## pcalg: A brief overview

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#### 1 General Introduction

In this document, we will give a short overview of the R-package pcalg. The package can be used to analyze dependencies among random variables. The main functions are

- pcAlgo: (also see ?pcAlgo) Use the PC-Algorithm to estimate the skeleton of a DAG. A DAG is a Directed Acyclic Graph and the skeleton of a DAG is the DAG without the arrowheads. In practice, the skeleton will only be a good estimate, if you have way more samples than nodes. If not, you will only get the edges, that show the most obvious "direct" dependencies ("direct" means: The dependence didn't go away, even when conditioning on any subset of the remaining variables). Using the plot option zvalue.lwd you can further qualify the edges by choosing their line width according to their reliability in the statistical tests. I.e., thick lines show reliable dependencies. The value of this function is an object of S4 class "pcAlgo" (see ?pcAlgo-class) on which you can use plot, summary and show.
- corGraph: (see also ?corGraph) Each pair of variables is tested for zero correlation on a given significance level. If a test is rejected, the corresponding edge in the graph is kept. I.e., the graph shows an edge between correlated nodes.

Furthermore, there are some functions that generate random data and perform statistical tests.

## 2 Some examples

In this section, we will show some simple examples with the most important functions. First, we generate a data set that will be used later on:

```
> library(pcalg)
> p <- 10
> n <- 10000
> s <- 0.4
> set.seed(42)
> g <- randomDAG(p, s)
> d <- rmvDAG(n, g)</pre>
```

The DAG that was produced by g <- randomDAG(p,s) is shown in Figure 1.

d contains the datamatrix you would start from in practice. Let's try to estimate the skeleton of the DAG and also compute the correlation graph:

```
> gSkel <- pcAlgo(d, alpha = 0.05)
> gCor <- corGraph(d, alpha = 0.05)</pre>
```

The result of corGraph is a graph object and can be plotted with plot(gCor) (see Figure 2). The result of pcAlgo is a pcAlog object. To get an overview, you can use the summary function:

```
> summary(gSkel)
```

#### > plot(g)

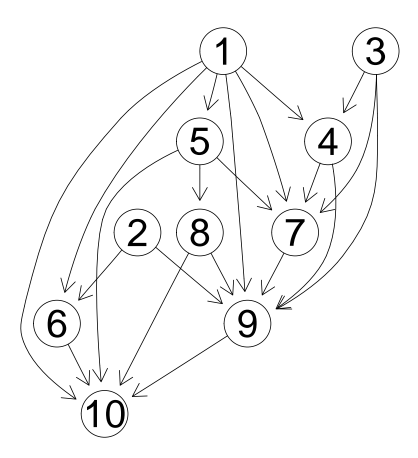


Figure 1: True DAG

#### > plot(gCor)

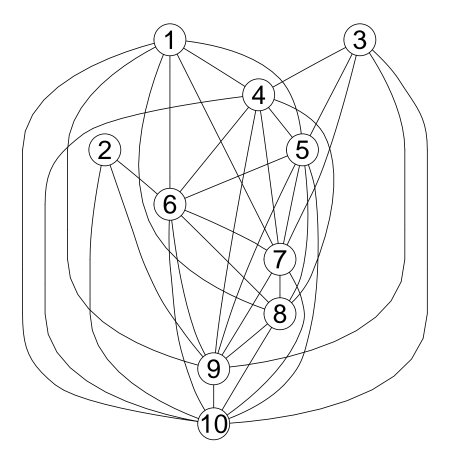


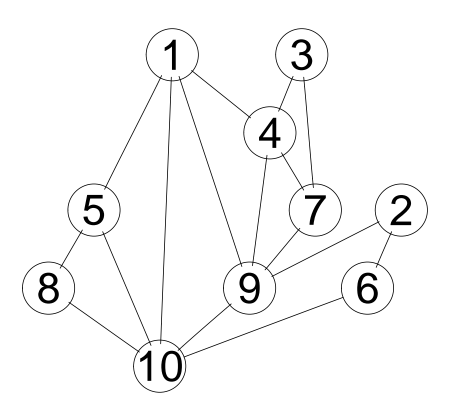
Figure 2: Correlation Graph

```
Object of class 'pcAlgo', from Call:
  pcAlgo(dm = d, alpha = 0.05)

Fitted skeleton based on 10000 observations:
A graphNEL graph with undirected edges
Number of Nodes = 10
Number of Edges = 16

Max. order of algorithm: 5
Number of edgetests from m = 0 up to m = 5 : 81 295 329 277 36 0
Max. number of neighbours: 5 at node(s) 9 10
Avg. number of neighbours: 3.2

> plot(gSkel)
```



pcAlgo(dm = d, alpha = 0.05)

Figure 3: Estimated skeleton

When plotting the pcAlgo object by using plot(gSkel), we obtain Figure 3; choosing the option zvalue.lwd = TRUE, we can adapt the line width of the graph according to the reliability of the corresponding statistical test (see Figure 4).

To compare the estimate of the skeleton with the true skeleton, use

> compareGraphs(gSkel@graph, g)

tpr fpr tdr 0.7619048 0.0000000 1.0000000 > plot(gSkel, zvalue.lwd = TRUE)

# pcAlgo(dm = d, alpha = 0.05)

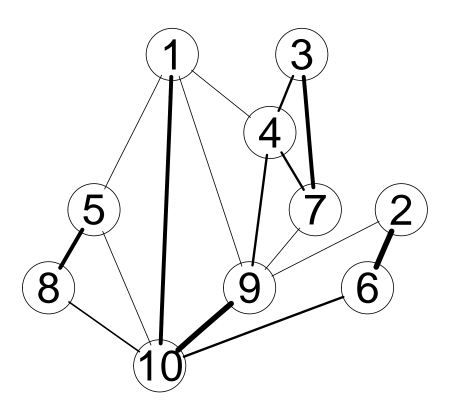


Figure 4: Estimated skeleton using variable line width