KEGGgraph: a graph approach to KEGG PATHWAY in R and Bioconductor

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Abstract

We demonstrate the capabilities of the KEGGgraph package, an interface between KEGG pathways and graph model in R as well as a collection of tools for these graphs. Superior to preceding approaches, KEGGgraph maintains the pathway topology and allows further analysis or dissection of pathway graphs. It parses the regularly updated KGML (KEGG XML) files into graph models maintaining all essential pathway attributes.

1 Introduction

Since its first introduction in 1995, KEGG PATHWAY has been widely used as a reference knowledge base for understanding biological pathways and functions of cellular processes. The knowledge from KEGG has proven of great value by numerous work in a wide range of fields [Kanehisa et al., 2008].

Pathways are stored and presented as graphs on the KEGG server side, where nodes are molecules (protein, compound, etc) and edges represent relation types between the nodes, e.g. activation or phosphorylation. The graph nature raised our interest to investigate them with powerful graph tools implemented in R and Bioconductor [Gentleman et al., 2004], including graph, RBGL and Rgraphviz [Carey et al., 2005]. While it is barely possible to query the graph characteristics by manual parsing, a native and straightforward client-side tool is currently missing to parse pathways and analyze them consequently with tools for graph in R.

To address this problem, we developed the open-source software package KEGGgraph, an interface between KEGG pathway and graph object as well as a collection of tools to analyze, dissect and visualize these graphs.

The package requires KGML (KEGG XML) files, which can be downloaded from KEGG REST web service (https://rest.kegg.jp/) without license permission for academic purposes. To demonstrate the functionality, in “extdata” sub-directory of KEGGgraph we have pre-installed several KGML files.
2 Software features

KEGGgraph offers the following functionalities:

Parsing: It should be noted that one 'node' in KEGG pathway does not necessarily map to merely one gene product, for example the node 'ERK' in the human TGF-Beta signaling pathway contains two homologues, MAPK1 and MAPK3. Therefore, among several parsing options, user can set whether to expand these nodes topologically. Beyond facilitating the interpretation of pathways in a gene-oriented manner, the approach also entitles unique identifiers to nodes, enabling merging graphs from different pathways.

Graph operations: Two common operations on graphs are subset and merge. A sub-graph of selected nodes and the edges in between are returned when subsetting, while merging produces a new graph that contains nodes and edges of individual ones. Both are implemented in KEGGgraph.

Visualization: KEGGgraph provides functions to visualize KEGG graphs with custom style. Nevertheless users are not restricted by them, alternatively they are free to render the graph with other tools like the ones in Rgraphviz.

Besides the functionalities described above, KEGGgraph also has tools for remote KGML file retrieval, graph feature study and other related tasks. We will demonstrate them later in this vignette.

3 Case studies

We load the KEGGgraph by typing or pasting the following codes in R command line:

> library(KEGGgraph)

3.1 Get KGML files

There are at least two possibilities to get KGML (KEGG XML) files:

• Manual download from KEGG REST API, documented at at [https://www.kegg.jp/kegg/rest/keggapi.html]

• Automatic retrieval from KEGG REST API, available at [https://rest.kegg.jp] with the function retrieveKGML.

To retrieve KGML file automatically, one has to know the pathway identifier. KEGG pathway identifiers have the form of [a-z]3-4[0-9]5, where the three- or four-alphabet code represent the organism and the five digits represent pathways. The same pathway in different species share the five-digit code, which is the first parameter accepted by retrieveKGML. The second parameter required is the alphabet-code of the organism.

For example, the following codes retrieve p53 signaling pathway, which has the digit-code 04115, of the species Macaca fascicularis, or crab-eating macaque, which has the three-alphabet code mcf:
Pathways and their identifiers can be browsed in the KEGG website [https://genome.jp/kegg](https://genome.jp/kegg), or found with any search engine.

### 3.2 Parsing and graph feature query

First we read in KGML file for human MAPK signaling pathway (with KEGG ID hsa04010):

```r
> mapkKGML <- system.file("extdata/hsa04010.xml",
+ package="KEGGgraph")
```

Once the file is ready, we can either parse them into an object of `KEGGPathway` or an object of `graph`. `KEGGPathway` object maintains the information of the pathway (title, link, organism, etc), while `graph` objects are more natural approach and can be directly plugged in many other tools. To convert the pathway into graph, we use

```r
> mapkG <- parseKGML2Graph(mapkKGML,expandGenes=TRUE)
> mapkG
```

A graphNEL graph with directed edges

- Number of Nodes = 265
- Number of Edges = 876

Alternatively we can parse the KGML file first into an object of `KEGGpathway`, which can be later converted into the graph object, as the following lines show:

```r
> mapkpathway <- parseKGML(mapkKGML)
> mapkpathway
```

**KEGG Pathway**

- **Title**: MAPK signaling pathway
- **Name**: path:hsa04010
- **Organism**: hsa
- **Number**: 04010
- **Image**: [http://www.genome.jp/kegg/pathway/hsa/hsa04010.gif](http://www.genome.jp/kegg/pathway/hsa/hsa04010.gif)
- **Link**: [http://www.genome.jp/dbget-bin/show_pathway?hsa04010](http://www.genome.jp/dbget-bin/show_pathway?hsa04010)

---

Statistics:

- 136 node(s)
- 171 edge(s)
- 0 reaction(s)

---
> mapkG2 <- KEGGpathway2Graph(mapkpathway, expandGenes=TRUE)

A graphNEL graph with directed edges
Number of Nodes = 265
Number of Edges = 876

There is no difference between graph objects derived from two approaches.

The option 'expandGenes' in parsing controls whether the nodes of paralogues in pathways should be expanded or not. Since one ‘node’ in KEGG pathway does not necessarily map to only one gene/gene product (e.g. 'ERK' maps to MAPK1 and MAPK3), the option allows expanding these nodes and takes care of copying existing edges.

Another option users may find useful is 'genesOnly', when set to TRUE, the nodes of other types than 'gene' (compounds, for example) are neglected and the result graph consists only gene products. This is especially desired when we want to query network characteristics of gene products. Its value is set to 'TRUE' by default.

The following commands extract node and edge information:

> mapkNodes <- nodes(mapkG)
> nodes(mapkG)[1:3]
[1] "hsa:5923" "hsa:5924" "hsa:11072"

> mapkEdges <- edges(mapkG)
> edges(mapkG)[1]

`hsa:5923`
[1] "hsa:22800" "hsa:22808" "hsa:3265" "hsa:3845" "hsa:4893" "hsa:6237"

Edges in KEGG pathways are directional, that is, an edge starting at node A pointing to node B does not guarantee a reverse relation, although reciprocal edges are also allowed. When listing edges, a list indexed with node names is returned. Each item in the list records the nodes pointed to.

We can also extract the node attributes specified by KEGG with getKEGGnodeData:

> mapkGnodedata <- getKEGGnodeData(mapkG)
> mapkGnodedata[[2]]

KEGG Node (Entry 'hsa:5924）:
------------------------------------------------------------
[ displayName ]: RASGRF1, GRF1...
[ Name ]: hsa:5924
[ Type ]: gene
[ Link ]: http://www.genome.jp/dbget-bin/www_bget?hsa+5923+5924
------------------------------------------------------------
An alternative to use `getKEGGnodeData` is

```r
> getKEGGnodeData(mapkG, 'hsa:5924')
```

KEGG Node (Entry 'hsa:5924'):

```
[ displayName ]: RASGRF1, GRF1...
[ Name ]: hsa:5924
[ Type ]: gene
[ Link ]: http://www.genome.jp/dbget-bin/www_bget?hsa+5923+5924
```

, returning identical results.

Similarly the `getKEGGedgeData` is able to extract edge information:

```r
> mapkGedgedata <- getKEGGedgeData(mapkG)
> mapkGedgedata[[4]]
```

KEGG Edge (Type: PPrel):

```
[ Entry 1 ID ]: hsa:5923
[ Entry 2 ID ]: hsa:3845
[ Subtype ]:
  [ Subtype name ]: activation
  [ Subtype value ]: -->
```

Alternatively the query above can be written as:

```r
> getKEGGedgeData(mapkG,'hsa:627~hsa:4915')
```

KEGG Edge (Type: PPrel):

```
[ Entry 1 ID ]: hsa:627
[ Entry 2 ID ]: hsa:4915
[ Subtype ]:
  [ Subtype name ]: activation
  [ Subtype value ]: -->
```

For `KEGGNode` and `KEGGEdge` objects, methods are implemented to fetch their attributes, for example `getName`, `getType` and `getDisplayName`. Guides to use these methods as well as examples can be found in help pages.
This case study finishes with querying the degree attributes of the nodes in the graph. We ask the question which nodes have the highest out- or in-degrees. Roughly speaking the out-degree (number of out-going edges) reflects the regulatory role, while the in-degree (number of in-going edges) suggests the subjectability of the protein to intermolecular regulations.

```r
> mapkGoutdegrees <- sapply(edges(mapkG), length)
> mapkGindegrees <- sapply(inEdges(mapkG), length)
> topouts <- sort(mapkGoutdegrees, decreasing=T)
> topins <- sort(mapkGindegrees, decreasing=T)
> topouts[1:3]
hsa:5594  hsa:5595  hsa:1432
   26     26     13

> topins[1:3]
hsa:5923  hsa:5924  hsa:10125
   26     26     26
```

3.3 Graph subset and merge

We demonstrate the subsetting of the graph with 25 randomly chosen nodes of MAPK pathway graph:

```r
> library(Rgraphviz)
> set.seed(123)
> randomNodes <- sample(nodes(mapkG), 25)
> mapkGsub <- subGraph(randomNodes, mapkG)
> mapkGsub

A graphNEL graph with directed edges
Number of Nodes = 25
Number of Edges = 6
```

The subgraph is visualized in figure [1] where nodes with in-degree or out-degree in red and others in grey [2]. And in the example we also demonstrate how to convert KEGG ID into other other identifiers via the Entrez GeneID. More details on the conversion of IDs can be found on page [3].

Another common operation on graphs is merging, that is, combining different graphs together. It is inspired by the fact that many KEGG pathways embed other pathway, for example MAPK

---

1The makeAttr function is used to assign nodes with rendering attributes, whose code can be found in the Rnw file.
> outs <- sapply(edges(mapkGsub), length) > 0
> ins <- sapply(inEdges(mapkGsub), length) > 0
> ios <- outs | ins
> ## translate the KEGG IDs into Gene Symbol
> if(require(org.Hs.eg.db)) {
+   ioGeneID <- translateKEGGID2GeneID(names(ios))
+   nodesNames <- sapply(mget(ioGeneID, org.Hs.egSYMBOL, ifnotfound=NA), "[[",1)
+ } else {
+   nodesNames <- names(ios)
+ }
> names(nodesNames) <- names(ios)
> nAttrs <- list();
> nAttrs$fillcolor <- makeAttr(mapkGsub, "lightgrey", list(orange=names(ios)[ios]))
> nAttrs$label <- nodesNames
> plot(mapkGsub, "neato", nodeAttrs=nAttrs,
+     attrs=list(node=list(fillcolor="lightgreen",
+                 width="0.75", shape="ellipse"),
+                 edge=list(arrowsize="0.7")))

Figure 1: A random subgraph of MAPK signaling pathway
signaling pathway embeds 6 pathways including Wnt signaling pathway. mergeGraphs provides the possibility to merge them into one graph for further analysis. Next we merge MAPK and Wnt signaling pathway into one graph. The graphs to be merged should be organized into a list, and it is commandary to use ‘expandGenes=TRUE’ option when parsing to make sure the nodes are unique and indexed by KEGGID.

```r
> wntKGML <- system.file("extdata/hsa04310.xml",package="KEGGgraph")
> wntG <- parseKGML2Graph(wntKGML)
> graphs <- list(mapk=mapkG, wnt=wntG)
> merged <- mergeGraphs(graphs)
> merged
A graphNEL graph with directed edges
Number of Nodes = 386
Number of Edges = 1628
```

We observe that the node number in the merged graph (386) is less than the sum of two graphs (265 and 148 for MAPK and Wnt pathway respectively), reflecting the crosstalk between the pathways by sharing nodes.

### 3.4 Using other graph tools

In R and Bioconductor there are powerful tools for graph algorithms and operations, including *graph*, *Rgraphviz* and *RBGL*. The KEGG graphs can be analyzed with their functionalities to describe characteristics of the pathway and to answer biological relevant questions.

Here we demonstrate the use of other graph tools with asking the question which nodes are of the highest importance in MAPK signalling pathway. To this end we turn to relative betweenness centrality [Aittokallio and Schwikowski, 2006][Carey et al., 2005]. Betweenness is a centrality measure of a node within a graph. Nodes that occur on many shortest paths between other vertices have higher betweenness than those that do not. It is scaled by the factor of (n-1)(n-2)/2 to get relative betweenness centrality, where n is the number of nodes in the graph. Both measurements estimate the importance or the role of the node in the graph.

With the function implemented in *RBGL*, our aim is to identify most important nodes in MAPK signalling pathway.

*Note:* the current version of RBGL (version 1.59.5) reports the error that BGL_brandes_betweeness_centrality not available for .Call() for package “RBGL”. Therefore the execution has been suppressed for now.

```r
> library(RBGL)
> bcc <- brandes.betweenness.centrality(mapkG)
> rbccs <- bcc$relative.betweenness.centrality.vertices[1L,]
> toprbccs <- sort(rbccs,decreasing=TRUE)[1:4]
> toprbccs
```
We identify the top 4 important nodes judged by betweenness centrality as MAP3K1 (hsa:4214), GRB2 (hsa:2885), MAP2K2 (hsa:5605) and MAP2K1 (hsa:5604) (the mapping between KEGG ID and gene symbol is done using biomaRt, see page [14]). In figure 2, we illustrate them as well as their interacting partners in MAPK pathway.

### 4 Other functionalities

Besides the ability to parse and operate on KEGG PATHWAY graph objects, the KEGGgraph package also provides functionalities to complement tasks related to deal with KEGG pathways. We introduce some of them here, for a full list of functions please see the package help file:

```r
> help(package=KEGGgraph)
```

#### 4.1 Parsing chemical compound reaction network

KEGG PATHWAY captures two kinds of network: the protein network and the chemical network. The protein network consists relations (edges) between gene products, while the chemical network illustrate the relations between chemical compounds. Since the metabolic pathway can be viewed both as a network of proteins (enzymes) and as a network of chemical compounds, metabolic pathways can be viewed as both protein networks and chemical networks, whereas regulatory pathways are always viewed as protein networks only. KEGGPathway provides methods to access this network.

We show the example of Glycine, serine and threonine metabolism pathway.

```r
> mapfile <- system.file("extdata/map00260.xml",package="KEGGgraph")
> map <- parseKGML(mapfile)
> map

KEGG Pathway
[ Title ]: Glycine, serine and threonine metabolism
[ Name ]: path:map00260
[ Organism ]: map
[ Number ]: 00260
[ Image ]: http://www.genome.jp/kegg/pathway/map/map00260.gif
[ Link ]: http://www.genome.jp/dbget-bin/show_pathway?map00260

------------------------------------------------------------

Statistics:
144 node(s)
371 edge(s)
```
> toprbccName <- names(toprbccs)
> toprin <- sapply(toprbccName, function(x) inEdges(mapkG)[x])
> toprout <- sapply(toprbccName, function(x) edges(mapkG)[x])
> toprSubnodes <- unique(unname(c(unlist(toprin), unlist(toprout), toprbccName)))
> toprSub <- subGraph(toprSubnodes, mapkG)
> nAttrs <- list()
> tops <- c("MAPK3K1","GRB2","MAP2K2","MAP2K1")
> topLabels <- lapply(toprbccName, function(x) x); names(topLabels) <- tops
> nAttrs$label <- makeAttr(toprSub, "", topLabels)
> nAttrs$fillcolor <- makeAttr(toprSub, "lightblue", list(orange=toprbccName))
> nAttrs$width <- makeAttr(toprSub,"",list("0.8"=toprbccName))
> plot(toprSub, "twopi", nodeAttrs=nAttrs, attrs=list(graph=list(start=2)))

Figure 2: Nodes with the highest relative betweenness centrality in MAPK pathway (in orange) and their interacting partners (in blue).
> reactions <- getReactions(map)
> reactions[[1]]

KEGG Reaction(rn:R08211)

[ Name ]: rn:R08211
[ Type ]: irreversible
[ Substrate Name ]: cpd:C00576
[ Product Name ]: cpd:C00719

Figure 3 shows how to extract reactions from the pathway and to build a directed graph with them.

4.2 Expand embedded pathways

Function parseKGMLexpandMaps is a function to handle with pathways embedding other pathways. For example, pancreatic cancer pathway embeds 9 other pathways including MAPK and ErbB signaling pathway, cell cycle and apoptosis pathway, etc. To parse them into one graph, the users only have to download the KGML file and feed the file name to parseKGMLexpandMaps, the function parses the file, analyze the embedded pathways, download their files from KEGG REST API service automatically (alternatively a local repository can be specified for KGML files) and merge the individual pathways into a single graph. For example, the following single line parses MAPK signaling pathway with all its embedded pathways

> mapkGembed <- parseKGMLexpandMaps(mapkKGML)

As its name suggests, function subGraphByNodeType subsets the graph by node type, the nodes to subset are those of the type given by the user. It is useful when the KGML file was parsed with 'genesOnly=FALSE' option and later on the user wants only certain kind of node, 'gene' for example, remained. The following example shows how to use it.

> mapkGall <- parseKGML2Graph(mapkKGML, genesOnly=FALSE)
> mapkGall

A graphNEL graph with directed edges
Number of Nodes = 277
Number of Edges = 891

> mapkGsub <- subGraphByNodeType(mapkGall, "gene")
> mapkGsub
Figure 3: Reaction network built of chemical compounds: the orange nodes are the three compounds with maximum out-degree in this network.
A graphNEL graph with directed edges
Number of Nodes = 265
Number of Edges = 876

4.3 Annotation

translateKEGGID2GeneID translates KEGG identifiers (KEGGID) into Entrez GeneID. For example, if we want to find the Entrez GeneID of the nodes in MAPK pathway having the highest relative betweenness centrality, the following codes do the job.

```r
toprbccKEGGID <- names(toprbccs)
toprbccKEGGID
toprbccGeneID <- translateKEGGID2GeneID(toprbccKEGGID)
toprbccGeneID
```

To convert GeneID to other identifiers, we recommend genome wide annotation packages, for human it is `org.Hs.eg.db` and the packages for other organisms can be found at [http://www.bioconductor.org/packages/release/data/annotation/](http://www.bioconductor.org/packages/release/data/annotation/). To demonstrate its use, we draw the sub-network in the figure 2 again, whereas nodes are now labeled with gene symbols.

```r
if(require(org.Hs.eg.db)) {
  tnodes <- nodes(toprSub)
tgeneids <- translateKEGGID2GeneID(tnodes)
tgensymbols <- sapply(mget(tgeneids, org.Hs.egSYMBOL, ifnotfound=NA), 
                        "[[", 
toprSubSymbol <- toprSub
  nodes(toprSubSymbol) <- tgensymbols
  plot(toprSubSymbol, "neato", attrs=list(node=list(font=5, fillcolor="lightblue")))
}
```
Alternatively, users could use R package biomaRt [Durinck et al., 2005, Durinck and Huber, 2008] for ID conversion, whereas it assumes that the user has an internet connection. The following example shows how to translate the node hits we acquired in the example above into HGNC symbols:

```r
> library(biomaRt)
> hsapiens <- useMart("ensembl","hsapiens_gene_ensembl")
> filters <- listFilters(hsapiens)
> getBM(attributes=c("entrezgene","hgnc_symbol"),
+       filters="entrezgene",
+       values=toprbccGeneID, mart=hsapiens)
```

5 Acknowledgement

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Many users have provided very helpful feedback. Here is an incomplete list of them: Martin Morgan, Paul Shannon, Juliane Manitz and Alexander Gulliver Bjørnholt Gjønning.

6 Conclusion

Before the release of KEGGgraph, several R/Bioconductor packages have been introduced and proven their usefulness in understanding biological pathways with KEGG. However, KEGGgraph is the first package able to parse any KEGG pathways from KGML files into graphs. In comparison, existing tools can not achieve the results we present here. They either neglects the graph topology (KEGG.db), do not parse pathway networks (keggorth), or are specialized for certain pathways (cMAP and pathRender).

With KEGGgraph, we contribute a direct and natural approach to KEGG pathways, and the possibilities to study them in R and Bioconductor.

7 Session Info

The script runs within the following session:

R version 4.4.1 (2024-06-14)
Platform: x86_64-pc-linux-gnu
Running under: Ubuntu 24.04.1 LTS

Matrix products: default
BLAS: /home/biocbuild/bbs-3.20-bioc/R/lib/libRblas.so
LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.12.0

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

time zone: America/New_York
tzcode source: system (glibc)

attached base packages:
[1] stats4   grid     stats   graphics  grDevices utils   datasets
[8] methods  base
other attached packages:

- RBGL_1.82.0
- org.Hs.eg.db_3.20.0
- AnnotationDbi_1.68.0
- IRanges_2.40.0
- S4Vectors_0.44.0
- Biobase_2.66.0
- Rgraphviz_2.50.0
- graph_1.84.0
- BiocGenerics_0.52.0
- KEGGgraph_1.66.0

loaded via a namespace (and not attached):

- crayon_1.5.3
- vctrs_0.6.5
- httr_1.4.7
- cli_3.6.3
- rlang_1.1.4
- DBI_1.2.3
- UCSC.utils_1.2.0
- png_0.1-8
- jsonlite_1.8.9
- bit_4.5.0
- RCurl_1.98-1.16
- Biostrings_2.74.0
- XML_3.99-0.17
- KEGGREST_1.46.0
- bitops_1.0-9
- fastmap_1.2.0
- GenomeInfoDb_1.42.0
- memoise_2.0.1
- compiler_4.4.1
- RSQLite_2.3.7
- blob_1.2.4
- pkgconfig_2.0.3
- XVector_0.46.0
- R6_2.5.1
- GenomeInfo_DbData_1.2.13
- tools_4.4.1
- bit64_4.5.2
- zlibbioc_1.52.0
- cachem_1.1.0

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