# Assignment 4

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### Problem 1

### Problem 2

```
# import file
data <- read.table("./data.txt", header = TRUE, sep = " ", stringsAsFactors = FALSE)</pre>
```

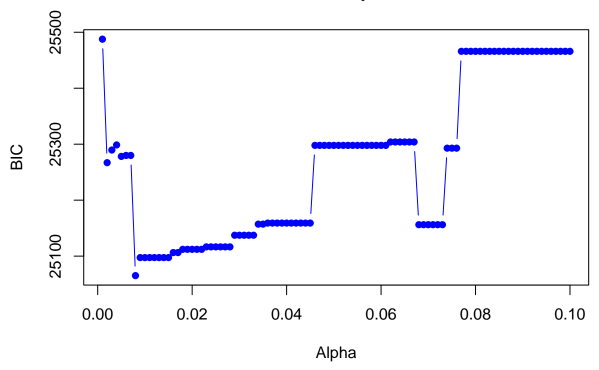
#### PC algorithm

```
# Function to calculate log likelihood
compute_log_likelihood <- function(data, adj_matrix) {</pre>
  sample_cov <- cov(data) # Sample covariance matrix</pre>
  adj_cov <- sample_cov * adj_matrix # Adjust covariance for graph</pre>
  # Regularization for numerical stability
  p <- ncol(data)</pre>
  regularization <- diag(10, p)
  adj_cov <- adj_cov + regularization</pre>
  # Log-likelihood calculation
  tryCatch({
    log_det <- log(det(adj_cov)) # Log determinant</pre>
    inv_cov <- solve(adj_cov)</pre>
                                 # Inverse covariance
    log_likelihood <- -0.5 * nrow(data) * (log_det + sum(diag(inv_cov %*% sample_cov)))</pre>
   return(log_likelihood)
  }, error = function(e) {
    return(NA) # Return NA if numerical issues occur
  })
}
# Function to calculate BIC
compute_bic <- function(pc_fit, data) {</pre>
  adj_matrix <- as(pc_fit@graph, "matrix") # Adjacency matrix</pre>
```

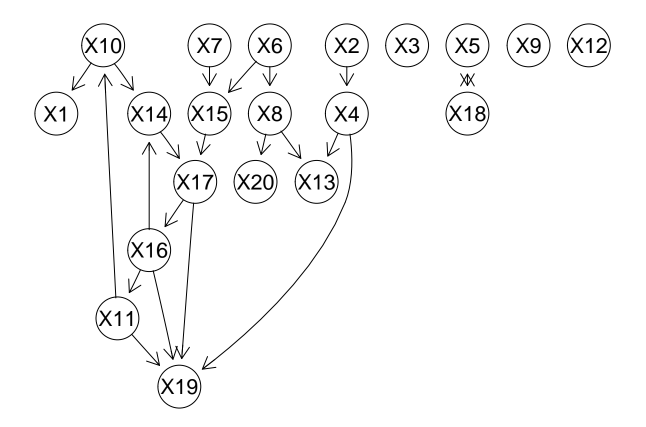
```
num_params <- sum(adj_matrix) # Number of edges (parameters)</pre>
  log_lik <- compute_log_likelihood(data, adj_matrix)</pre>
  if (is.na(log_lik)) return(Inf) # Return infinite BIC for invalid graphs
 n <- nrow(data) # Sample size</pre>
 p <- ncol(data) # Number of variables</pre>
 bic <- -2 * log_lik + log(n) * (num_params + p) # BIC formula
 return(bic)
}
# Generate a sequence of alpha values
alphas \leftarrow seq(0.001, 0.1, by = 0.001)
# Fit PC algorithm and compute BIC for each alpha
bic_values <- sapply(alphas, function(alpha) {</pre>
 pc_fit <- pc(</pre>
    suffStat = list(C = cor(data), n = nrow(data)),
   indepTest = gaussCItest,
   alpha = alpha,
   labels = colnames(data),
   verbose = FALSE
  compute_bic(pc_fit, data)
})
# Find optimal alpha
optimal_alpha <- alphas[which.min(bic_values)]</pre>
cat("Optimal alpha based on BIC:", optimal_alpha, "\n")
## Optimal alpha based on BIC: 0.008
# Plot BIC vs Alpha
```

```
# Plot BIC vs Alpha
plot(
  alphas, bic_values, type = "b", pch = 16, col = "blue",
  xlab = "Alpha", ylab = "BIC",
  main = "BIC vs Alpha"
)
```

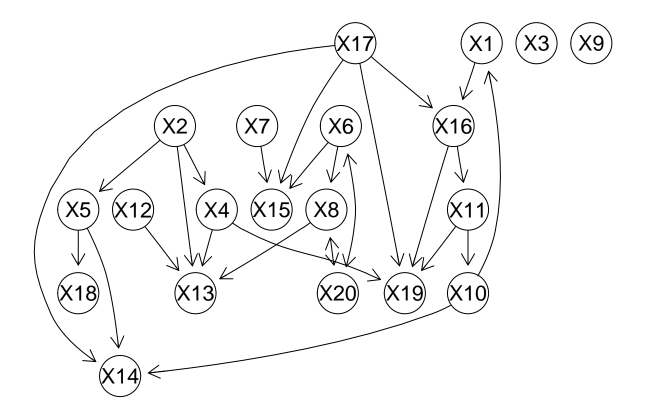
## **BIC** vs Alpha



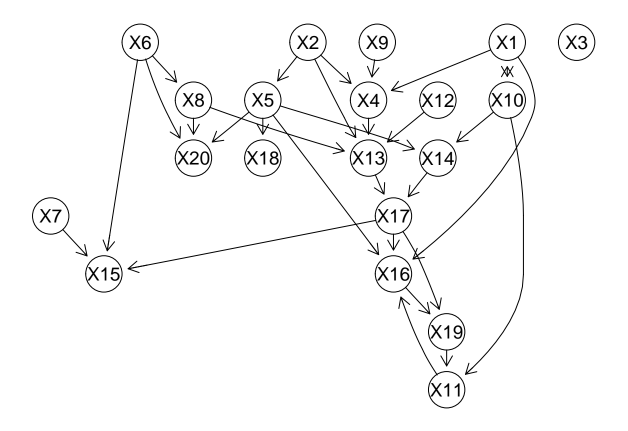
```
# Plot CPDAG
plot(pc.fit.001@graph)
```



plot(pc.fit.008@graph)



plot(pc.fit.10graph)



```
# Check the number of edges
pc.fit.001
## Object of class 'pcAlgo', from Call:
## pc(suffStat = list(C = cor(data), n = nrow(data)), indepTest = gaussCItest,
       alpha = 0.001, labels = colnames(data), verbose = TRUE)
## Number of undirected edges: 1
## Number of directed edges:
## Total number of edges:
                                20
pc.fit.008
## Object of class 'pcAlgo', from Call:
## pc(suffStat = list(C = cor(data), n = nrow(data)), indepTest = gaussCItest,
##
       alpha = 0.008, labels = colnames(data), verbose = TRUE)
## Number of undirected edges: 2
## Number of directed edges:
                                23
## Total number of edges:
                                25
```

#### pc.fit.1

```
## Object of class 'pcAlgo', from Call:
## pc(suffStat = list(C = cor(data), n = nrow(data)), indepTest = gaussCItest,
## alpha = 0.1, labels = colnames(data), verbose = TRUE)
## Number of undirected edges: 1
## Number of directed edges: 28
## Total number of edges: 29
```

I used the PC algorithm to estimate CPDAGs for a range of  $\alpha$  values (0.001 to 0.1) to identify the optimal  $\alpha$  that minimizes the Bayesian Information Criterion (BIC). For each  $\alpha$ , the graph structure  $G(\alpha)$  was used to compute the log-likelihood  $-2\ell(\hat{\Sigma}_{G'},\hat{\mu})$ , where  $\ell(\cdot)$  is the log-likelihood of a Gaussian model based on the graph-constrained covariance matrix. The BIC for each graph was computed as  $-2\ell + \log(n)(k+p)$ , with k representing the number of edges and p the number of variables.

The optimal  $\alpha = 0.008$  was selected as the value minimizing BIC. This model is considered to have an optimal balance between complexity and fit.

Estimated CPDAGs were plotted for specific  $\alpha$  values (0.001, 0.008, 0.1), and the number of edges was examined for each plot. We see that the smaller  $\alpha$  value led to sparser graph with fewer edges, and the larger  $\alpha$  value led to denser graphs with more edges.

#### Graphical lasso method

I will conduct model selection using EBIC, RIC and StARS.

```
# Apply graphical lasso method
out.glasso <- huge(as.matrix(data), method = "glasso")</pre>
```

## Conducting the graphical lasso (glasso) with lossless screening....in progress: 9%Conducting the graphical lasso (glasso)....done.

```
# Perform model selection using EBIC
out.select <- huge.select(out.glasso, criterion = "ebic")</pre>
```

## Conducting extended Bayesian information criterion (ebic) selection....done

```
# Plot the selected graph
plot(out.glasso)
```

parsity vs. Regularization

0.2

0.0

lambda = 0.684

lambda = 0.53



lambda = 0.41



Regularization Parameter

0.2

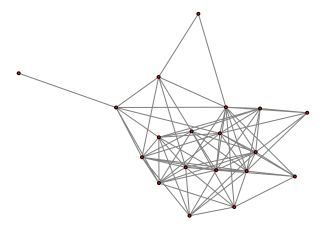
0.5

 $\begin{tabular}{lll} \# \ Extract \ adjacency \ matrix \ of \ the \ selected \ graph \\ adj\_matrix <- \ out.select$refit \end{tabular}$ 

# Check selected lambda
out.select\$opt.lambda

## [1] 0.08835911

# Visualize the selected graph
huge.plot(adj\_matrix)

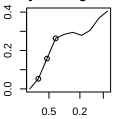


•

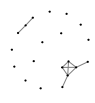
```
# Do the same using RIC
out.select <- huge.select(out.glasso, criterion = "ric")</pre>
```

plot(out.glasso)

parsity vs. Regularization



lambda = 0.684



lambda = 0.53



lambda = 0.41

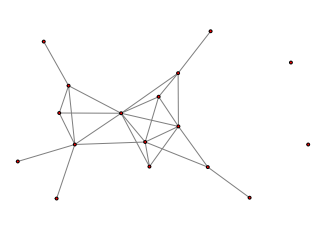


Regularization Parameter

adj\_matrix <- out.select\$refit</pre> out.select\$opt.lambda

## [1] 0.5592347

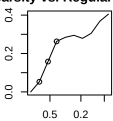
huge.plot(adj\_matrix)



```
# Do the same using STARS
out.select <- huge.select(out.glasso, criterion = "stars")</pre>
```

## Conducting Subsampling....in progress:5% Conducting Subsampling....in progress:10% Conducting Subsampling....inplot(out.glasso)

parsity vs. Regularization



Regularization Parameter

lambda = 0.684



lambda = 0.53



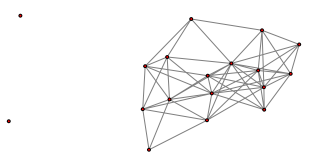
lambda = 0.41



adj\_matrix <- out.select\$refit</pre> out.select\$opt.lambda

## [1] 0.2458645

huge.plot(adj\_matrix)



First, I estimated a sequence of graphs corresponding to different regularization parameters ( $\lambda$ ). Larger  $\lambda$  values produced sparser graphs with many isolated edges, while smaller  $\lambda$  values led to highly connected graphs. Subsequently, three model selection criteria were applied to identify the optimal graph that balances fit and sparsity. EBIC balances log-likelihood and penalizes model complexity in high-dimensional data ( $\lambda=0.0884$ ). RIC uses a simpler penalty that generally favors fewer edges, resulting in a moderate sparsity ( $\lambda=0.4997$ ). StARS prioritizes the stability of edges across subsampled data ( $\lambda=0.1904$ ). We need to choose the criteria based on the nature of data and underlying relationships in the data.