# Multivariate Analysis and Statistical Learning PC Algorithm implementation

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21/12/2018



#### Theoretical references (1)

- Bayesian Networks can be rappresented as a directed acyclic graph (DAG);
- "acyclic" means that there are no paths starting from a node v that ends with v itself, ∀v ∈ G.

# Theoretical references (2)

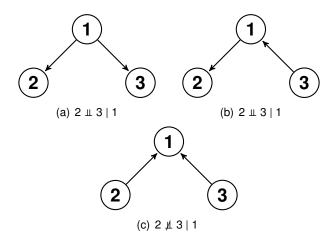
Let G = (V, E) be a DAG relative to a finite set  $X = \{X_v, v \in V\}$  of casual variables, then:

$$\forall u, v \in V \text{ non adjacent } | v \in nd(u) \Rightarrow$$

$$\Rightarrow u \perp v \mid nd(u) - v$$

Where nd(u) is the set of **non-descendant** nodes of u, that are all those nodes u' for which there is no path from u to u'.

# Theoretical references (3)



# **PC-Algorithm**

Given a set of variables with a joint Gaussian probability distribution, it is possible to learn the DAG closer to the sample through the use of **PC-Algorithm**.

It is composed of two sub-functions that solve two different problems:

- The construction of the skeleton from the moral graph;
- 2 The construction of the DAG from a given skeleton.

#### Step 1: read the dataset

- Import pandas library;
- call pandas.read\_csv() function to read dataset;
- define alpha;
- call **get\_skeleton** on dataset and alpha as arguments.

```
import pandas as pd
alpha = .10
dataset = pd.read_csv("marks.dat",sep=",")
(g,sep_set) = get_skeleton(dataset, alpha, dataset.columns)
```

# Step 2: initialization

- Read names of the dataset variables accessing dataset.columns field;
- retrieve the correlation matrix of the given dataset with dataset.corr().values;
- initialize N,n as the number of sampling and the number of variables;
- initialize G as the complete graph of dimension n;
- initalize the separation\_set as a list of list;
- initialize I = 0, stop = false.

```
labels = dataset.columns
corr_matrix = dataset.corr().values
N = dataset.values.shape[0]
n = len(corr_matrix[0])
G = complete(n)
sep_set = [[[] for i in range(n)] for j in range(n)]
stop = False
l = 0
```



#### Step 3: define adj() function

 Define the adj() function in order to get the adjacents of a node in a given graph.

```
def adj(x,G):
    adjacents = list()
    for j in range(0,len(G[x])):
        if G[x][j] == 1:
            adjacents.append(j)
    return adjacents
```

#### Step 4: how many variables are actually dependent?

- set stop condition to true
- retrieve dependent variables: i,j are actually dependent if the adjacence matrix[i][j] is equal to 1
- call the set of dependent variables act\_dep

```
while stop == False and any(G):
    stop = True
    act_dep = []
    for i in range(len(G)):
        for j in range(len(G[i])):
        if G[i][j] == 1:
        act_dep.append((i,j))
```

#### Step 5: variables needed for independence test

- For x,y in act\_dep;
- retrieve the neighbors of x calling the adj() function;
- remove y from the neighbors set;
- if **neighbors** set has dimension ≥ I then
  - if neighbors set has dimension > I go ahead.

#### Step 6: conditional independence test

- Foreach set K of neighbors of dimension I;
- test independence of x and y given K;
- if the p value is greater than alpha:
  - remove the edge x,y setting G[x][y] = 0;
  - set K as the separation\_set[x][y].

```
for K in set(combinations(neighbors, 1)):
    p_value = indep_test(corr_matrix, N, x, y, list(K))
    if p_value >= alpha:
        G[x][y] = 0
        G[y][x] = 0
        sep_set[x][y] = list(K)
        break

l = l + 1
```

#### Step 7: from the skeleton to the CPDAG

- Return G and separation\_set;
- call to\_cpdag(G, separation\_set).

```
(g,sep_set) = get_skeleton(dataset, alpha, dataset.columns)
g = to_cpdag(g,sep_set)
```

#### Step 8: define the getDependents() function

- Define getDependents(adj\_matrix,reqij, reqji);
- this function retrieve all the variables i,j such that:
   adj\_matrix[i][j] == reqij and adj\_matrix[j][i] == reqji.

#### Step 9: CPDAG initialization

- Set the cpdag as the skeleton;
- set dep as the set of variables i,j for which exists an edge from i to j.

```
cpdag = skeleton.tolist()
dep = getDependents(skeleton,1,None)
```

# Step 10: rule "zero" (1)

- For each pair x,y in dep:
- add to allZ all the variables z for which exists an egde from z to y and z is not x;
- if:
   there is no edge between x and z
   there is a separation set between x and z
   there is a separation set between z and x
   y is not in separation set between x and z or in separation set between z and x, then:
- remove the edge from y to x and from z to y.



#### Step 10: rule "zero" (2)

#### Step 11: apply rules

- Using the same logic we apply the known rules 1,2 and 3;
  - **Rule 1:** orient j k into  $j \rightarrow k$  whenever there is an arrow  $i \rightarrow j$  such that i and k are not adjacent;
  - Rule 2: orient i j into  $i \rightarrow j$  whenever there is a chain  $i \rightarrow k \rightarrow j$ ;
  - Rule 3: orient i j into  $i \rightarrow j$  whenever there are two chains  $i \rightarrow k \rightarrow j$  and  $i \rightarrow l \rightarrow j$  such that k and l are nonadjacent.
- Return the resulting cpdag;
- using matplotlib and networkx we are able to plot the resulting cpdag.



Consider this R code:

• it gives:



# 2 1 3

```
> delta/100
Time difference of 0.00781296 secs
> plot(eq.dag, main="PC DAG") #acc
```

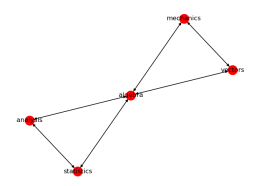


Consider this python code:

```
def test_butterfly_model():
    alpha = .10
    dataset = pd.read_csv("marks.dat",sep=",")
    deltas = 0
    import time
    for i in range(100):
        t0 = time.time()
        (g,sep_set) = get_skeleton(dataset, alpha)
        g = to_cpdag(g,sep_set)
        tf = time.time()
        deltas += (tf-t0)
    print "Elapsed "+str((deltas)/100)+" sec"
    plot(g,dataset.columns)
```

• it gives:





C:\Users\Tommaso\Desktop\MASL\contest>python pcalgorithm.py
Elapsed 0.0074799990654 sec



# Github repository





https://github.com/alexfoglia1/MASL