## Advanced Machine Learning - Lab 01

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## 1 Hill Climbing

**Task:** Show that multiple runs of the hill-climbing algorithm can return non-equivalent Bayesian network (BN) structures. Explain why this happens. Use the Asia dataset which is included in the bnlearn package. To load the data, run data("asia").

Hint: Check the function hc in the bnlearn package. Note that you can specify the initial structure, the number of random restarts, the score, and the equivalent sample size (a.k.a imaginary sample size) in the BDeu score. You may want to use these options to answer the question. You may also want to use the functions plot, arcs, vstructs, cpdag and all.equal.

Answer: First we will load the data set and have a look at it.

#### head(asia)

```
Α
         S
## 1 no yes no no yes
                           no yes
## 2 no yes no no
                           no
## 3 no no yes no
                   no yes yes yes
## 4 no no
            no no yes
                           no yes
## 5 no no no no
                   no
                           no yes
                       no
## 6 no yes
            no no
                   no
                           no yes
```

To show that different runs of the hill climbing algorithm yield in different results, we will use two different approaches.

- 1. We will create one random graph, and then run the hc() function multiple times, also utilising the different options offered.
- 2. We will use different random graphs which should give us different results.

The hill climbing algorithm is basically finding the next global maximum (or minimum), which is a really easy approach but will also not give us good results. As different measurement metrics will obviously give us different results, we will not further investigate this option.

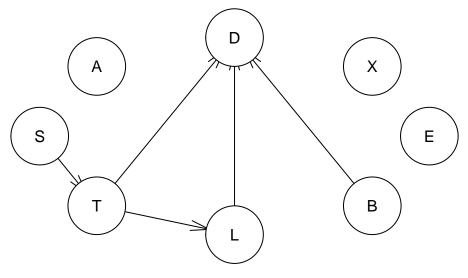
#### 1.1 Same Graph

So we will first create a random graph using the dataset.

```
bayes_net1 = random.graph(colnames(asia))
bayes_net1
```

```
##
##
     Random/Generated Bayesian network
##
     model:
##
      [A] [S] [B] [E] [X] [T|S] [L|T] [D|T:L:B]
##
                                               8
##
     nodes:
##
                                               5
     arcs:
##
       undirected arcs:
                                               0
##
                                               5
       directed arcs:
##
     average markov blanket size:
                                               1.75
                                               1.25
##
     average neighbourhood size:
##
     average branching factor:
                                               0.62
##
##
     generation algorithm:
                                               Full Ordering
                                               0.2857143
     arc sampling probability:
##
score(bayes_net1, asia)
```

```
## [1] -13931.42
plot(bayes_net1)
```



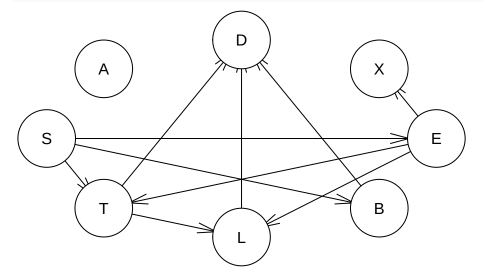
Now we will use the hill climbing algorithm to optimise our random graph

```
bayes_net1_hc = hc(asia, start = bayes_net1)
bayes_net1_hc
```

##

```
##
     Bayesian network learned via Score-based methods
##
##
     model:
##
      [A] [S] [B|S] [E|S] [T|S:E] [X|E] [L|T:E] [D|T:L:B]
##
     nodes:
##
     arcs:
                                               10
##
       undirected arcs:
                                               0
                                               10
##
       directed arcs:
##
     average markov blanket size:
                                               3.00
##
                                               2.50
     average neighbourhood size:
##
     average branching factor:
                                               1.25
##
##
     learning algorithm:
                                               Hill-Climbing
##
                                               BIC (disc.)
     score:
##
     penalization coefficient:
                                               4.258597
##
     tests used in the learning procedure:
                                               81
                                               TRUE
     optimized:
score(bayes_net1_hc, asia)
```

# ## [1] -11133.45 plot(bayes\_net1\_hc)



We observe that the network has changed and has a better score. Another call.

```
bayes_net1_hc = hc(asia, start = bayes_net1_hc)
score(bayes_net1_hc, asia)
```

## [1] -11133.45

This time the score has not been updated, as we're *stuck* in the the same local optima.

What we can do is use the parameters restart and and perturb. According to the documentation and the source code the parameters perturb specifies how many edges or nodes (in this terminology arcs) to change after each restart, the default is one. The parameter restart specifies how often to do that. This usually results in slightly better graphs. If perturb = 1 all combinations are quickly investigated and a different for restart does not really matter that much any more:

```
bayes_net1_hc_restart_low = hc(asia, start = bayes_net1_hc, restart = 10)
bayes_net1_hc_restart_high = hc(asia, start = bayes_net1_hc, restart = 100)
bayes_net1_hc_restart_vhigh = hc(asia, start = bayes_net1_hc, restart = 1000)

score(bayes_net1_hc_restart_low, asia)

## [1] -11133.45

score(bayes_net1_hc_restart_high, asia)

## [1] -11119.26

score(bayes_net1_hc_restart_vhigh, asia)

## [1] -11119.26
```

#### 1.2 Different Graphs

If we use different random graphs in the beginning, the hill climbing algorithm will find different local optima.

```
bayes_net2_1_hc = hc(asia, start = random.graph(colnames(asia)))
bayes_net2_2_hc = hc(asia, start = random.graph(colnames(asia)))
score(bayes_net2_1_hc, asia)
## [1] -11120.65
score(bayes_net2_2_hc, asia)
```

```
## [1] -11131.01
```

As we can see, they are different. If we specify the **restart** parameter again, it will eventually find the same optima, at least for this case, as this is a very small problem.

```
bayes_net2_1_hc = hc(asia, start = random.graph(colnames(asia)), restart = 100)
bayes_net2_2_hc = hc(asia, start = random.graph(colnames(asia)), restart = 100)
score(bayes_net2_1_hc, asia)
### [1] -11107 29
```

```
## [1] -11107.29
score(bayes_net2_2_hc, asia)
```

## [1] -11107.29

#### 2 Inference

Task: Learn a BN from 80 % of the Asia dataset. The dataset is included in the bnlearn package. To load the data, run data("asia"). Learn both the structure and the parameters. Use any learning algorithm and settings that you consider appropriate. Use the BN learned to classify the remaining 20 % of the Asia dataset in two classes: S = yes and S = no. In other words, compute the posterior probability distribution of S for each case and classify it in the most likely class. To do so, you have to use exact or approximate inference with the help of the bnlearn and gRain packages, i.e. you are not allowed to use functions such as predict. Report the confusion matrix, i.e. true/false positives/negatives. Compare your results with those of the true Asia BN, which can be obtained by running dag = model2network("[A][S][T|A][L|S][B|S][D|B:E][E|T:L][X|E]")

Hint: You already know the Lauritzen-Spiegelhalter algorithm for inference in BNs, which is an exact algorithm. There are also approximate algorithms for when the exact ones are too demanding computationally. For exact inference, you may need the functions bn.fit and as.grain from the bnlearn package, and the functions compile, setFinding and querygrain from the package gRain. For approximate inference, you may need the functions prop.table, table and cpdist from the bnlearn package. When you try to load the package gRain, you will get an error as the package RBGL cannot be found. You have to install this package by running the following two commands (answer no to any offer to update packages):

source("https://bioconductor.org/biocLite.R") biocLite("RBGL")

**Answer:** First we split into training and test, by sampling randomly.

```
# Exercise 2)
asia = asia %>% mutate(id = row_number())
train = asia %>% sample_frac(.8)
test = anti_join(asia, train, by = 'id')
train = select(train, -id)
test = select(test, -id)
testX = select(test, -S)
testY = test %>% select(S)
# Helper Functions
## Training the network
train_bayesian_network = function(structure = NULL,
                             data = train,
                             learning_algorithm = iamb) {
 # Network
 if (is.null(structure)) {
   bayesian_network = iamb(data)
 else {
   bayesian_network = structure
 # Parameters
 bayesian_network_fit = bn.fit(bayesian_network, data)
 bayesian_network_grain = as.grain(bayesian_network_fit)
 return(list(bn_grain=bayesian_network_grain,
            bn_fit=bayesian_network_fit,
            bn_structure=bayesian_network))
}
## Predicting with the network
predict_bayesian_network = function(bayesian_network,
                              testX = testX,
                              testY_ = testY,
                              markov_blanket = NULL) {
```

```
# Parallel setup
  no_cores = detectCores()
  cl = makeCluster(no cores)
  clusterExport(cl, list("querygrain", "setEvidence", "bayesian_network"),
                envir=environment())
  # predict for each data point of the test data
  if (!is.null(markov_blanket)) {
    # When predicting, only use the nodes from the Markow blanket
   res = t(parApply(cl, testX, 1, FUN = function(x) {
      return(querygrain(setEvidence(bayesian_network, markov_blanket,
                                    x[markov_blanket]))$S)
   }))
  }
  else {
    # Predict using all nodes
   res = t(parApply(cl, testX_, 1, FUN = function(x) {
     return(querygrain(setEvidence(bayesian_network, names(x), x))$S)
   }))
  }
  # classify
  pred = parApply(cl, res, 1, FUN = function(x) {
   if (x[2] > 0.5) return("yes")
   return("no")
  })
  # Factorize
  testY_factor = testY_[,1]
  pred_factor = factor(pred)
  # Call to a library to calculate interesting metrics
  confusion_matrix = confusionMatrix(pred_factor,
                                     testY_factor, mode="everything")
  stopCluster(cl)
  return(list(res=res, pred=pred, cf=confusion_matrix))
}
```

First we learn the network structure and then we fit the parameters. Afterwards we transform the object into a grain object. We do this by using the custom built function.

Using the given evidence we calculate the posterior probability for each case. We then take the posterior probability to classify the test data. As a comparison we also take the built in predict() function.

```
predicate_2 = predict_bayesian_network(bn_2$bn_grain)

# Metrics
predicate_2$cf
```

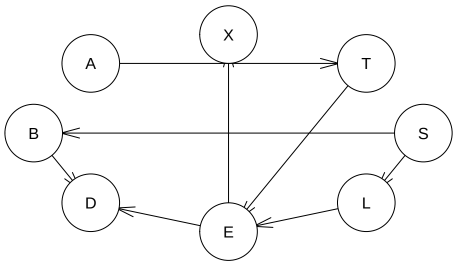
```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction no yes
##
          no 360 130
##
          yes 155 355
##
##
                  Accuracy: 0.715
##
                    95% CI: (0.6859, 0.7428)
##
       No Information Rate: 0.515
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.4303
##
##
    Mcnemar's Test P-Value: 0.1551
##
##
               Sensitivity: 0.6990
##
               Specificity: 0.7320
##
            Pos Pred Value: 0.7347
            Neg Pred Value: 0.6961
##
##
                 Precision: 0.7347
##
                    Recall: 0.6990
##
                        F1: 0.7164
##
                Prevalence: 0.5150
##
            Detection Rate: 0.3600
##
      Detection Prevalence: 0.4900
##
         Balanced Accuracy: 0.7155
##
##
          'Positive' Class : no
##
# We will use predict to compare our own inference with the build in version
predicate_2_sol = predict(bn_2$bn_fit, "S", data = test, method="bayes-lw")
# Factorize
testY_factor = testY[,1]
confusionMatrix(predicate_2_sol, testY_factor, mode="everything")
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction no yes
          no 361 131
##
##
          yes 154 354
##
##
                  Accuracy: 0.715
##
                    95% CI: (0.6859, 0.7428)
       No Information Rate: 0.515
##
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.4303
##
##
    Mcnemar's Test P-Value: 0.1925
##
```

```
##
               Sensitivity: 0.7010
##
               Specificity: 0.7299
##
            Pos Pred Value: 0.7337
            Neg Pred Value: 0.6969
##
##
                 Precision: 0.7337
##
                    Recall : 0.7010
##
                        F1: 0.7170
                Prevalence: 0.5150
##
##
            Detection Rate: 0.3610
##
      Detection Prevalence : 0.4920
##
         Balanced Accuracy: 0.7154
##
##
          'Positive' Class : no
##
```

As we can see, we observe the very similar results.

The true Bayesian network is given by the following:

```
# Define true network, train and use for prediction
dag = model2network("[A][S][T|A][L|S][B|S][D|B:E][E|T:L][X|E]")
plot(dag)
```



```
bn_2_true = train_bayesian_network(dag)
predicate_2_true = predict_bayesian_network(bn_2_true$bn_grain)
predicate_2_true$cf
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction no yes
##
          no 357 115
##
          yes 158 370
##
##
                  Accuracy: 0.727
##
                    95% CI: (0.6982, 0.7544)
##
       No Information Rate: 0.515
       P-Value [Acc > NIR] : < 2e-16
```

```
##
##
                     Kappa: 0.4549
##
    Mcnemar's Test P-Value : 0.01102
##
##
##
               Sensitivity: 0.6932
               Specificity: 0.7629
##
            Pos Pred Value: 0.7564
##
##
            Neg Pred Value: 0.7008
##
                 Precision: 0.7564
##
                    Recall: 0.6932
                        F1: 0.7234
##
##
                Prevalence: 0.5150
            Detection Rate: 0.3570
##
##
      Detection Prevalence: 0.4720
##
         Balanced Accuracy: 0.7280
##
##
          'Positive' Class : no
##
```

We see that the true network just has a slightly better F1 score compared to the trained versions from the data.

#### 3 Markov Blanket

**Task:** In the previous exercise, you classified the variable S given observations for all the rest of the variables. Now, you are asked to classify S given observations only for the so-called Markov blanket of S, i.e. its parents plus its children plus the parents of its children minus S itself. Report again the confusion matrix.

**Hint:** You may want to use the function mb from the bnlearn package.

**Answer:** First we will extract the *Markov Blanket* using the function mb(). The *Markov Blanket* are all nodes that shield the node in question from the rest of the network. Then we apply the same procedure as before and look at the results.

```
# Exercise 3)
predicate_3_mb =
 predict_bayesian_network(bn_2$bn_grain,
                  markov_blanket = mb(bn_2$bn_structure, "S"))
predicate_3_mb$cf
## Confusion Matrix and Statistics
##
##
        Reference
##
 Prediction no yes
##
      no 360 130
##
      ves 155 355
##
            Accuracy: 0.715
##
             95% CI: (0.6859, 0.7428)
##
##
    No Information Rate: 0.515
```

```
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.4303
##
##
   Mcnemar's Test P-Value: 0.1551
##
               Sensitivity: 0.6990
##
               Specificity: 0.7320
##
##
            Pos Pred Value: 0.7347
##
            Neg Pred Value: 0.6961
##
                 Precision: 0.7347
                    Recall: 0.6990
##
##
                        F1: 0.7164
##
                Prevalence: 0.5150
##
            Detection Rate: 0.3600
##
      Detection Prevalence: 0.4900
##
         Balanced Accuracy: 0.7155
##
##
          'Positive' Class : no
##
```

As we can see the F1 score (and so the results) remain unchanged when only considering the markov blanket.

### 4 Naive Bayes

**Task:** Repeat the exercise (2) using a naive Bayes classifier, i.e. the predictive variables are independent given the class variable. See p. 380 in Bishop's book or Wikipedia for more information on the naive Bayes classifier. Model the naive Bayes classifier as a BN. You have to create the BN by hand, i.e. you are not allowed to use the function naive.bayes from the bnlearn package.

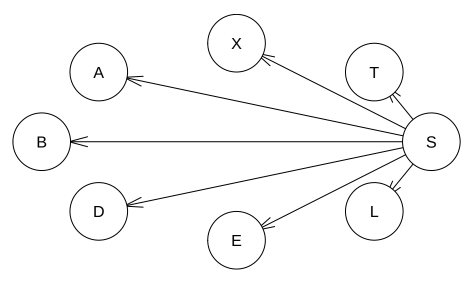
Hint: Check http://www.bnlearn.com/examples/dag/ to see how to create a BN by hand.

**Answer:** The Naive Bayes classifier assumes independence between all the predictive variables. Therefore the classifier is given by

$$p(S|x_i) \propto p(S) \prod_{i=1}^{N} p(x_i|S)$$

 $x_i$  are the features, therefore our Bayesian network structure is based on the following formula:

$$p(S) \propto p(S)p(A|S)p(T|S)p(L|S)p(B|S)p(E|S)p(X|S)p(D|S)$$



As we can see, according to our model there exists no dependence between the features. Let's train the parameters and look at the results.

```
##
##
             Reference
##
  Prediction no yes
##
          no 380 174
##
          yes 135 311
##
                  Accuracy: 0.691
##
                    95% CI : (0.6613, 0.7195)
##
##
       No Information Rate: 0.515
##
       P-Value [Acc > NIR] : < 2e-16
##
##
                     Kappa: 0.38
##
##
   Mcnemar's Test P-Value: 0.03064
##
##
               Sensitivity: 0.7379
##
               Specificity: 0.6412
            Pos Pred Value: 0.6859
##
            Neg Pred Value: 0.6973
##
##
                 Precision: 0.6859
                    Recall: 0.7379
##
##
                        F1: 0.7109
                Prevalence: 0.5150
##
            Detection Rate: 0.3800
##
##
      Detection Prevalence: 0.5540
```

Balanced Accuracy: 0.6896

##

```
##
## 'Positive' Class : no
##
```

## 5 Explanation of Different Results

Task: Explain why you obtain the same or different results in the exercises (2-4).

**Answer:** As the observations of the Markov Blanket make S independent from all other variables, we should and we do observe the exact same result in 2) and 3) (for the same trained network). For the Naive Bayes classifier we see, that we get a slightly worse F1 score, which is to be expected, as our assumption does not hold, when we compare with the true network given. Despite this fact, the Naive Base classifier does quite well.

#### 6 Source Code

```
knitr::opts_chunk$set(echo = TRUE)
library(bnlearn)
library(tibble)
library(dplyr)
library(gRain)
library(caret)
library(parallel)
#if (!requireNamespace("BiocManager", quietly = TRUE))
    install.packages("BiocManager")
#BiocManager::install("RBGL")
set.seed(42)
# Exercise 1)
as_tibble("asia")
head(asia)
bayes_net1 = random.graph(colnames(asia))
bayes_net1
score(bayes_net1, asia)
plot(bayes_net1)
bayes_net1_hc = hc(asia, start = bayes_net1)
bayes_net1_hc
score(bayes_net1_hc, asia)
plot(bayes_net1_hc)
bayes_net1_hc = hc(asia, start = bayes_net1_hc)
```

```
score(bayes_net1_hc, asia)
bayes_net1_hc_restart_low = hc(asia, start = bayes_net1_hc, restart = 10)
bayes_net1_hc_restart_high = hc(asia, start = bayes_net1_hc, restart = 100)
bayes_net1_hc_restart_vhigh = hc(asia, start = bayes_net1_hc, restart = 1000)
score(bayes net1 hc restart low, asia)
score(bayes_net1_hc_restart_high, asia)
score(bayes_net1_hc_restart_vhigh, asia)
bayes_net2_1_hc = hc(asia, start = random.graph(colnames(asia)))
bayes_net2_2_hc = hc(asia, start = random.graph(colnames(asia)))
score(bayes_net2_1_hc, asia)
score(bayes_net2_2_hc, asia)
bayes_net2_1_hc = hc(asia, start = random.graph(colnames(asia)), restart = 100)
bayes_net2_2_hc = hc(asia, start = random.graph(colnames(asia)), restart = 100)
score(bayes net2 1 hc, asia)
score(bayes_net2_2_hc, asia)
# Exercise 2)
asia = asia %>% mutate(id = row_number())
train = asia %>% sample_frac(.8)
test = anti_join(asia, train, by = 'id')
train = select(train, -id)
test = select(test, -id)
testX = select(test, -S)
testY = test %>% select(S)
# Helper Functions
## Training the network
train_bayesian_network = function(structure = NULL,
                              data = train,
                              learning_algorithm = iamb) {
 # Network
 if (is.null(structure)) {
   bayesian_network = iamb(data)
 }
 else {
   bayesian_network = structure
```

```
# Parameters
  bayesian_network_fit = bn.fit(bayesian_network, data)
  bayesian_network_grain = as.grain(bayesian_network_fit)
 return(list(bn_grain=bayesian_network_grain,
              bn fit=bayesian network fit,
              bn_structure=bayesian_network))
}
## Predicting with the network
predict_bayesian_network = function(bayesian_network,
                                    testX_ = testX,
                                    testY_ = testY,
                                    markov_blanket = NULL) {
  # Parallel setup
  no_cores = detectCores()
  cl = makeCluster(no_cores)
  clusterExport(cl, list("querygrain", "setEvidence", "bayesian_network"),
                envir=environment())
  # predict for each data point of the test data
  if (!is.null(markov blanket)) {
    # When predicting, only use the nodes from the Markow blanket
   res = t(parApply(cl, testX, 1, FUN = function(x) {
     return(querygrain(setEvidence(bayesian_network, markov_blanket,
                                    x[markov_blanket]))$S)
   }))
  else {
    # Predict using all nodes
   res = t(parApply(cl, testX_, 1, FUN = function(x) {
     return(querygrain(setEvidence(bayesian_network, names(x), x))$S)
   }))
  }
  # classify
  pred = parApply(cl, res, 1, FUN = function(x) {
   if (x[2] > 0.5) return("yes")
   return("no")
  })
  # Factorize
  testY_factor = testY_[,1]
  pred_factor = factor(pred)
  # Call to a library to calculate interesting metrics
  confusion_matrix = confusionMatrix(pred_factor,
                                     testY_factor, mode="everything")
  stopCluster(cl)
  return(list(res=res, pred=pred, cf=confusion_matrix))
```

```
}
bn_2 = train_bayesian_network(structure = NULL,
                 data = train,
                 learning_algorithm = iamb)
predicate_2 = predict_bayesian_network(bn_2$bn_grain)
# Metrics
predicate_2$cf
# We will use predict to compare our own inference with the build in version
predicate_2_sol = predict(bn_2$bn_fit, "S", data = test, method="bayes-lw")
# Factorize
testY_factor = testY[,1]
confusionMatrix(predicate_2_sol, testY_factor, mode="everything")
# Define true network, train and use for prediction
dag = model2network("[A][S][T|A][L|S][B|S][D|B:E][E|T:L][X|E]")
plot(dag)
bn_2_true = train_bayesian_network(dag)
predicate_2_true = predict_bayesian_network(bn_2_true$bn_grain)
predicate_2_true$cf
predicate_3_mb =
 predict_bayesian_network(bn_2$bn_grain,
                    markov_blanket = mb(bn_2$bn_structure, "S"))
predicate_3_mb$cf
# Exercise 4)
bn_naive_bayes = model2network("[S][A|S][T|S][L|S][B|S][E|S][X|S][D|S]")
plot(bn_naive_bayes)
```