# **KEGGgraph: Application Examples**

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#### **Abstract**

In this vignette, we demonstrate the application of *KEGGgraph* as flexible module in analysis pipelines targeting heterogenous biological questions. For basic use of the *KEGGgraph* package, please refer to the vignette *KEGGgraph*: a graph approach to *KEGG PATHWAY* in *R* and *Bioconductor*.

### 1 Introduction

In many cases, *KEGGgraph* is used as stand-alone software package to parse KEGG pathway [Kanehisa *et al.*, 2008] into graph models and consequently to analyze them. However, *KEGGgraph* can also be combined with other tools (within or out of the scope of R and Bioconductor [Gentleman *et al.*, 2004, Carey *et al.*, 2005]), to build flexible analysis pipelines. In this vignette we demonstrate the cooperation between *KEGGgraph* and other tools with several examples.

## 2 Parse created or edited pathway

KGML-ED [Klukas and Schreiber, 2007] is a tool designed for the dynamic visualization, interactive navigation and editing of KEGG pathway diagrams. The program is able to modify or even create *de novo* new pathways and export them into KGML (KEGG XML) files. Since *KEGGgraph* captures the graph information of KEGG PATHWAY by parsing KGML files, it is also capable to parse the KGML files exported from KGML-ED program. The joint use of the two programs allows user edit KGML files from KEGG (for example, by adding/removing new nodes/edges), or create new pathways from scratch, and then analyse them.

Here we demontrate the parse of a toy pathway (network motif) created by KGML-ED, which illustrates a feed-forward loop (FFL) [Mangan and Alon, 2003, Alon, 2007].

```
> library(KEGGgraph)
```

<sup>&</sup>gt; toyKGML <- system.file("extdata/kgml-ed-toy.xml", package="KEGGgraph")</pre>

<sup>&</sup>gt; toyGraph <- parseKGML2Graph(toyKGML, genesOnly=FALSE)</pre>

<sup>&</sup>gt; toyGraph

```
A graphNEL graph with directed edges

Number of Nodes = 5

Number of Edges = 6

> nodes(toyGraph)

[1] "CRP" "GalS" "galETK" "cAMP" "galactose"
```

We visualize the graph with *Rgraphviz* in Figure 1.

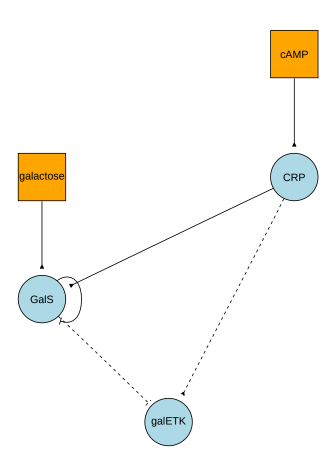


Figure 1: Visualization of the toy network (incoherent type-1 feed-forward loop, IL1-FFL) created by *KGML-ED* and parsed by *KEGGgraph*. The rectangle nodes represent compounds while ellipse ones represent gene/gene products. Solid and dashed red arrows represent activation and expression, whereas blue represent inhibition and repression. Note that self-inhibition of GalS is presented by a node and an edge pointing to its self.

Similarly, users could create new or modify existing pathways by modifying KGML files with tools including KGML-ED, and these pathways can be consequently parsed, visualized and analyzed consequently by *KEGGgraph* and other tools for graphs implemented in Bioconductor. In this sense *KEGGgraph* is not only able to study existing biological pathways constructed by KEGG, but also a tool for modelling pathways and biological networks in R.

### 3 Microarray analysis

KEGG PATHWAY contains pathway maps for nearly 1000 organisms and has been intensively expanded with pathways including signaling transduction, cellular processes and human diseases in recent years. As the graph-theory based approach to this valuable knowledge-base, *KEGGgraph* can be built into analysis pipelines where the pathway information could be useful, especially with the graph attributes. To demonstrate this we show one example of microarray analysis, cooperating with another package SPIA [Tarca *et al.*, 2009], a package implementing the Signaling Pathway Impact Analysis (SPIA) algorithm.

The microarray data is decribed in the vignette of *SPIA*, here it is only briefly summarized: Affemetrix GeneChip (GEO accession: GSE4107), containing 10 normal samples and 12 colorectal cancer samples, is normalized with *affy* and *limma* package. The result of topTable in *limma* is used as the input data.

```
> if(require(SPIA)) {
    data(colorectalcancer, package="SPIA")
+ } else {
+
    load(system.file("extdata/colorectalcancerSPIA.RData", package="KEGGgraph"),
+ }
> head(top)
                     logFC
                                                   P. Value
                                                               adj.P.Val
               ID
                           AveExpr
                                            t
10738
        201289_at 5.960206 6.226928 23.94388 1.789221e-17 9.782565e-13 25.40124
        209189 at 5.143502 7.487305 17.42995 1.560212e-14 2.843486e-10 21.02120
18604
11143 201694_s_at 4.148081 7.038281 16.46040 5.153117e-14 7.043667e-10 20.14963
10490 201041_s_at 2.429889 9.594413 14.06891 1.293706e-12 1.414667e-08 17.66883
10913 201464_x_at 1.531126 8.221044 10.98634 1.685519e-10 1.151947e-06 13.61189
11463
        202014 at 1.429269 5.327647 10.45906 4.274251e-10 2.418771e-06 12.80131
      ENTREZ
10738
        3491
18604
        2353
11143
        1958
10490
        1843
        3725
10913
      23645
11463
```

We only consider the probes annotated with EntrezGeneID and differentially expressed with FDR *p*-value less than 0.05.

```
> library(hgu133plus2.db)
> x <- hgu133plus2ENTREZID
> top$ENTREZ <- unlist(as.list(x[top$ID]))
> top <- top[!is.na(top$ENTREZ),]
> top <- top[!duplicated(top$ENTREZ),]
> tg1 <- top[top$adj.P.Val < 0.05,]
> DE_Colorectal <- tg1$logFC
> names(DE_Colorectal) <- as.vector(tg1$ENTREZ)
> ALL Colorectal <- top$ENTREZ</pre>
```

The SPIA algorithm takes as input two vectors: log2 fold changes of differentially expressed genes (DE\_Colorectal) and those of all EntrezGeneID annotated genes (ALL\_Colorectal), and returns a table of pathways ranked from the most to the least significant. The results show that Colorectal cancer pathway (ID:05210) is significantly activated (ranked the 5th of the result table). To visualize the results, we visualize the pathway with differentially expressed genes marked with color.

For the first time we could use the retrieveKGML function to retrieve the KGML remotely from the KEGG FTP site.

```
> tmp <- "hsa05210.xml"
> retrieveKGML(pathwayid="05210", organism="hsa", destfile=tmp)
```

We have attached this file in the *KEGGgraph*, we parse it into graph and observe that around 30% percent of the genes in the pathway are differentially expressed. The pathway map (figure) can be found at http://www.genome.jp/dbget-bin/show\_pathway?hsa05210.

We visualize the pathway with pseudo-colors representing the log2 fold change of the differentially expressed genes in figure 2.

In the figure 2, we observe that:

• Not all the genes are connected with each other, 40.4761904761905% (34/84) nodes in the pathway have a degree of 0 – they are not conencted to any other node. There are several reasons for this. One of them is the genes with genetic alterations are indicated in the pathway map, but not interweaved into the pathway (cf. http://www.genome.jp/dbget-bin/show\_pathway?hsa05210). Another important factor is that in some pathways, especially human disease pathways, KEGG records genes involved in the disease

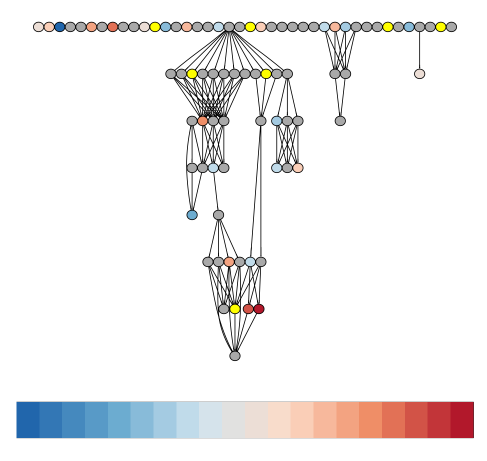


Figure 2: Overview of colorectal cancer pathway with pseudo-colored nodes representing expression change. Red nodes represented up-regulated mRNAs in cancer samples and blue ones are down-regulated. Grey nodes are detected not to be differentially expressed, yellow ones are not reported (possibly due to under detection threshold). The gradual colors indicate the log2 fold change of the expression - the darker the colors, the larger fold change. One distinct feature is the existence of many non-connected nodes, which could have been omitted when the graph information is discarded and only the membership of the gene is considered.

but does not provide any edge in that pathway, although one can find their interaction partners in other pathways (e.g., Grb2 has no edges in colorectal cancer pathway, however in other maps including MAPK signaling pathway, both up- and down-stream interactors are found). One solution to the later problem is to *merge* (*union*) the related pathways together, to this end *KEGGgraph* provides the function mergeKEGGgraphs.

• From the visualization in 2, it is difficult to recognize any patterns. To examine the pathway in more details, it is necessary to use *Divide and Conquer* strategy, namely to *subset* the graph into subgraphs. To this end *KEGGgraph* provides the function subKEGGgraph, which maintains the KEGG information while subsetting the graph.

Out of 34 nonconnected genes, 7 is up-regulated and 4 downregulated. Notably, out of 10 *Frizzled* homologues (Wnt receptors), four are up-regulated (*FZD2*, *FZD4*, *FZD7* and *FZD10*), and only one is down-regulated(*FZD5*), in accordance with the former report [Vincan *et al.*, 2007].

Next we concentrate on upregulated genes, the result given by SPIA package indicates that colorectal pathway is activated.

We can use degree centrality (the sum of in- and out-degree of each node) as a measure of the relative importance of the node compared to others. In this subgraph, 8 nodes have an degree equal or larger than three (in decreasing order: AKT3, TCF7L1, CCND1, FOS, MAPK3, MAPK1, JUN, MAPK10), and 5 of them are upreguated to different extent. This reinforces the conclusion of *SPIA* that the pathway is activated - it seems that the relative important nodes are up-regulated. Especially we note the upregulation of Jun and Fos, two transcription factors with many targeting genes, although their up-stream interactors (MAPK3 and MAPK1) are either not significantly differentially expressed or slightly down-regulated (MAPK1, logFC=-0.381).

Further analysis could done, for example, by merging the colorectal pathway with linked pathways (Wnt signaling, apoptosis, etc) and investigate the graph characteristics of differentially expressed genes and their links.

### References

[Gentleman *et al.*, 2004] Gentleman *et al.* (2004). Bioconductor: open software development for computational biology and bioinformatics, *Genome Biology*, **5**, R80.

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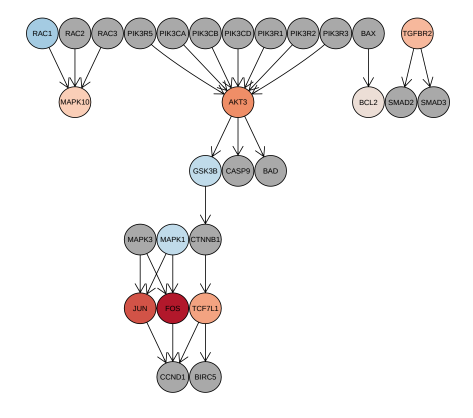


Figure 3: Up-regulated genes (seven, in red) and their neighborhood genes in colorectal cancer pathway, the color follows the legend in figure 2. Un-reported genes are hidden from the figure.

[Aittokallio and Schwikowski, 2006] Aittokallio and Schwikowski. (2006). Graph-based methods for analysing networks in cell biology, *Briefings in Bioinformatics*, **7**, 243-255.

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