Project 8

Santiago Castro Dau, June Monge, Rachita Kumar, Sarah Lötscher

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Problem 20: Classical NEMs

1. For each model, construct the transitive closure (by adding edges) and define the corresponding adjacency matrices Φ and Θ , which represent the signalling pathways and the E-gene attachments. Determine the corresponding expected effect patterns (F).

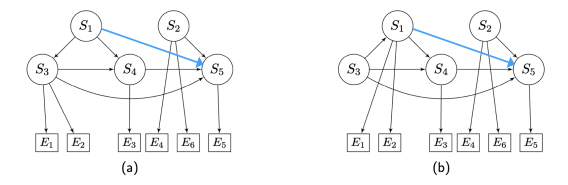


Figure 1: New edges drawn in blue.

Construct phi

```
phi1
a) Φ
     S1 S2 S3 S4 S5
## S1 1 0 1 1 1
## S2 0 1 0 0 1
## S3 0 0 1 1 1
## S4 0 0 0 1 1
## S5 0 0 0 0 1
phi2
b) Φ
     S1 S2 S3 S4 S5
##
## S1 1 0 0 1 1
## S2 0 1 0 0 1
## S3 1 0 1 1 1
## S4 0 0 0 1 1
## S5 0 0 0 0 1
theta1 = array(dim = c(5,6), dimnames = list(c("S1", "S2", "S3", "S4", "S5"),
                                           c("E1", "E2", "E3", "E4", "E5", "E6")))
theta1["S1",] = c(0,0,0,0,0,0)
theta1["S2",] = c(0,0,0,1,0,1)
theta1["S3",] = c(1,1,0,0,0,0)
theta1["S4",] = c(0,0,1,0,0,0)
theta1["S5",] = c(0,0,0,0,1,0)
theta2 = array(dim = c(5,6), dimnames = list(c("S1", "S2", "S3", "S4", "S5"),
                                           c("E1", "E2", "E3", "E4", "E5", "E6")))
theta2["S1",] = c(1,1,0,0,0,0)
theta2["S2",] = c(0,0,0,1,0,1)
theta2["S3",] = c(0,0,0,0,0,0)
theta2["S4",] = c(0,0,1,0,0,0)
theta2["S5",] = c(0,0,0,0,1,0)
```

Construct thetha

```
theta1
```

```
a) \Theta
##
    E1 E2 E3 E4 E5 E6
## S1 0 0 0 0 0 0
## S2 0 0 0 1 0 1
## S3 1 1 0 0 0 0
## S4 0 0 1 0 0 0
## S5 0 0 0 0 1 0
```

```
theta2
b) Θ
##
     E1 E2 E3 E4 E5 E6
## S1 1 1 0 0 0
## S2 0
        0 0 1 0 1
## S3 0
        0 0 0 0
## S4 0 0 1 0 0 0
## S5 0 0 0 0 1 0
F1 = phi1%*%theta1
F2 = phi2%*%theta2
Calculate F = \Phi\Theta
F1
a) F
##
     E1 E2 E3 E4 E5 E6
## S1 1 1 1 0 1 0
## S2 0 0 0 1 1 1
## S3 1 1 1 0 1 0
## S4 0 0 1 0 1 0
## S5 0 0 0 0 1 0
F2
b) F
##
     E1 E2 E3 E4 E5 E6
## S1 1 1 1 0 1 0
## S2 0
        0 0 1 1 1
## S3
     1 1 1 0 1 0
## S4
     0
        0
          1
             0 1
                  0
## S5 0 0 0 0 1 0
```

2. Assuming no noise, determine the discrete data D1 and D2 from both models. Given only the data, can you tell apart the two models?

```
D1 = array(dim = c(6, 5), dimnames = list(c("E1", "E2", "E3", "E4", "E5", "E6"), c("S1", "S2", "S3", "S4", "S5")))

D1["E1",] = c(1,0,1,0,0)

D1["E2",] = c(1,0,1,0,0)

D1["E3",] = c(1,0,1,1,0)

D1["E4",] = c(0,1,0,0,0)

D1["E5",] = c(1,1,1,1,0)

D1["E6",] = c(0,1,0,0,1)

D2 = array(dim = c(6, 5), dimnames = list(c("E1", "E2", "E3", "E4", "E5", "E6"),
```

```
c("S1", "S2", "S3", "S4", "S5")))
D2["E1",] = c(1,0,1,0,0)
D2["E2",] = c(1,0,1,0,0)
D2["E3",] = c(1,0,1,1,0)
D2["E4",] = c(0,1,0,0,0)
D2["E5",] = c(1,1,1,1,0)
D2["E6",] = c(0,1,0,0,1)
D1
a) D1
##
     S1 S2 S3 S4 S5
## E1
      1
         0
            1
              0
## E2
      1
         0
            1
               0
## E3
      1
         0
            1
               1
                  0
## E4
     0 1 0 0 0
## E5
     1 1 1 1 0
## E6 0 1
           0 0
D2
2) D2
     S1 S2 S3 S4 S5
## E1
         0 1 0
     1
## E2 1
         0
## E3 1
         0
            1
              1
                  0
## E4
      0
         1
            0
               0
                  0
## E5
     1 1
           1
                  Λ
              1
```

Since the Data matrices D1 and D2 are identical, we cannot tell the two models apart.

3. Use the mnem1 package for this question: Take D1 and D2 from the previous question. For each model, calculate the marginal log-likelihood ratio (network score) given the data by setting the false positive rate to be 5 and the false negative rate to be 1.

```
library(mnem)
## Registered S3 methods overwritten by 'RcppEigen':
##
    method
                          from
##
    predict.fastLm
                          RcppArmadillo
##
    print.fastLm
                          RcppArmadillo
##
     summary.fastLm
                          RcppArmadillo
     print.summary.fastLm RcppArmadillo
network_score_1 = scoreAdj(D1,adj = phi1,method="disc",fpfn=c(0.05,0.01))$score
network_score_2 = scoreAdj(D2,adj = phi2,method="disc",fpfn=c(0.05,0.01))$score
network_score_1
```

```
Network score of (a)
```

```
## [1] 51.68914
```

```
network_score_2
```

Network score of (b)

```
## [1] 51.68914
```

Problem 21: Hidden Markov NEMs

1. Using the definitions for HM-NEMs from the lecture, compute the transition probabilities from $G_t = u$ to $G_{t+1} \in v1, v2$ for different smoothness parameter $\lambda \in 0.1, \ldots 0.9$.

```
u = t(array(c(c(1,1,1,0)),
              c(0,1,1,1),
              c(0,0,1,1),
              c(0,0,0,1)),
              dim = c(4, 4), dimnames = list(c("S1", "S2", "S3", "S4"),
                                              c("S1", "S2", "S3", "S4"))))
v1 = t(array(c(c(1,1,1,0),
              c(0,1,1,1),
              c(0,0,1,0),
              c(0,0,0,1)),
              dim = c(4, 4), dimnames = list(c("S1", "S2", "S3", "S4"),
                                              c("S1", "S2", "S3", "S4"))))
v2 = t(array(c(c(1,0,0,0)),
              c(1,1,1,0),
              c(1,0,1,0),
              c(1,0,0,1)),
              dim = c(4, 4), dimnames = list(c("S1", "S2", "S3", "S4"),
                                              c("S1", "S2", "S3", "S4"))))
lambda = seq(0.1, 0.9, by=0.1)
s_uv1 = sum(u!=v1)
s_uv2 = sum(u!=v2)
w = mnem:::enumerate.models(c("S1", "S2", "S3", "S4"), trans.close = FALSE)
```

Generated 4096 unique models (out of 4096)

```
s_uw = array(dim = c(length(w), 1))
for(i in 1:length(w)){
    s_uw[i] = sum(u!=w[[i]])
}

power <- function(x, y) sign(x) * abs(x)^y

T = array(dim = c(9,2), dimnames = list(lambda,c("v1", "v2")))
C = array(dim = c(9,1), dimnames = list(lambda,c("C")))
for(i in lambda){
    C[as.character(i),] = sum(power((1-i),s_uw)*i)
    T[as.character(i),"v1"] = (1/C[as.character(i),])*((1-i)^s_uv1)*i
    T[as.character(i),"v2"] = (1/C[as.character(i),])*((1-i)^s_uv2)*i</pre>
```

```
}
```

Т

Transition probabilities

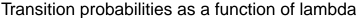
```
## v1 v2
## 0.1 0.0004066299 2.160998e-04
## 0.2 0.0006915442 1.812842e-04
## 0.3 0.0012014646 1.413511e-04
## 0.4 0.0021316282 9.945325e-05
## 0.5 0.0038536733 6.021365e-05
## 0.6 0.0070554312 2.889905e-05
## 0.7 0.0128765947 9.387038e-06
## 0.8 0.0224313310 1.435605e-06
## 0.9 0.0318630818 3.186308e-08
```

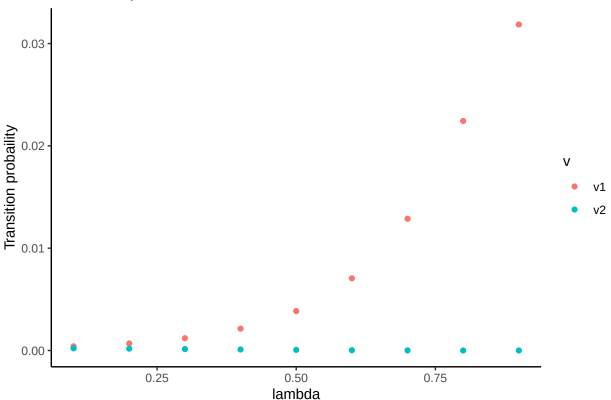
2. Plot the transition probabilities for v_1 and v_2 as a function of λ . Describe the transition probabilities as a function of λ .

```
library(reshape2)
library(ggplot2)
library(RColorBrewer)

## Warning: package 'RColorBrewer' was built under R version 4.0.5

data = data.frame(melt(T))
colnames(data)<-c("lambda","v","T")
plot<-ggplot(data,aes(x=lambda,y=T,color=v))+
    geom_point()+
    theme_classic()+
    ylab("Transition probabilities as a function of lambda")
plot.</pre>
```





As λ increases the similarity between networks becomes more relevant for the transition probabilities, that is, dissimilar networks get more highly penalized with a lower transition probability as λ increases. Since v_1 is more similar to u than v_2 , the probability to transition into v_1 increases as we increase λ . Conversely as we bring λ close to zero, network similarity becomes less relevant and we see the transition probabilities of v_1 and v_2 converge.

Problem 22: Mixture NEMs

S2

0 1

1. Determine the the cellular perturbation map ρ , where $\rho_{ic} = 1$ if cell c is perturbed by a knock-down of S-gene i.

2. Assume that C_1, C_2 are generated from F_1 and C_3, C_4 are generated from F_2 , compute the noiseless log odds matrix R, where $R_{jc} > 0$ means that the perturbation on cell c has an effect on E-gene j:

```
phi_F1["S1",] = c(1,1)
phi_F1["S2",] = c(0,1)
phi_F2 = array(dim = c(2,2), dimnames = list(c("S1", "S2"),
                                             c("S1", "S2")))
phi_F2["S1",] = c(1,0)
phi_F2["S2",] = c(1,1)
theta_F1 = array(\dim = c(2,2), \dim = list(c("S1","S2"),
                                               c("E1","E2")))
theta_F1["S1",] = c(1,0)
theta_F1["S2",] = c(0,1)
theta_F2 = array(\dim = c(2,2), \dim = list(c("S1","S2"),
                                              c("E1","E2")))
theta_F2["S1",] = c(0,1)
theta_F2["S2",] = c(1,0)
EEP_F1 = t(t(rho)%*%phi_F1%*%theta_F1)
EEP_F1[EEP_F1>1] = 1
EEP_F2 = t(t(rho)%*%phi_F2%*%theta_F2)
EEP_F2[EEP_F2>1] = 1
```

(a) For each component k, compute the expected effect pattern $(\rho^T \phi^k \theta^k)^T$. Replace all non-zeros by 1.

```
EEP_F1
```

Expected effect pattern of F1

```
## C1 C2 C3 C4
## E1 1 0 1 0
## E2 1 1 1 1
```

```
EEP_F2
```

Expected effect pattern of F2

```
## C1 C2 C3 C4
## E1 0 1 1 1
## E2 1 1 1 1
```

```
R= cbind(EEP_F1[,1:2],EEP_F2[,3:4])
R[R==0] = -1
R
```

(b) Based on the component assignment for each cell, extract the corresponding column from the expected effect patterns computed above and put it into R. Replace all zeros by -1.

```
## C1 C2 C3 C4
## E1 1 -1 1 1
## E2 1 1 1 1
```

3. Take R from the previous question. Given the vector of mixture weights $\pi = (0.44, 0.56)$, calculate the responsibilities Γ . Then, update the mixture weights.

```
L1 = t(EEP_F1)\%*\%R
L2 = t(EEP_F2)\%*\%R
print("L1")
## [1] "L1"
L1
     C1 C2 C3 C4
##
## C1 2 0 2 2
## C2 1 1 1 1
## C3 2 0 2 2
## C4 1 1 1 1
print("L2")
## [1] "L2"
L2
     C1 C2 C3 C4
##
## C1 1 1 1 1
## C2 2 0 2 2
## C3 2 0 2 2
## C4 2 0 2 2
pi = c(0.44, 0.56)
gamma = array(dim = c(2,4), dimnames = list(c("F1","F2"),
                                           c("C1", "C2", "C3", "C4")))
gamma["F1",] = pi[1]*exp(diag(L1))/(pi[1]*exp(diag(L1))+pi[2]*exp(diag(L2)))
gamma["F2",] = pi[2]*exp(diag(L2))/(pi[2]*exp(diag(L2))+pi[1]*exp(diag(L1)))
Responsibilities gamma
pi[1] = sum(gamma["F1",])/(sum(gamma["F1",])+sum(gamma["F2",]))
pi[2] = sum(gamma["F2",])/(sum(gamma["F1",])+sum(gamma["F2",]))
```

Updated mixture weights

рi

[1] 0.5066091 0.4933909