## \#\#

\#\# Attaching package: 'igraph'
\#\# The following objects are masked from 'package:gRbase':
\#\#
\#\# edges, is_dag, topo_sort
\#\# The following objects are masked from 'package:stats':
\#\#
\#\# decompose, spectrum
\#\# The following object is masked from 'package:base':
\#\#
\#\# union

## Chapter 1

## Graphs and Conditional Independence

> As of major version 2 of $g R b a s e$, that is versions 2.x.y, gRbase no longer depends on the packages graph, Rgraphviz, and RBGL packages. Graph functionality in these packages now relies either on the igraph package or on graph algorithms implemented in gRbase. This document reflects these changes.
> As a consequence, this document provides an up-to-date version of Chapter 1 in the book Graphical Models with $R$ (2012); hereafter abbreviated GMwR, see Højsgaard et al. [2012].
> This document also reflects that since GMwR was published in 2012, some packages that are mentioned in GMwR are no longer on CRAN. This includes the packages lcd and sna.
> In this document it has been emphasized if a function has been imported from igraph or if it is native function from gRbase by writing 'igraph: :this_function()' and 'gRbase: : this_function()',

One notable feature that is not available in this version of $g R b$ ase are functions related to maximal prime subgraph decomposition. They may be reimplented at a later stage.

## Contents

1 Graphs and Conditional Independence ..... 3
1.1 Introduction ..... 6
1.2 Graphs ..... 6
1.2.1 Undirected Graphs ..... 6
1.2.2 Directed Acyclic Graphs ..... 14
1.2.3 Mixed Graphs ..... 19
1.3 Conditional Independence and Graphs ..... 23
1.4 More About Graphs ..... 25
1.4.1 Special Properties ..... 25
1.4.2 The igraph package ..... 30
1.4.3 Operations on Graphs in Different Representations ..... 36

### 1.1 Introduction

A graph as a mathematical object may be defined as a pair $\mathcal{G}=(V, E)$, where $V$ is a set of vertices or nodes and $E$ is a set of edges. Each edge is associated with a pair of nodes, its endpoints. Edges may in general be directed, undirected, or bidirected. Graphs are typically visualized by representing nodes by circles or points, and edges by lines, arrows, or bidirected arrows. We use the notation $\alpha-\beta, \alpha \rightarrow \beta$, and $\alpha \leftrightarrow \beta$ to denote edges between $\alpha$ and $\beta$. Graphs are useful in a variety of applications, and a number of packages for working with graphs are available in R .

In statistical applications we are particularly interested in two special graph types: undirected graphs and directed acyclic graphs (often called DAGs).

The gRbase package supplements igraph by implementing some algorithms useful in graphical modelling. gRbase also provides two wrapper functions, ug() and $\operatorname{dag}()$, for easily creating undirected graphs and DAGs represented either as igraph objects or adjacency matrices.

The first sections of this chapter describe some of the most useful functions available when working with graphical models. These come variously from the gRbase and igraph, but it is not usually necessary to know which.

As statistical objects, graphs are used to represent models, with nodes representing model variables (and sometimes model parameters) in such a way that the independence structure of the model can be read directly off the graph. Accordingly, a section of this chapter is devoted to a brief description of the key concept of conditional independence and explains how this is linked to graphs. Throughout the book we shall repeatedly return to this in more detail.

### 1.2 Graphs

Our graphs have a finite node set $V$ and for the most part they are simple graphs in the sense that they have no loops nor multiple edges. Two vertices $\alpha$ and $\beta$ are said to be adjacent, written $\alpha \sim \beta$, if there is an edge between $\alpha$ and $\beta$ in $\mathcal{G}$, i.e. if either $\alpha-\beta$, $\alpha \rightarrow \beta$, or $\alpha \leftrightarrow \beta$.

In this chapter we primarily represent graphs as igraph objects, and except where stated otherwise, the functions we describe operate on these objects.

### 1.2.1 Undirected Graphs

An undirected graph may be created using the $u g()$ function. The graph can be specified using a list of formulas, a single formula or a list of vectors. Thus the following forms are equivalent:

```
library(gRbase)
ug0 <- gRbase::ug(~a:b, ~}\textrm{b}:\textrm{c:d, ~
```

```
ug0 <- gRbase::ug(~a:b + b:c:d + e)
ug0 <- gRbase::ug(~ a*b + b*c*d + e)
ug0 <- gRbase::ug(c("a", "b"), c("b", "c", "d"), "e")
ug0
## IGRAPH 3e18bf5 UN-- 5 4 --
## + attr: name (v/c)
## + edges from 3e18bf5 (vertex names):
## [1] a--b b--c b--d c--d
```



The default size of vertices and their labels is quite small. This is easily changed by setting certain attributes on the graph, see Sect. 1.4.2 for examples. However, to avoid changing these attributes for all the graphs shown in the following we have defined a small plot function myplot(). There are also various facilities for controlling the layout. For example, we may use a layout algorithm called layout.fruchterman.reingold as follows:

```
myplot <- function(x, layout=layout.fruchterman.reingold(x), ...) {
    V(x)$size <- 30
    V(x)$label.cex <- 3
    plot(x, layout=layout, ...)
```

```
    return(invisible())
}
```

The graph ug0i is then displayed with:

```
myplot(ug0)
```



Per default the $u g$ () function returns an igraph object, but the option result="matrix" lead it to return an adjacency matrix instead. For example,

```
ugOi <- gRbase::ug(~a:b + b:c:d + e, result="matrix")
ug0i
## a b c d e
## a 0 1 0 0 0
## b 1 0 1 1 0
## c 0 1 0 1 0
## d 0 1 1 0 0
## e 0 0 0 0 0
```

Different represents of a graph can be obtained by coercion:

```
as(ug0, "matrix")
## a b c d e
## a 0 1 0 0 0
## b 1 0 1 1 0
## c 0 1 0 1 0
## d 0 1 1 0 0
## e 0 0 0 0 0
as(ug0, "dgCMatrix")
## 5 x 5 sparse Matrix of class "dgCMatrix"
## a b c d e
## a . 1 . . .
## b 1 . 1 1.
## c . 1 . 1.
## d . 1 1 . .
## e . . . . .
as(ug0i, "igraph")
## IGRAPH 98eadae UN-- 5 4 --
## + attr: name (v/c), label (v/c)
## + edges from 98eadae (vertex names):
## [1] a--b b--c b--d c--d
```

Edges can be added and deleted using the addEdge() and removeEdge() functions:
\#\# Using gRbase
ug0a <- gRbase::addEdge("a", "c", ug0)
ug0a <- gRbase::removeEdge("c", "d", ug0)

```
## Using igraph
ug0a <- igraph::add_edges(ug0, c("a", "c"))
ug0a <- igraph::delete_edges(ug0, c("c|d"))
```

The nodes and edges of a graph can be retrieved with nodes() and edges() functions.

```
## Using gRbase
gRbase::nodes(ug0)
## [1] "a" "b" "c" "d" "e"
gRbase::edges(ug0) |> str()
## List of 5
## $ a: chr "b"
## $ b: chr [1:3] "a" "c" "d"
## $ c: chr [1:2] "b" "d"
## $ d: chr [1:2] "b" "c"
## $ e: chr(0)
```

\#\# Using igraph
igraph::V(ug0)
\#\# + 5/5 vertices, named, from 3e18bf5:
\#\# [1] a b c d e
igraph::V(ug0) |> attr("names")
\#\# [1] "a" "b" "c" "d" "e"
igraph::E(ug0)
\#\# + 4/4 edges from 3e18bf5 (vertex names):
\#\# [1] a--b b--c b--d c--d
igraph::E(ug0) |> attr("vnames")
\#\# [1] "a|b" "b|c" "b|d" "c|d"
gRbase::maxClique(ug0) \#\# |> str()
\#\# \$maxCliques

```
## $maxCliques[[1]]
## [1] "e"
##
## $maxCliques[[2]]
## [1] "a" "b"
##
## $maxCliques[[3]]
## [1] "b" "c" "d"
gRbase::get_cliques(ug0) |> str()
## List of 3
## $ : chr "e"
## $ : chr [1:2] "a" "b"
## $ : chr [1:3] "b" "c" "d"
## Using igraph
igraph::max_cliques(ug0) |>
    lapply(function(x) attr(x, "names")) |> str()
## List of 3
## $ : chr "e"
## $ : chr [1:2] "a" "b"
## $ : chr [1:3] "b" "c" "d"
```

A path (of length $n$ ) between $\alpha$ and $\beta$ in an undirected graph is a set of vertices $\alpha=\alpha_{0}, \alpha_{1}, \ldots, \alpha_{n}=\beta$ where $\alpha_{i-1}-\alpha_{i}$ for $i=1, \ldots, n$. If a path $n \alpha=\alpha_{0}, \alpha_{1}, \ldots, \alpha_{n}=\beta$ has $\alpha=\beta$ then the path is said to be a cycle of length $n$. nnn A subset $D \subset V$ in an undirected graph is said to separate $A \subset V$ from $B \subset V$ if every path between a vertex in $A$ and a vertex in $B$ contains a vertex from $D$.
gRbase::separates("a", "d", c("b", "c"), ug0)
\#\# [1] TRUE
This shows that $\{b, c\}$ separates $\{a\}$ and $\{d\}$.
The graph $\mathcal{G}_{0}=\left(V_{0}, E_{0}\right)$ is said to be a subgraph of $\mathcal{G}=(V, E)$ if $V_{0} \subseteq V$ and $E_{0} \subseteq E$. For $A \subseteq V$, let $E_{A}$ denote the set of edges in $E$ between vertices in $A$. Then $\mathcal{G}_{A}=\left(A, E_{A}\right)$ is the subgraph induced by $A$. For example

```
ug1 <- gRbase::subGraph(c("b", "c", "d", "e"), ug0)
```

```
ug12 <- igraph::subgraph(ug0, c("b", "c", "d", "e"))
```

$\operatorname{par}(\operatorname{mfrow}=c(1,2), \operatorname{mar}=c(0,0,0,0))$
myplot(ug1); myplot(ug12)


The boundary $\operatorname{bd}(\alpha)=\operatorname{adj}(\alpha)$ is the set of vertices adjacent to $\alpha$ and for undirected graphs the boundary is equal to the set of neighbours ne $(\alpha)$. The closure $\operatorname{cl}(\alpha)$ is $\operatorname{bd}(\alpha) \cup$ $\{\alpha\}$.

```
gRbase::adj(ug0, "c")
## $c
## [1] "b" "d"
gRbase::closure("c", ug0)
## [1] "c" "b" "d"
```


### 1.2.2 Directed Acyclic Graphs

A directed graph as a mathematical object is a pair $\mathcal{G}=(V, E)$ where $V$ is a set of vertices and $E$ is a set of directed edges, normally drawn as arrows. A directed graph is acyclic if it has no directed cycles, that is, cycles with the arrows pointing in the same direction all the way around. A $D A G$ is a directed graph that is acyclic.

A DAG may be created using the $d a g()$ function. The graph can be specified by a list of formulas or by a list of vectors. The following statements are equivalent:

```
dag0 <- gRbase::dag(~a, ~
dag0 <- gRbase::dag(~a + b*a + c*a*b + d*c*e + e*a + g*f)
dag0 <- gRbase::dag(~a + b|a + c|a*b + d|c*e + e|a + g|f)
dag0 <- gRbase::dag("a", c("b", "a"), c("c", "a", "b"), c("d", "c", "e"),
    c("e", "a"), c("g", "f"))
dag0
## IGRAPH 4074b4e DN-- 7 7 --
## + attr: name (v/c)
## + edges from 4074b4e (vertex names):
## [1] a->b a->c b->c c->d e->d a->e f->g
```

Note that ~a means that "a" has no parents while ~d*b*c means that "d" has parents "b" and "c". Instead of "*", a ":" can be used in the specification. If the specified graph contains cycles then dag() returns NULL.

Per default the $\underline{d a g()}$ function returns an igraph object, but the option result="matrix" leads it to return an adjacency matrix instead.


The nodes and edges of a DAG can be retrieved with the nodes() and edges() functions.

```
gRbase::nodes(dag0)
## [1] "a" "b" "c" "d" "e" "f" "g"
gRbase::edges(dag0) |> str()
## List of 7
```

```
## $ a: chr [1:3] "b" "c" "e"
## $ b: chr "c"
## $ c: chr "d"
## $ d: chr(0)
## $ e: chr "d"
## $ f: chr "g"
## $ g: chr(0)
```

Thus edges () gives the children of each node. Alternatively a list of (ordered) pairs can be optained with edgeList()

```
edgeList(dag0) |> str()
## List of 7
## $ : chr [1:2] "a" "b"
## $ : chr [1:2] "a" "c"
## $ : chr [1:2] "a" "e"
## $ : chr [1:2] "b" "c"
## $ : chr [1:2] "c" "d"
## $ : chr [1:2] "e" "d"
## $ : chr [1:2] "f" "g"
```

The upar() function returns a list, with an element for each node together with its parents:

```
vpardag0 <- gRbase::vpar(dag0)
vpardag0 |> str()
## List of 7
## $ a: chr "a"
## $ b: chr [1:2] "b" "a"
## $ c: chr [1:3] "c" "a" "b"
## $ d: chr [1:3] "d" "c" "e"
## $ e: chr [1:2] "e" "a"
## $ f: chr "f"
## $ g: chr [1:2] "g" "f"
vpardag0$c
## [1] "c" "a" "b"
```

A path (of length $n$ ) from $\alpha$ to $\beta$ is a sequence of vertices $\alpha=\alpha_{0}, \ldots, \alpha_{n}=\beta$ such that $\alpha_{i-1} \rightarrow \alpha_{i}$ is an edge in the graph. If there is a path from $\alpha$ to $\beta$ we write $\alpha \mapsto \beta$. The
parents $\mathrm{pa}(\beta)$ of a node $\beta$ are those nodes $\alpha$ for which $\alpha \rightarrow \beta$. The children $\operatorname{ch}(\alpha)$ of a node $\alpha$ are those nodes $\beta$ for which $\alpha \rightarrow \beta$. The ancestors $\operatorname{an}(\beta)$ of a node $\beta$ are the nodes $\alpha$ such that $\alpha \mapsto \beta$. The ancestral set an $(A)$ of a set $A$ is the union of $A$ with its ancestors. The ancestral graph of a set $A$ is the subgraph induced by the ancestral set of $A$.

```
gRbase::parents("d", dag0)
## [1] "c" "e"
gRbase::children("c", dag0)
## [1] "d"
gRbase::ancestralSet(c("b", "e"), dag0)
## [1] "a" "b" "e"
ag <- gRbase::ancestralGraph(c("b", "e"), dag0)
myplot(ag)
```



An important operation on DAGs is to (i) add edges between the parents of each node, and then (ii) replace all directed edges with undirected ones, thus returning an undirected graph. This operation is used in connection with independence interpretations of the DAG, see Sect. 1.3, and is known as moralization. This is implemented by the moralize() function:

```
dag0m <- gRbase::moralize(dag0)
myplot(dag0m)
```



### 1.2.3 Mixed Graphs

Although the primary focus of this book is on undirected graphs and DAGs, it is also useful to consider mixed graphs. These are graphs with at least two types of edges, for example directed and undirected, or directed and bidirected.

A sequence of vertices $v_{1}, v_{2}, \ldots v_{k}, v_{k+1}$ is called a path if for each $i=1 \ldots k$, either $v_{i}-v_{i+1}, v_{i} \leftrightarrow v_{i+1}$ or $v_{i} \rightarrow v_{i+1}$. If $v_{i}-v_{i+1}$ for each $i$ the path is called undirected, if $v_{i} \rightarrow v_{i+1}$ for each $i$ it is called directed, and if $v_{i} \rightarrow v_{i+1}$ for at least one $i$ it is called semi-directed. If $v_{i}=v_{k+1}$ it is called a cycle.

Mixed graphs are represented in the igraph package as directed graphs with multiple edges. In this sense they are not simple. A convenient way of defining them (in lieu of model formulae) is to use adjacency matrices. We can construct such a matrix as follows:

```
adjm <- matrix(c(0, 1, 1, 1,
        1, 0, 0, 1,
        1, 0, 0, 1,
        0, 1, 0, 0), byrow=TRUE, nrow=4)
rownames(adjm) <- colnames(adjm) <- letters[1:4]
adjm
## a b c d
## a 0 1 1 1
## b 1 0 0 1
## c 1 0 0 1
## d 0 1 0 0
```

Note that igraph interprets symmetric entries as double-headed arrows and thus does not distinguish between bidirected and undirected edges. However we can persuade igraph to display undirected instead of bidirected edges:

```
gG1 <- gG2 <- as(adjm, "igraph")
lay <- layout.fruchterman.reingold(gG1)
E(gG2)$arrow.mode <- c(2,0)[1+is.mutual(gG2)]
```

$\operatorname{par}(m f r o w=c(1,2), \operatorname{mar}=c(0,0,0,0))$
myplot(gG1, layout=lay); myplot(gG2, layout=lay)


A chain graph is a mixed graph with no bidirected edges and no semi-directed cycles. Such graphs form a natural generalisation of undirected graphs and DAGs, as we shall see later. The following example is from Frydenberg [1990]:

```
d1 <- matrix(0, 11, 11)
d1[1,2] <- d1[2,1] <- d1[1,3] <- d1[3,1] <- d1[2,4] <- d1[4,2] <-
    d1[5,6] <- d1[6,5] <- 1
d1[9,10] <- d1[10,9] <- d1[7,8] <- d1[8,7] <- d1[3,5] <-
    d1[5,10] <- d1[4,6] <- d1[4,7] <- 1
d1[6,11] <- d1[7,11] <- 1
```

```
rownames(d1) <- colnames(d1) <- letters[1:11]
cG1 <- as(d1, "igraph")
E(cG1)$arrow.mode <- c(2,0)[1+is.mutual(cG1)]
myplot(cG1, layout=layout.fruchterman.reingold)
```



The components of a chain graph $\mathcal{G}$ are the connected components of the graph formed after removing all directed edges from $\mathcal{G}$. All edges within a component are undirected, and all edges between components are directed. Also, all arrows between any two components have the same direction. The graph constructed by identifying its nodes with the components of $\mathcal{G}$, and joining two nodes with an arrow whenever there is an arrow between the corresponding components in $\mathcal{G}$, is a DAG, the so-called component $D A G$ of $\mathcal{G}$, written $\mathcal{G}_{C}$.

The anterior set of a vertex set $S \subseteq V$ is defined in terms of the component DAG. Write the set of components of $\mathcal{G}$ containing $S$ as $S_{c}$. Then the anterior set of $S$ in $\mathcal{G}$ is defined as the union of the components in the ancestral set of $S_{c}$ in $\mathcal{G}_{C}$. The anterior graph of $S \subseteq V$ is the subgraph of $\mathcal{G}$ induced by the anterior set of $S$.

The moralization operation is also important for chain graphs. Similar to DAGs, unmarried parents of the same chain components are joined and directions are then removed.

### 1.3 Conditional Independence and Graphs

The concept of statistical independence is presumably familiar to all readers but that of conditional independence may be less so. Suppose that we have a collection of random variables $\left(X_{v}\right)_{v \in V}$ with a joint density. Let $A, B$ and $C$ be subsets of $V$ and let $X_{A}=\left(X_{v}\right)_{v \in A}$ and similarly for $X_{B}$ and $X_{C}$. Then the statement that $X_{A}$ and $X_{B}$ are conditionally independent given $X_{C}$, written $A \Perp B \mid C$, means that for each possible value of $x_{C}$ of $X_{C}, X_{A}$ and $X_{B}$ are independent in the conditional distribution given $X_{C}=x_{c}$. So if we write $f()$ for a generic density or probability mass function, then one characterization of $A \Perp B \mid C$ is that

$$
f\left(x_{A}, x_{B} \mid x_{C}\right)=f\left(x_{A} \mid x_{C}\right) f\left(x_{B} \mid x_{C}\right) .
$$

An equivalent characterization [Dawid, 1998] is that the joint density of ( $X_{A}, X_{B}, X_{C}$ ) factorizes as

$$
\begin{equation*}
f\left(x_{A}, x_{B}, x_{C}\right)=g\left(x_{A}, x_{C}\right) h\left(x_{B}, x_{C}\right), \tag{1.1}
\end{equation*}
$$

that is, as a product of two functions $g()$ and $h()$, where $g()$ does not depend on $x_{B}$ and $h()$ does not depend on $x_{A}$. This is known as the factorization criterion.

Parametric models for $\left(X_{v}\right)_{v \in V}$ may be thought of as specifying a set of joint densities (one for each admissible set of parameters). These may admit factorisations of the form just described, giving rise to conditional independence relations between the variables. Some models give rise to patterns of conditional independences that can be represented as an undirected graph. More specifically, let $\mathcal{G}=(V, E)$ be an undirected graph with cliques $C_{1}, \ldots C_{k}$. Consider a joint density $f()$ of the variables in $V$. If this admits a factorization of the form

$$
f\left(x_{V}\right)=\prod_{i=1}^{k} g_{i}\left(x_{C_{i}}\right)
$$

for some functions $g_{1}() \ldots g_{k}()$ where $g_{j}()$ depends on $x$ only through $x_{C_{j}}$ then we say that $f()$ factorizes according to $\mathcal{G}$.

If all the densities under a model factorize according to $\mathcal{G}$, then $\mathcal{G}$ encodes the conditional independence structure of the model, through the following result (the global Markov property): whenever sets $A$ and $B$ are separated by a set $C$ in the graph, then $A \Perp B \mid C$ under the model. Thus for example

```
myplot(ug0)
```


gRbase::separates("a", "d", "b", ug0)
\#\# [1] TRUE
shows that under a model with this dependence graph, $a \Perp d \mid b$.
If we want to find out whether two variable sets are marginally independent, we ask whether they are separated by the empty set, which we specify using a character vector of length zero:

```
gRbase::separates("a", "d", character(0), ug0)
## [1] FALSE
```

Model families that admit suitable factorizations are described in later chapters in this book. These include: log-linear models for multivariate discrete data, graphical Gaussian models for multivariate Gaussian data, and mixed interaction models for mixed discrete and continuous data.

Other models give rise to patterns of conditional independences that can be represented by DAGs. These are models for which the variable set $V$ may be ordered in such way that the joint density factorizes as follows

$$
\begin{equation*}
f\left(x_{V}\right)=\prod_{v \in V} f\left(x_{v} \mid x_{\mathrm{pa}(v)}\right) \tag{1.2}
\end{equation*}
$$

for some variable sets $\{\mathrm{pa}(v)\}_{v \in V}$ such that the variables in $\mathrm{pa}(v)$ precede $v$ in the ordering. Again the vertices of the graph represent the random variables, and we can identify the sets $\mathrm{pa}(v)$ with the parents of $v$ in the DAG.

With DAGs, conditional independence is represented by a property called d-separation. That is, whenever sets $A$ and $B$ are $d$-separated by a set $C$ in the graph, then $A \Perp B \mid C$ under the model. The notion of $d$-separation can be defined in various ways, but one characterisation is as follows: $A$ and $B$ are $d$-separated by a set $C$ if and only if they are separated in the graph formed by moralizing the anterior graph of $A \cup B \cup C$.

So we can easily define a function to test this:

```
d_separates <- function(a, b, c, dag_) {
    ##ag <- ancestralGraph(union(union(a, b), c), dag_)
    ag <- ancestralGraph(c(a, b, c), dag_)
    separates(a, b, c, moralize(ag))
}
d_separates("c", "e", "a", dag0)
## [1] TRUE
```

So under dag0 it holds that $c \Perp e \mid a$.
Still other models correspond to patterns of conditional independences that can be represented by a chain graph $\mathcal{G}$. There are several ways to relate Markov properties to chain graphs. Here we describe the so-called LWF Markov properties, associated with Lauritzen, Wermuth and Frydenberg.

For these there are two levels to the factorization requirements. Firstly, the joint density needs to factorize in a way similar to a DAG, i.e.

$$
f\left(x_{V}\right)=\prod_{C \in \mathcal{C}} f\left(x_{C} \mid x_{\mathrm{pa}(C)}\right)
$$

where $\mathcal{C}$ is the set of components of $\mathcal{G}$. In addition, each conditional density $f\left(x_{C} \mid x_{\mathrm{pa}(C)}\right)$ must factorize according to an undirected graph constructed in the following way. First form the subgraph of $\mathcal{G}$ induced by $C \cup \mathrm{pa}(C)$, drop directions, and then complete $\mathrm{pa}(C)$ (that is, add edges between all vertices in $\mathrm{pa}(C))$ ).

For densities which factorize as above, conditional independence is related to a property called c-separation: that is, $A \Perp B \mid C$ whenever sets $A$ and $B$ are $c$-separated by $C$ in the graph. The notion of $c$-separation in chain graphs is similar to that of $d$-separation in DAGs. $A$ and $B$ are $c$-separated by a set $C$ if and only if they are separated in the graph formed by moralizing the anterior graph of $A \cup B \cup C$.

### 1.4 More About Graphs

### 1.4.1 Special Properties

A node in an undirected graph is simplicial if its boundary is complete.

```
gRbase::is.simplicial("b", ug0)
## [1] FALSE
gRbase::simplicialNodes(ug0)
## [1] "a" "c" "d" "e"
```

To obtain the connected components of a graph:

```
gRbase::connComp(ug0) |> str()
## List of 2
## $ : chr [1:4] "a" "b" "c" "d"
## $ : chr "e"
## Using igraph
igraph::components(ug0) |> str()
## List of 3
## $ membership: Named num [1:5] 1 1 1 1 2
## ..- attr(*, "names")= chr [1:5] "a" "b" "c" "d" ...
## $ csize : num [1:2] 4 1
## $ no : int 2
```

If a cycle $\alpha=\alpha_{0}, \alpha_{1}, \ldots, \alpha_{n}=\alpha$ has adjacent elements $\alpha_{i} \sim \alpha_{j}$ with $j \notin\{i-1, i+1\}$ then it is said to have a chord. If it has no chords it is said to be chordless. A graph with no chordless cycles of length $\geq 4$ is called triangulated or chordal:

```
gRbase::is.triangulated(ug0)
## [1] TRUE
igraph::is_chordal(ug0)
## $chordal
## [1] TRUE
##
## $fillin
## NULL
##
## $newgraph
## NULL
```

Triangulated graphs are of special interest for graphical models as they admit closedform maximum likelihood estimates and allow considerable computational simplification by decomposition.

A triple $(A, B, D)$ of non-empty disjoint subsets of $V$ is said to decompose $\mathcal{G}$ into $\mathcal{G}_{A \cup D}$ and $\mathcal{G}_{B \cup D}$ if $V=A \cup B \cup D$ where $D$ is complete and separates $A$ and $B$.

```
gRbase::is.decomposition("a", "d", c("b", "c"), ug0)
## [1] FALSE
```

Note that although $\{d\}$ is complete and separates $\{a\}$ and $\{b, c\}$ in ug0, the condition fails because $V \neq\{a, b, c, d\}$.

A graph is decomposable if it is complete or if it can be decomposed into decomposable subgraphs. A graph is decomposable if and only if it is triangulated.

An ordering of the nodes in a graph is called a perfect ordering if $\operatorname{bd}(i) \cap\{1, \ldots, i-1\}$ is complete for all $i$. Such an ordering exists if and only if the graph is triangulated. If the graph is triangulated, then a perfect ordering can be obtained with the maximum cardinality search (or mcs) algorithm. The mcs() function will produce such an ordering if the graph is triangulated; otherwise it will return NULL.

```
myplot(ug0)
```


a


```
gRbase::mcs(ug0)
## [1] "a" "b" "c" "d" "e"
```

```
igraph::max_cardinality(ug0)
## $alpha
## [1] 5 4 2 3 1
##
## $alpham1
## + 5/5 vertices, named, from 3e18bf5:
## [1] e c d b a
igraph::max_cardinality(ug0)$alpham1 |> attr("names")
## [1] "e" "c" "d" "b" "a"
```

Sometimes it is convenient to have some control over the ordering given to the variables:

```
gRbase::mcs(ug0, root=c("d", "c", "a"))
```

\#\# [1] "d" "c" "b" "a" "e"

Here mcs() tries to follow the ordering given and succeeds for the first two variables but then fails afterwards.

The cliques of a triangulated undirected graph can be ordered as $\left(C_{1}, \ldots, C_{Q}\right)$ to have the running intersection property (also called a RIP ordering). The running intersection property is that $C_{j} \cap\left(C_{1} \cup \ldots \cup C_{j-1}\right) \subset C_{i}$ for some $i<j$ for $j=2, \ldots, Q$. We define the sets $S_{j}=C_{j} \cap\left(C_{1} \cup \ldots \cup C_{j-1}\right)$ and $R_{j}=C_{j} \backslash S_{j}$ with $S_{1}=\emptyset$. The sets $S_{j}$ are called separators as they separate $R_{j}$ from $\left(C_{1} \cup \ldots \cup C_{j-1}\right) \backslash S_{j}$. Any clique $C_{i}$ where $S_{j} \subset C_{i}$ with $i<j$ is a possible parent of $C_{i}$. The $\operatorname{rip()}$ function returns such an ordering if the graph is triangulated (otherwise, it returns list()):

```
gRbase::rip(ug0)
## cliques
## 1 : a b
## 2 : b c d
## 3 : e
## separators
## 1 :
## 2 : b
## 3 :
## parents
## 1 : 0
## 2 : 1
## 3:0
```

If a graph is not triangulated it can be made so by adding extra edges, so called fill-ins, using triangulate():

```
ug2 <- gRbase::ug(~a:b:c + c:d + d:e + a:e)
ug2 <- gRbase::ug(~a:b:c + c:d + d:e + e:f + a:f)
gRbase::is.triangulated(ug2)
## [1] FALSE
igraph::is_chordal(ug2) |> str()
## List of 3
## $ chordal : logi FALSE
## $ fillin : NULL
## $ newgraph: NULL
myplot(ug2)
```


ug3 <- gRbase::triangulate (ug2)
gRbase::is.triangulated(ug3)
\#\# [1] TRUE

```
zzz <- igraph::is_chordal(ug2, fillin=TRUE, newgraph=TRUE)
V(ug2)[zzz$fillin]
## + 4/6 vertices, named, from b261a21:
## [1] d a e a
ug32 <- zzz$newgraph
par(mfrow=c(1,3), mar=c(0,0,0,0))
lay <- layout.fruchterman.reingold(ug2)
myplot(ug2, layout=lay);
myplot(ug3, layout=lay);
myplot(ug32, layout=lay)
```



The Markov blanket of a vertex $v$ in a DAG may be defined as the minimal set that $d$-separates $v$ from the remaining variables. It is easily derived as the set of neighbours to $v$ in the moral graph of $\mathcal{G}$. For example, the Markov blanket of vertex e in dag0 is

```
adj(moralize(dag0), "e")
```

It is easily seen that the Markov blanket of $v$ is the union of $v$ 's parents, $v$ 's children, and the parents of $v$ 's children.

### 1.4.2 The igraph package

It is possible to create igraph objects using the graph.formula() function:

```
ug4 <- graph.formula(a -- b:c, c--b:d, e -- a:d)
ug4
## IGRAPH 605b3d9 UN-- 5 6 --
## + attr: name (v/c)
## + edges from 605b3d9 (vertex names):
## [1] a--b a--c a--e b--c c--d d--e
myplot(ug4)
```



The same graph may be created from scratch as follows:

```
ug4.2 <- graph.empty(n=5, directed=FALSE)
V(ug4.2)$name <- V(ug4.2)$label <- letters[1:5]
ug4.2 <- add.edges(ug4.2, 1+c(0,1, 0,2, 0,4, 1,2, 2,3, 3,4))
ug4.2
## IGRAPH 7b46544 UN-- 5 6 --
## + attr: label (v/c), name (v/c)
## + edges from 7b46544 (vertex names):
## [1] a--b a--c a--e b--c c--d d--e
```

The graph is displayed using the plot() function, with a layout determined using the graphopt method. A variety of layout algorithms are available: type ?layout for an overview. Note that per default the nodes are labelled $0,1, \ldots$ and so forth. We show how to modify this shortly.

As mentioned previously we have created a custom function myplot() which creates somewhat more readable plots:

```
myplot(ug4, layout=layout.graphopt)
```

Objects in igraph graphs are defined in terms of node and edge lists. In addition, they have attributes: these belong to the vertices, the edges or to the graph itself. The following example sets a graph attribute, layout, and two vertex attributes, label and color. These are used when the graph is plotted. The name attribute contains the node labels.

```
ug4$layout <- layout.graphopt(ug4)
V(ug4)$label <- V(ug4)$name
V(ug4)$color <- "red"
V(ug4)[1]$color <- "green"
V(ug4)$size <- 40
V(ug4)$label.cex <- 3
plot(ug4)
```



Note the use of array indices to access the attributes of the individual vertices. Currently, the indices are zero-based, so that $V(\mathrm{ug} 4)$ [1] refers to the second node (B). (This may change). Edges attributes are accessed similarly, using a container structure E(ug4): also here the indices are zero-based (currently).

It is easy to extend igraph objects by defining new attributes. In the following example we define a new vertex attribute, discrete, and use this to color the vertices.

```
ug5 <- set.vertex.attribute(ug4, "discrete", value=c(T, T, F, F, T))
V(ug5)[discrete]$color <- "green"
V(ug5)[!discrete]$color <- "red"
plot(ug5)
```



A useful interactive drawing facility is provided with the tkplot() function. This causes a pop-up window to appear in which the graph can be manually edited. One use of this is to edit the layout of the graph: the new coordinates can be extracted and re-used by the $p$ lot () function. For example
> tkplot(ug4)
2


The $t k p l o t$ () function returns a window id (here 2). While the popup window is open, the current layout can be obtained by passing the window id to the tkplot.getcoords() function, as for example

```
xy <- tkplot.getcoords(2)
plot(g, layout=xy)
```

It is straightforward to reuse layout information with igraph objects. The layout functions when applied to graphs return a matrix of $(x, y)$ coordinates:
layout.fruchterman.reingold(ug4)

| \#\# | $[, 1]$ | $[, 2]$ |
| :--- | ---: | ---: |
| \#\# [1,] | -0.3670 | -0.004236 |
| \#\# [2,] | 0.2393 | 0.979433 |
| \#\# [3,] | 0.6976 | -0.081288 |
| \#\# [4,] | 0.6498 | -1.362600 |
| \#\# [5,] | -0.5042 | -1.279081 |

Most layout algorithms use a random generator to choose an initial configuration. Hence if we set the layout attribute to be a layout function, repeated calls to plot will use different layouts. For example, after

```
ug4$layout <- layout.fruchterman.reingold
```

repeated invocations of plot (ug4) will use different layouts. In contrast, after

```
ug4$layout <- layout.fruchterman.reingold(ug4)
```

the layout will be fixed. The following code fragment illustrates how two graphs with the same vertex set may be plotted using the same layout.

```
ug5 <- gRbase::ug(~A*B*C + B*C*D + D*E)
ug6 <- gRbase::ug(~A*B + B*C + C*D + D*E)
lay.fr <- layout.fruchterman.reingold(ug5)
ug6$layout <- ug5$layout <- lay.fr
V(ug5)$size <- V(ug6)$size <- 50
V(ug5)$label.cex <- V(ug6)$label.cex <- 3
par(mfrow=c(1,2), mar=c(0,0,0,0))
plot(ug5); plot(ug6)
```



An overview of attributes used in plotting can be obtained by typing ?igraph.plotting. A final example illustrates how more complex graphs can be displayed:

```
em1 <- matrix(c(0, 1, 1, 0,
    0, 0, 0, 1,
    1, 0, 0, 1,
    0, 1, 0, 0), nrow=4, byrow=TRUE)
iG <- graph.adjacency(em1)
V(iG)$shape <- c("circle", "square", "circle", "square")
V(iG)$color <- rep(c("red", "green"), 2)
V(iG)$label <- c("A", "B", "C", "D")
E(iG)$arrow.mode <- c(2,0)[1 + is.mutual(iG)]
E(iG)$color <- rep(c("blue", "black"), 3)
E(iG)$curved <- c(T, F, F, F, F, F)
iG$layout <- layout.graphopt(iG)
myplot(iG)
```



### 1.4.3 Operations on Graphs in Different Representations

The gRbase package has a function querygraph() which provides a common interface to the graph operations for undirected graphs and DAGs illustrated above. Moreover, querygraph() works on graphs represented as igraph objects and adjacency matrices. The general syntax is

```
args(querygraph)
## function (object, op, set = NULL, set2 = NULL, set3 = NULL)
## NULL
```

For example, we obtain:

```
ug_ <- gRbase::ug(~a:b + b:c:d + e)
gRbase::separates("a", "d", c("b", "c"), ug_)
## [1] TRUE
gRbase::querygraph(ug_, "separates", "a", "d", c("b", "c"))
## [1] TRUE
gRbase::qgraph(ug_, "separates", "a", "d", c("b", "c"))
## [1] TRUE
```


## Bibliography

Søren Højsgaard, David Edwards, and Steffen Lauritzen. Graphical models with $R$. Springer, 2012.
M. Frydenberg. The chain graph Markov property. Scandinavian Journal of Statistics, 17: 333-353, 1990.
A. P. Dawid. Conditional independence. In Samuel Kotz, Campbell B. Read, and David L.

Banks, editors, Encyclopedia of Statistical Sciences, Update Volume 2, pages 146-155. Wiley-Interscience, New York, 1998.

## Index

addEdge() [graph], 10
dag() [gRbase], 6, 14
edgeList() [gRbase], 16
edges() [graph], 11, 15, 16
graph.formula() [igraph], 30
mcs() [gRbase], 27, 28
moralize() [gRbase], 18
myplot() [gRbase], 31
myplot(), 8
nodes() [graph], 11, 15
plot() [igraph], 31, 33
querygraph() [gRbase], 36
removeEdge() [graph], 10
rip() [gRbase], 28
tkplot() [igraph], 33
tkplot.getcoords() [igraph], 33
triangulate() [gRbase], 29
ug() [gRbase], 6, 9
vpar() [gRbase], 16
adjacency matrix, 6,36
adjacent nodes, 6
ancestors, 16
ancestral graph, 16
ancestral set, 16
anterior graph, 22
anterior set, 22
boundary, 13
c-separation, 25
chain graph components, 22
chain graphs, 21
children, 16
chordal graphs, 26
chordless cycles, 26
closure, 13
component DAG, 22
conditional independence, 23
connected components, 26
cycle, 19
d-separation, 24
DAG, 14
decomposable graphs, 27
decomposition, 27
directed acyclic graph, 14
directed edges, 14
directed graph, 14
directed path, 19
edges, 6
factorizaton criterion, 23
fill-ins, 28
global Markov property, 23
induced subgraph, 12
Markov blanket, 30
maximum cardinality search, 27
mixed graphs, 19
moralization, 18, 22
neighbours, 13
nodes, 6
parents, 16
path, 16, 19
perfect vertex ordering, 27
RIP ordering, 28
running intersection property, 28
semi-directed path, 19
separation, 12
separators, 28
simple graphs, 6
simplicial node, 25
subgraph, 12
triangulated graphs, 26
undirected path, 19
vertices, 6

