ID	Count
GO:0003924	4
GO:0008135	3
GO:0000166	9
GO:0003746	2

```
GO:0005525      4
GO:0019001      4
```

- `enrichKEGG` for KEGG pathway enrichment analysis.

```
> z <- enrichKEGG(gene=gcSample[[3]],
+                 organism="human",
+                 pvalueCutoff=0.05,
+                 qvalueCutoff=0.05,
+                 readable=TRUE)
> head(summary(z))
```

ID	Description
hsa05130	Pathogenic Escherichia coli infection
hsa04145	Phagosome
hsa04540	Gap junction
hsa04962	Vasopressin-regulated water reabsorption

GeneRatio	BgRatio	pvalue	qvalue
hsa05130	4/17	58/5894	1.826892e-05
hsa04145	5/17	156/5894	5.827611e-05
hsa04540	4/17	90/5894	1.039489e-04
hsa04962	2/17	44/5894	6.898981e-03

geneID	Count	
hsa05130	TUBB2C/TUBB2A/TUBB3/TUBB6	4
hsa04145	TUBB2C/TUBB2A/TUBB3/RAB5B/TUBB6	5
hsa04540	TUBB2C/TUBB2A/TUBB3/TUBB6	4
hsa04962	NSF/RAB5B	2

With the demise of KEGG (at least without subscription), the pathway data used in *clusterProfiler* will not update, and we encourage user to use `enrichPathway` in Bioconductor package *ReactomePA*, which use Reactome as a source of pathway data.

The function calls of `groupGO`, `enrichGO` and `enrichKEGG` are similar. The input parameters of *gene* is a vector of entrezgene (for human and mouse) or ORF (for yeast) IDs, and *organism* must be one of "human", "mouse", and "yeast", according to the gene IDs.

For GO analysis, *ont* must be assigned to one of "BP", "MF", and "CC" for biological process, molecular function and cellular component, respectively. In `groupGO`, the *level* specify the GO level for gene projection.

In enrichment analysis, the *pvalueCutoff* is to restrict the result based on their p-values, and *qvalueCutoff* is to control false discovery rate (FDR) to prevent high FDR in multiple testing. The *readable* is a logical parameter to indicate the input gene IDs will map to gene symbols or not.

## 4 Biological theme comparison

*clusterProfiler* was developed for biological theme comparison, and it supplies a function, `compareCluster`, to automatically calculate enriched functional categories of each gene clusters.

As we demonstrated in Yu et al. (2012), we analyzed the publicly available expression dataset of breast tumour tissues from 200 patients (GSE11121, Gene Expression Omnibus) (Schmidt et al., 2008). We identified 8 gene clusters from differentially expressed genes, and using `compareCluster` to compare these gene clusters by their enriched biological process, with the strict cutoff of p-values < 0.01 and q-values < 0.05. The analysis result was illustrated in Figure 1. More details of this analysis are described in Yu et al. (2012).

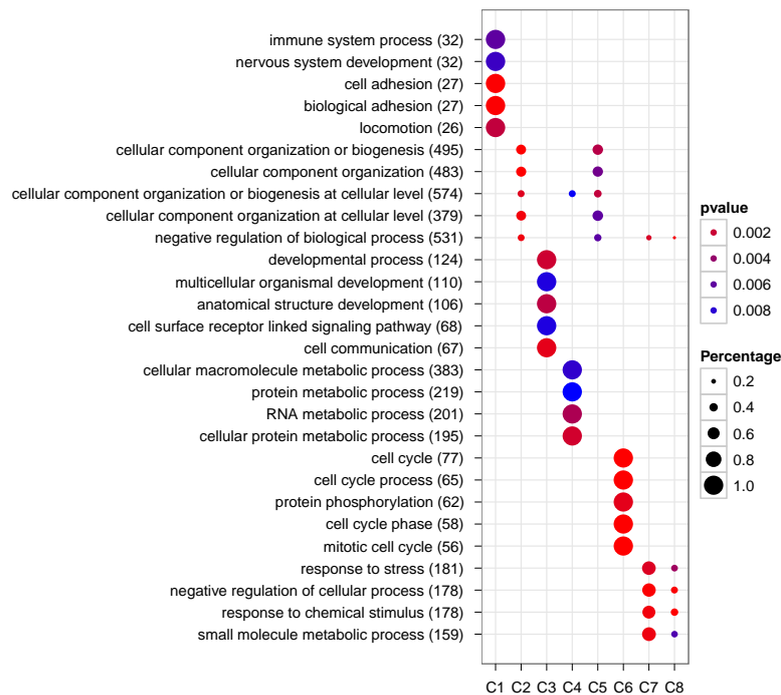


Figure 1: Comparison of GO enrichment of gene clusters

Another example was shown in Yu and He (2011), we calculated functional similarities among viral miRNAs using method described in Yu et al. (2011), and compared significant KEGG pathways regulated by different viruses using *clusterProfiler*.

The comparison function was designed as a general-package for comparing gene clusters of any kind of gene-ontology associations, not only GO and KEGG this package provided, but also other biological and biomedical ontologies.

For example, *compareCluster* can cooperate seamless with *DOSE* and *ReactomePA* and compare gene cluster in the context of disease and reactome pathway as demonstrated in the online vignette of *DOSE* and *ReactomePA* respectively.

## 5 Visualization

*clusterProfiler* implemented several methods for visualizing analyzed result.

- Bar Plot

Bar plot was used to visualized functional profile of the given collection of genes.

The plot function call was consistent for analysis results generated by *groupGO*, *enrichGO* and *enrichKEGG*.

Users can try the following command:

```
> plot(x, type="bar", order=FALSE, drop=TRUE)
> plot(z, type="bar", font.size=12)
```

```
> plot(y, type="bar", title="MF Enrichment analysis", showCategory=10)
```

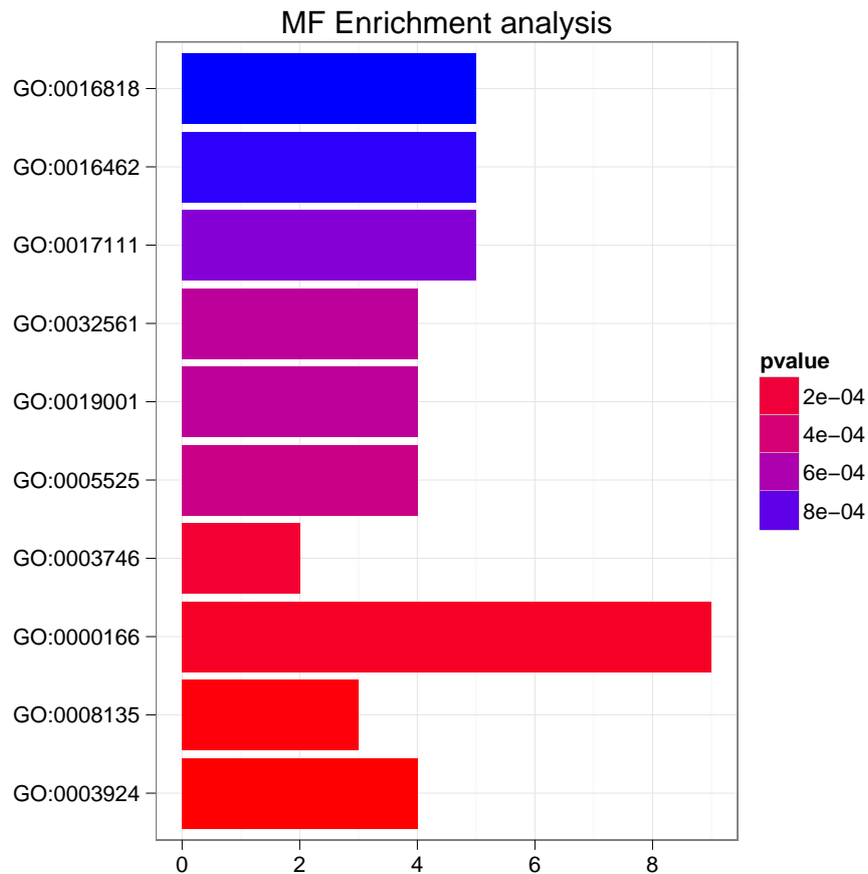


Figure 2: Example of plotting functional profiles

- Category Net Plot

Category-gene network model was also implemented to extract the complex relationships between genes and associated categories. It provides a high-level model to understand the functionalities of genes.

The plot function call was consistent for analysis results generated by `groupGO`, `enrichGO` and `enrichKEGG`.

Users can try the following command:

```
> plot(y, type="cnet", categorySize="geneNum")
> plot(z, type="cnet", categorySize="pvalue", output="interactive")
```

- Dot Plot

Dot plot was implemented for cluster comparison as shown in Figure 1. Here, we demonstrated the functional call of `compareCluster`.

```
> xx <- compareCluster(gcSample,
+                       fun="enrichGO",
```

```
> plot(x, type="cnet", showCategory=5, output="fixed")
```

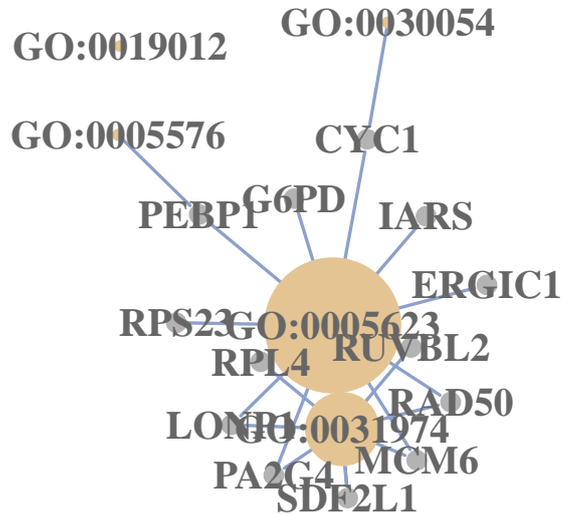


Figure 3: Example of plotting GO profiles using cnetplot

```
+ ont="CC",
+ organism="human",
+ pvalueCutoff=0.05,
+ qvalueCutoff=0.05)
> plot(xx)
```

Bar plot was also supported to visualize cluster comparison. User can try the following command to explore the usage:

```
> plot(xx, type="bar", by="percentage")
> plot(xx, type="bar", by="count")
```

By default, only top 5 (most significant) categories of each cluster was plotted. User can changes the parameter *showCategory* to specify how many categories of each cluster to be plotted, and if *showCategory* was set to *NULL*, the whole result will be plotted.

The dot sizes were based on their corresponding row percentage by default, and user can set the parameter

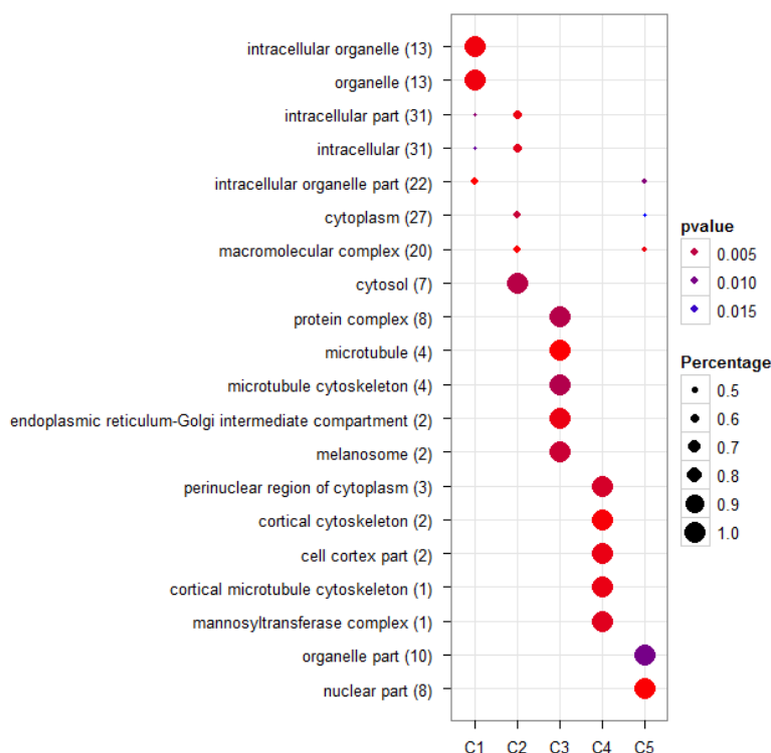


Figure 4: GO Enrichment Comparison

by to "count" to make the comparison based on gene counts. We choose "percentage" as default parameter to represent the size of dots, since some categories may contain a large number of genes, and make the dot sizes of those small categories too small to compare. To provide the full information, we also provide number of identified genes in each category (numbers in parentheses), as shown in Figure 3. If the dot sizes were based on "count", the row numbers will not shown.

The p-values indicate that which categories are more likely to have biological meanings. The dots in the plot are color-coded based on their corresponding p-values. Color gradient ranging from red to blue correspond to in order of increasing p-values. That is, red indicate low p-values (high enrichment), and blue indicate high p-values (low enrichment). P-values were filtered out by the threshold giving by parameter *pvalueCutoff*, and FDR was control by parameter *qvalueCutoff*.

`compareCluster` was designed as a general function for comparing gene clusters of any kind of gene-ontology associations, not only GO (`groupGO` and `enrichGO`) and KEGG (`enrichKEGG`) provided in this package, but also other biological or biomedical ontologies, including Disease Ontology (via `enrichDO` in *DOSE*) and Reactome Pathway (via `enrichPathway` in *ReactomePA*). More details can be found in the vignettes of *DOSE* and *ReactomePA*.

## 6 Session Information

The version number of R and packages loaded for generating the vignette were:

R Under development (unstable) (2012-02-29 r58536)  
Platform: i686-pc-linux-gnu (32-bit)

locale:

```
[1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8      LC_COLLATE=C
[5] LC_MONETARY=en_US.UTF-8  LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=C                LC_NAME=C
[9] LC_ADDRESS=C              LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
```

attached base packages:

```
[1] stats      graphics  grDevices  utils      datasets
[6] methods    base
```

other attached packages:

```
[1] GO.db_2.6.1           org.Hs.eg.db_2.6.4
[3] clusterProfiler_1.3.14 AnnotationDbi_1.17.26
[5] Biobase_2.15.4       BiocGenerics_0.1.13
[7] RSQLite_0.11.1       DBI_0.2-5
```

loaded via a namespace (and not attached):

```
[1] DO.db_2.3.0          DOSE_1.1.12
[3] IRanges_1.13.28     KEGG.db_2.6.1
[5] MASS_7.3-17         RColorBrewer_1.0-5
[7] colorspace_1.1-1    dichromat_1.2-4
[9] digest_0.5.1        ggplot2_0.9.0
[11] grid_2.15.0         igraph_0.5.5-4
[13] memoise_0.1         munsell_0.3
[15] plyr_1.7.1          proto_0.3-9.2
[17] qvalue_1.29.0       reshape2_1.2.1
[19] scales_0.2.0        stats4_2.15.0
[21] stringr_0.6         tcltk_2.15.0
[23] tools_2.15.0
```

## References

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Guangchuang Yu, Le-Gen Wang, Yanyan Han, and Qing-Yu He. clusterprofiler: an r package for comparing biological themes among gene clusters. *OMICS: A Journal of Integrative Biology*, 16:in press, 2012. ISSN 1536-2310.