

# Reactome Pathway Analysis

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## 1 Introduction

This package is designed for reactome pathway-based analysis. Reactome is an open-source, open access, manually curated and peer-reviewed pathway database.

In *ReactomePA*, we plan to implement:

- pathway enrichment analysis
- gene set enrichment analysis
- functional subpathway (active or perturbed subpathway) detection
- methods for visualization.

## 2 Pathway Enrichment Analysis

Enrichment analysis is a widely used approach to identify biological themes. Here, we implement hypergeometric model to assess whether the number of selected genes associated with reactome pathway is larger than expected. We also implement a category net model for viusalization.

- Hypergeometric model

Hypergeometric model was implemented to assess whether the number of selected genes associated with reactome pathway is larger than expected.

- Category Net Plot

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

- Case Study

Here, we used a vector of sample entrezgene ID, which was converted from an example list of genes from ProfCom <http://webclu.bio.wzw.tum.de/profcom/examples.php>.

```
> require(ReactomePA)
> data(sample)
> sample
```

```

[1] "10140"   "1917"    "1672"    "9445"    "5950"    "307"
[7] "978"     "7077"    "6539"    "3569"    "1545"    "1368"
[13] "6590"    "3081"    "3059"    "5797"    "5817"    "5095"
[19] "4118"    "6948"    "1282"    "1284"    "2261"    "4837"
[25] "1311"    "1428"    "3162"    "5004"    "728441"  "2678"
[31] "5744"    "3484"    "2745"    "2064"    "3486"    "9971"
[37] "8566"    "7453"    "11015"   "10397"   "4162"    "7980"
[43] "486"     "6696"    "3929"    "22795"   "4286"    "1410"
[49] "4653"    "3915"    "5358"    "3912"    "6781"    "1474"
[55] "6648"    "2719"    "3306"

> x <- enrichPathway(gene=sample, pvalueCutoff=0.05, qvalueCutoff=0.05, readable=T)
> head(summary(x))

  pathwayID
216083      216083
422475      422475
381426      381426
1430728     1430728
186797      186797
211859      211859

216083
422475
381426 Homo sapiens: Regulation of Insulin-like Growth Factor (IGF) Activity by IGF
1430728
186797
211859

  GeneRatio    BgRatio      pvalue      qvalue
216083      5/57 85/28363 7.948391e-07 4.936369e-05
422475      5/57 255/28363 1.614918e-04 5.014747e-03
381426      2/57 17/28363 5.292893e-04 1.095722e-02
1430728     8/57 983/28363 7.419227e-04 1.151933e-02
186797      3/57 128/28363 2.198867e-03 2.430808e-02
211859      3/57 131/28363 2.348408e-03 2.430808e-02

  geneID Count
216083      COL4A1/COL4A2/SPP1/LAMC1/LAMB1      5
422475      COL4A1/COL4A2/ERBB2/LAMC1/LAMB1      5
381426      IGFBP1/IGFBP3                      2
1430728     CDA/CYP1B1/HGD/NNMT/HMOX1/GGT1/GLRX/PDXK 8
186797      COL4A1/COL4A2/SPP1                      3
211859      CYP1B1/NNMT/GGT1                      3

> plot(x, showCategory=5)

```

- Compatibal with *clusterProfiler* Bioconductor package *clusterProfiler* designed visualization for comparing biological themes among gene clusters (Yu et al., 2012). More details and parameters are described in the documentation (Rfunction?compareCluster). Figure 2 has been generated using the data, as in Yu et al. (2012).

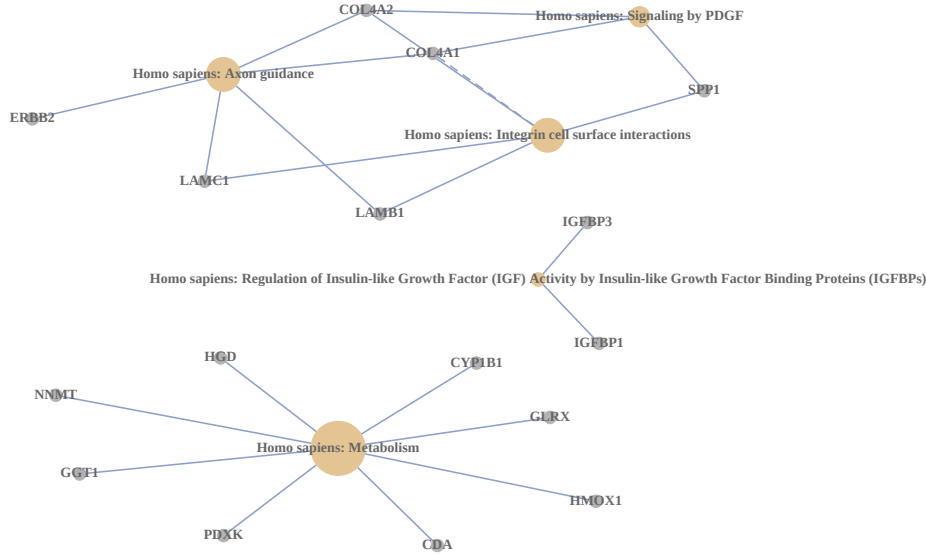


Figure 1: Visualization of Pathway enrichment analysis

### 3 Gene Set Enrichment Analysis

To be developed.

### 4 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] ReactomePA_0.2.4      AnnotationDbi_1.17.23
[3] Biobase_2.15.3        BiocGenerics_0.1.7
[5] RSQLite_0.11.1         DBI_0.2-5
```

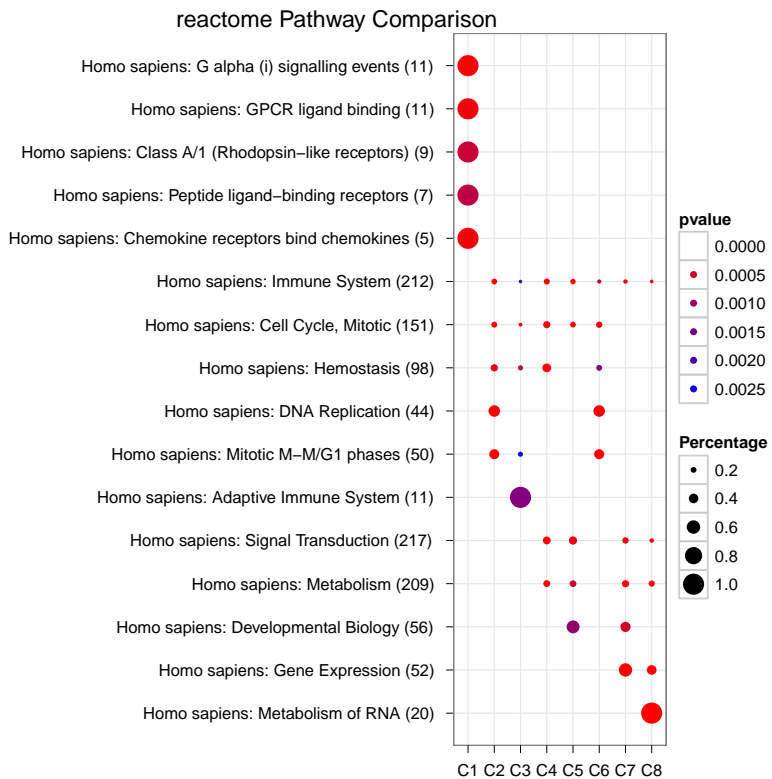


Figure 2: Example of working with clusterProfiler package

```
loaded via a namespace (and not attached):
[1] IRanges_1.13.26      igraph_0.5.5-4      org.Hs.eg.db_2.6.4
[4] plyr_1.7.1           qvalue_1.29.0       reactome.db_1.0.38
[7] tcltk_2.15.0          tools_2.15.0
```

## References

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.