# Validação - Somente COVID-19

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#### Variáveis selecionadas

```
## Rows: 76,666
## Columns: 10
           <fct> "(37,73]", "(37,73]", "(73,109]", "(37,73]", "(73,109]", "(~
## $ IDADE
## $ EVOLUCAO
           <fct> 2, 1, 1, 1, 2, 1, 2, 2, 1, 2, 1, 2, 2, 2, 1, 1, 1, 1, 1, 1, 1, ~
## $ RENAL
           <fct> 2, 1, 2, 1, 2, 2, 2, 2, 1, 1, 1, 1, 1, 1, 1, 2, 2, 2, 2, 2, ~
## $ DIABETES
## $ OBESIDADE <fct> 2, 2, 2, 2, 2, 2, 2, 2, 2, 2, 1, 2, 2, 1, 2, 2, 9, 2,~
<fct> 1, 1, 2, 1, 2, 2, 1, 2, 2, 2, 2, 2, 1, 1, 2, 2, 1, 2, 2, 2,~
## $ CARDIOPATI <fct> 2, 2, 2, 1, 1, 2, 2, 1, 1, 1, 1, 1, 1, 1, 2, 2, 1, 1, 1, 1, -
## $ SUPORT_VEN <fct> 2, 2, 2, 9, 3, 3, 2, 2, 3, 2, 2, 3, 1, 1, 3, 3, 2, 3, 9, 3,~
```

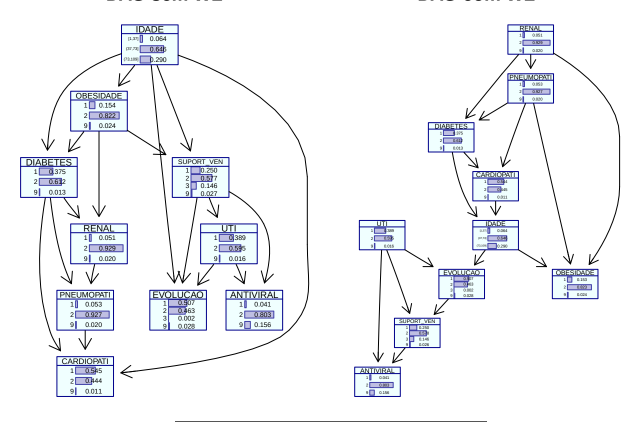
# Imposição de estrutura com arcos que fazem sentido clínico (White list)

# DAG (*Directed Acyclic Graphic*) usando ou não *whitelisting* no algoritmo de aprendizado da estrutura da rede causal

```
par(mfrow = c(1, 2))
graphviz.plot(bn2, shape='rectangle', highlight = NULL, main = 'DAG sem WL')
graphviz.plot(bn1, shape='rectangle', highlight = list(arcs = w1), main = 'DAG com imposição de uma WL'
                                            DAG com imposição de uma WL
          DAG sem WL
                      RENAL
                                                             IDADE
                   PNEUMOPATI
                                                    OBESIDADE
         DIABETES
                                             DIABETES
                                                                   SUPORT_VEN
              CARDIOPATI
              IDADE
   UTI
                                                    RENAL
                                                                        UTI
           EVOLUCAO
                           OBESIDADE
                                                              EVOLUCAO
                                                 PNEUMOPATI
                                                                          ANTIVIRAL
     SUPORT_VEN
                                                 CARDIOPATI
ANTIVIRAL
fitted.1 <- bn.fit(bn1, s1)</pre>
fitted.2 <- bn.fit(bn2, s1)</pre>
par(mfrow=c(1,2))
graphviz.chart(fitted.1,type = "barprob", col = "darkblue", bg = "azure", bar.col = "darkblue", main =
graphviz.chart(fitted.2, type = "barprob", col = "darkblue", bg = "azure", bar.col = "darkblue", main
```

### DAG sem WL

### DAG com WL



# Bootstrapping com WL

```
boots.trap <- 300
str.diff = suppressMessages(boot.strength(s1, R = boots.trap, algorithm = "mmhc"))
cat(paste('Threshold: ', attr(str.diff, "threshold")))
## Threshold: 0.72333333333333
avg.diff = averaged.network(str.diff)
strength.plot(avg.diff, str.diff, shape = "rectangle", main = paste("Iterações = ", boots.trap))</pre>
```

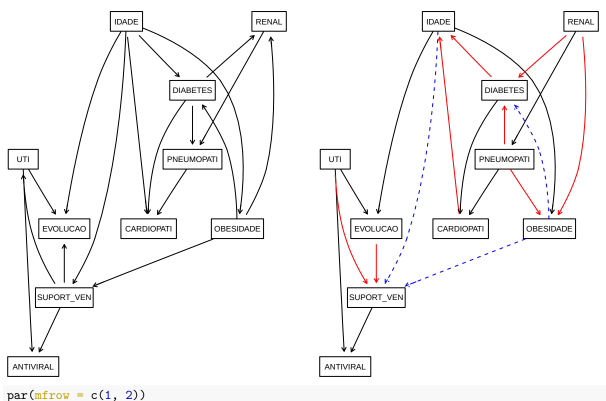
# Iterações = 300 RENAL UTI EVOLUCAO OBESIDADE DIABETES SUPORT\_VEN CARDIOPATI ANTIVIRAL

How can we compare the averaged network (avg.diff) with the network we originally learned in from all the data? The most qualitative way is to plot the two networks side by side, with the nodes in the same positions, and highlight the arcs that appear in one network and not in the other, or that appear with different directions.

```
par(mfrow = c(1, 2))
graphviz.compare(bn1, bn2, shape = "rectangle", main = c("DAG único sem WL", "DAG único com WL"))
```

# DAG único sem WL

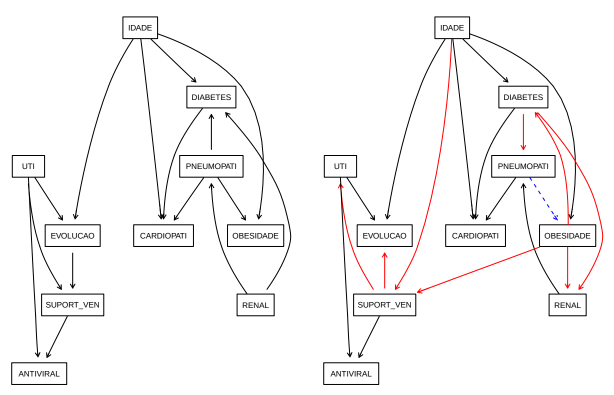
# DAG único com WL



graphviz.compare(avg.diff, bn1, shape = "rectangle", main = c("DAG médio sem WL", "DAG médio com WL"))

# DAG médio sem WL

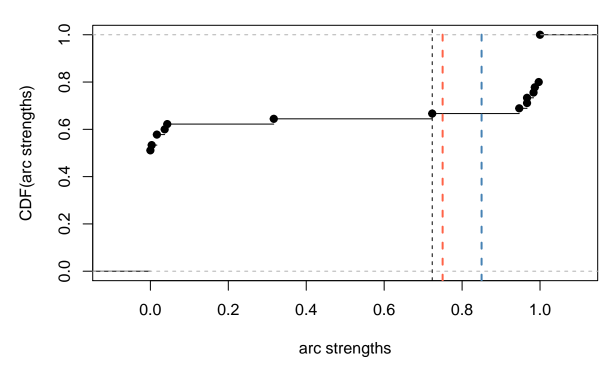
# DAG médio com WL



It is also a good idea to look at the threshold with respect to the distribution of the arc strengths

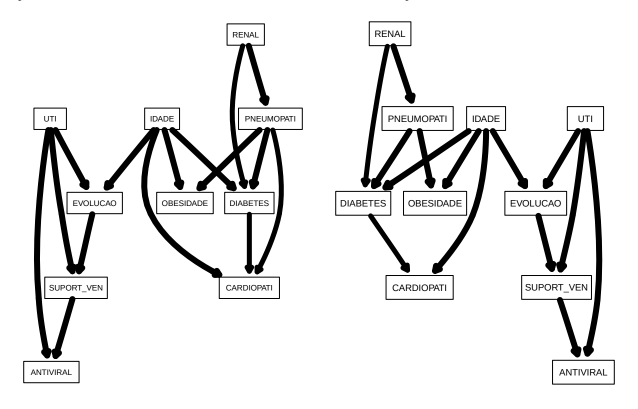
```
plot(str.diff)
abline(v = 0.75, col = "tomato", lty = 2, lwd = 2)
abline(v = 0.85, col = "steelblue", lty = 2, lwd = 2)
```

## threshold = 0.723



The simpler network we obtain by setting threshold = 0.8 in averaged.network() is shown below; it is certainly easier to reason with from a qualitative point of view.

ações = 300 Thr = 0.723333333333333 Iterações = 100 Thr = 0.95



# Markov Blanket da variável evolução

```
cat (mb(x = avg.diff, node = 'EVOLUCAO'))

## IDADE UTI SUPORT_VEN

##
cat (mb(x = avg.simpler, node = 'EVOLUCAO'))

## IDADE UTI SUPORT_VEN

#save the currente workspace
save(list = ls(all.names = TRUE), file = "kk_05_sem_neuro_90pc_300it.RData", envir = .GlobalEnv)
```