

# *RBGL*: R interface to boost graph library

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*Summary.* An interface from R to the Boost Graph Library (BGL, an alternative to STL programming for mathematical graph objects) is described.  
*This 2003 update employs the graph class of Bioconductor.*

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## 1 Working with the Bioconductor graph class

An example object representing file dependencies is included, as shown in Figure 1.

```
> library(RBGL)
```

```
Loading required package: graph
Loading required package: cluster
Loading required package: Ruuid
Creating a new generic function for "print" in "Ruuid"
Loading required package: Biobase
Welcome to Bioconductor
```

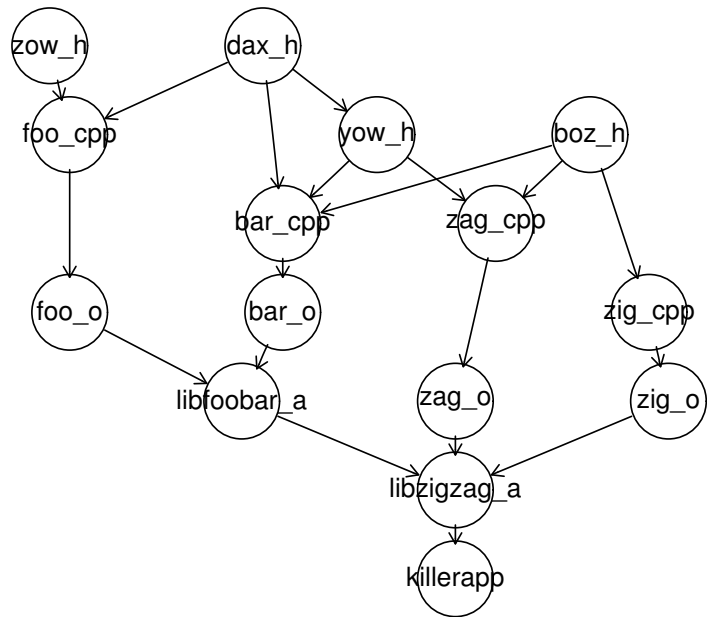


Figure 1: File dependency digraph example from Boost library.

```
Vignettes contain introductory material. To view,
simply type: openVignette()
For details on reading vignettes, see
the openVignette help page.
```

```
> library(Rgraphviz)
> data(FileDep)
> print(FileDep)

A graph with directed edges
Number of Nodes = 15
Number of Edges = 19
```

## 2 Algorithms supported by RBGL

### 2.1 Topological sort

The `tsort` function will return the indices of vertices in topological sort order:

```
> ts <- tsort(FileDep)
> print(nodes(FileDep)[ts + 1])

[1] "zow_h"        "boz_h"        "zig_cpp"      "zig_o"        "dax_h"
[6] "yow_h"        "zag_cpp"      "zag_o"        "bar_cpp"      "bar_o"
[11] "foo_cpp"      "foo_o"        "libfoobar_a" "libzigzag_a" "killerapp"
```

Note that if the input graph is not a DAG, BGL `topological_sort` will check this and throw 'not a dag'. This is crudely captured in the interface (a message is written to the console and zeroes are returned).

```
#FD2 <- FileDep
# now introduce a cycle
#FD2@edgeL[["bar_cpp"]]$edges <- c(8,1)
#tsort(FD2)
```

### 2.2 Kruskal's minimum spanning tree

Function `mstree.kruskal` just returns a list of edges, weights and nodes determining the minimum spanning tree (MST) by Kruskal's algorithm.

```
> km <- fromGXL(file(system.file("GXL/kmstEx.gxl", package = "graph")))
> print(mstree.kruskal(km))

$edgeList
 [,1] [,2] [,3] [,4]
[1,]    1    4    5    2
[2,]    3    5    1    4

$weights
 [,1] [,2] [,3] [,4]
[1,]    1    1    1    1

$nodes
[1] "A" "B" "C" "D" "E"
```

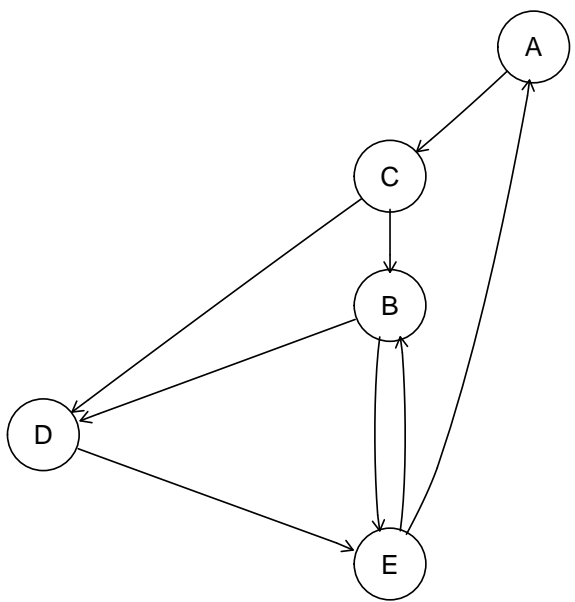


Figure 2: Kruskal MST example from Boost library.

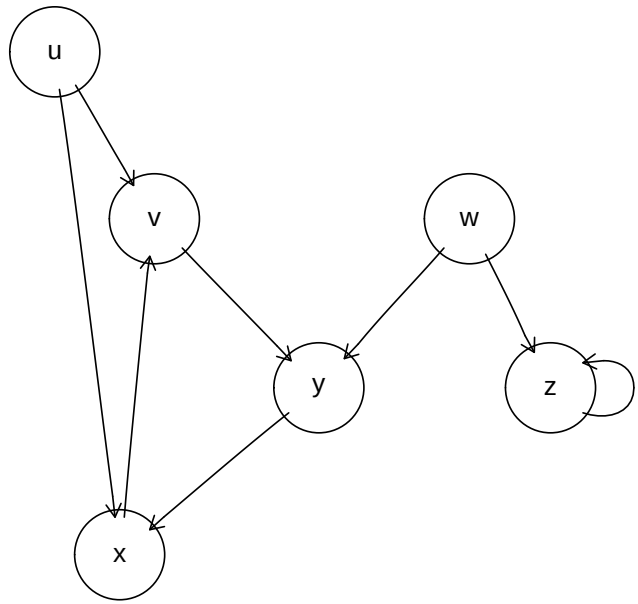


Figure 3: DFS example from Boost library.

### 2.3 Depth first search

The `dfs` function returns a list of node indices by discovery and finish order.

```

> df <- fromGXL(file(system.file("XML/dfsex.gxl", package = "RBGL")))
> print(o <- dfs(df))

$discovered
[1] 1 2 5 4 3 6

$finish
[1] 4 5 2 1 6 3

```

Here is the list of nodes in DFS discovery order.

```

> print(nodes(df)[o$discovered])
[1] "u" "v" "y" "x" "w" "z"

```

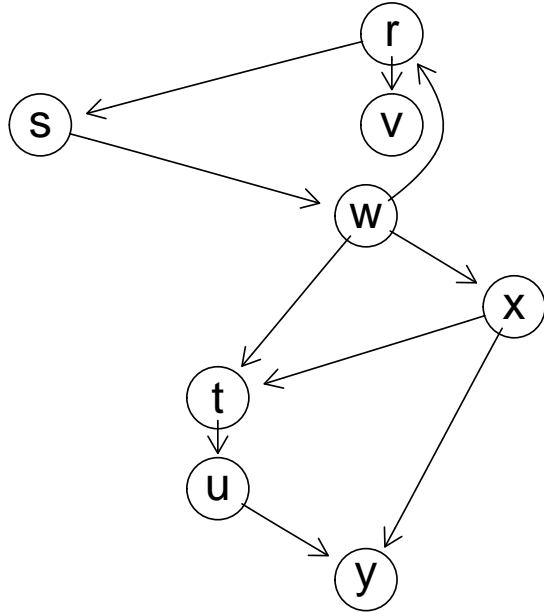
## 2.4 Breadth first search

The `bfs` function returns a vector of node indices for a breadth-first search (BFS) starting at the node indexed by `init.node`.

```
> bf <- fromGXL(file(system.file("XML/bfsex.gxl", package = "RBGL")))
> print(o <- bfs(bf, nodes(bf)[2]))
```

```
[1] 2 6 1 3 7 5 4 8
```

```
> z <- plot(bf)
```



order starting with the second node are

The nodes in BFS

```
> print(nodes(bf)[o])
```

```
[1] "s" "w" "r" "t" "x" "v" "u" "y"
```

## 2.5 Dijkstra's shortest paths

```
> dd <- fromGXL(file(system.file("XML/dijkex.gxl", package = "RBGL")))
> print(dijkstra.sp(dd))

$distances
A B C D E
0 6 1 4 5

$penult
A B C D E
1 5 1 3 4

$start
A
1

> ospf <- fromGXL(file(system.file("XML/ospf.gxl", package = "RBGL")))

> dijkstra.sp(ospf, nodes(ospf)[6])

$distances
RT1 RT2 RT3 RT4 RT5 RT6 RT7 RT8 RT9 RT10 RT11 RT12 N1 N2 N3 N4
  7   7   6   7   6   0   8   8   11   7   10   11   10   10   7   8
N6   N7   N8   N9   N10  N11  N12  N13  N14  N15   H1
  8   12   10   11   13   14   10   14   14   17   21

$penult
RT1 RT2 RT3 RT4 RT5 RT6 RT7 RT8 RT9 RT10 RT11 RT12 N1 N2 N3 N4
  15  15   6   15   6   6   17   17   20   6   19   20   1   2   3   3
N6   N7   N8   N9   N10  N11  N12  N13  N14  N15   H1
  10   8   10   11   12   9   7   5   5   7   12

$start
RT6
6

> sp.between(ospf, "RT6", "RT1")

$"RT6:RT1"
$"RT6:RT1$path
[1] "RT6" "RT3" "N3"  "RT1"
```

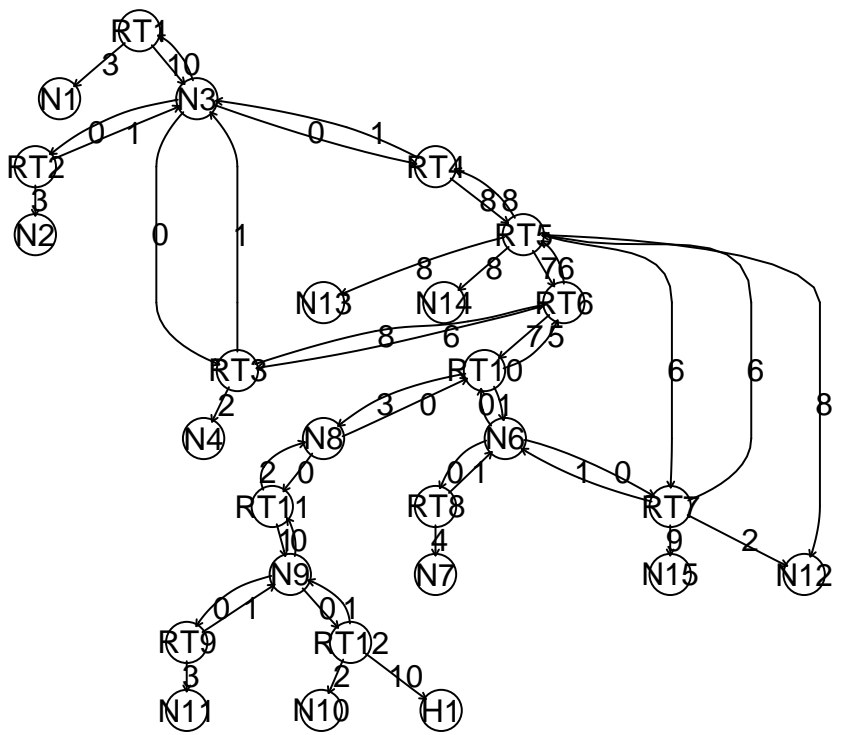


Figure 4: Network example from BGL.

```

$"RT6:RT1">$length
[1] NA

$"RT6:RT1">$pweights
RT6->RT3  RT3->N3  N3->RT1
      NA        NA        NA

> dd <- fromGXL(file(system.file("XML/dijkex.gxl", package = "RBGL")))
> print(dijkstra.sp(dd))

$distances
A B C D E
0 6 1 4 5

$penult
A B C D E
1 5 1 3 4

$start
A
1

```

## 2.6 Connected components

```

> km <- fromGXL(file(system.file("GXL/kmstEx.gxl", package = "graph")))
> km@nodes <- c(km@nodes, "F", "G", "H")
> km@edgeL$F <- list(edges = numeric(0))
> km@edgeL$G <- list(edges = 8, weights = 1)
> km@edgeL$H <- list(edges = 7, weights = 1)
> km@edgemode <- "undirected"
> print(connectedComp(ugraph(km)))

$"1"
[1] "A" "B" "C" "D" "E"

$"2"
[1] "F"

$"3"
[1] "G" "H"

```

## 2.7 Strongly connected components

```
> km <- fromGXL(file(system.file("GXL/kmstEx.gxl", package = "graph")))
> km@nodes <- c(km@nodes, "F", "G", "H")
> km@edgeL$F <- list(edges = numeric(0))
> km@edgeL$G <- list(edges = 8, weights = 1)
> km@edgeL$H <- list(edges = 7, weights = 1)
> km@edgemode <- "directed"
> print(strongComp(km))

$"1"
[1] "A" "B" "C" "D" "E"

$"2"
[1] "F"

$"3"
[1] "G" "H"
```

## 2.8 Edge connectivity and minimum disconnecting set

```
> coex <- fromGXL(file(system.file("XML/conn.gxl", package = "RBGL")))
> dcoex <- coex
> dcoex@edgemode <- "directed"
> udcoex <- ugraph(dcoex)

> print(edgeConnectivity(coex))

$connectivity
[1] 2

$minDisconSet
$minDisconSet[[1]]
[1] "D" "E"

$minDisconSet[[2]]
[1] "D" "H"
```

## 2.9 Min-Cut

```
> coex <- fromGXL(file(system.file("XML/conn.gxl", package = "RBGL")))
> dcoex <- coex
> dcoex@edgemode <- "directed"
> udcoex <- ugraph(dcoex)
```

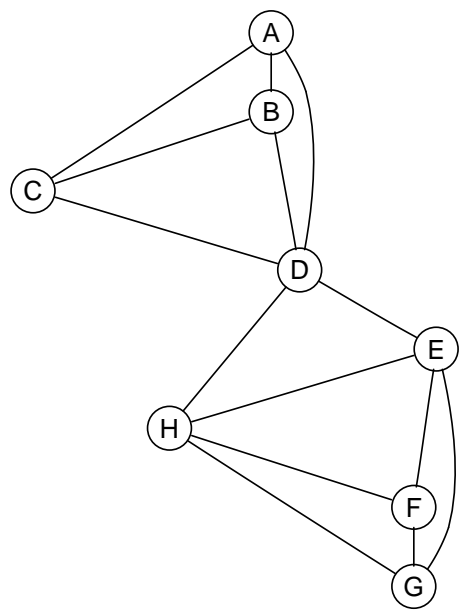


Figure 5: Edge connectivity example.

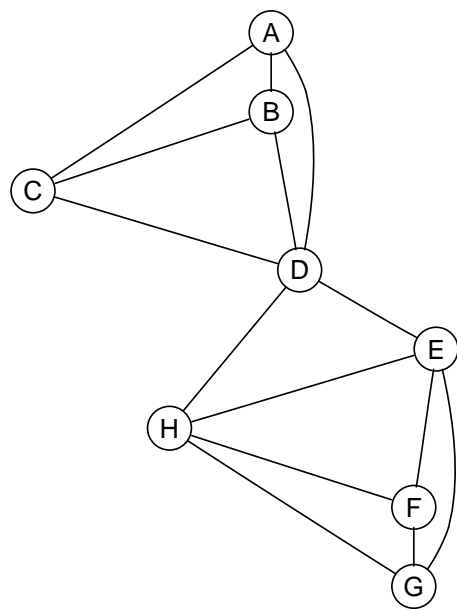


Figure 6: min-cut example.

```
> print(minCut(coex))
```

```
$mincut
```

```
[1] 2
```

```
$S
```

```
[1] 1 2 3 4
```

```
$"V-S"
```

```
[1] 5 6 7 8
```