```
[1] "# of discriminatory genes = 25"
```

- [1] "fold change for discriminatory genes: 0.3"
- [1] "fold change for nondiscriminatory genes: 0"

choose_k

2

100

The optimal number of clusters K is set as the most frequently found K from the 100 simulations tabulated above. It is found by using the BIC criterion after running the unpenalized EM algorithm on K spanning from 2 to 8.

Using the last set of simulated counts, I ran a grid search across varying tuning parameters. As done in Pan et al, I fixed $\lambda_1 = 1$, and searched over $\lambda_2 = (0.1, 0.2, ..., .2)$ and $\tau = (0.1, 0.2, ..., 2)$:

- [1] "tau= 0.2" "tau= 0.3" "tau= 0.1"
- [1] "lambda1= 1"
- [1] "lambda2= 0.2" "lambda2= 0.3" "lambda2= 0.4"

The results of the final run based on optimal tuning parameters are below: Below are the summary of results:

- [1] "Mean pi: 0.620869565217391" "Mean pi: 0.379130434782609"
- [1] "First 3 genes:"

[,1] [,2]

- [1,] 3.569660 4.153140
- [2,] 4.800947 5.389779
- [3,] 4.305897 4.902170
- [1] "Last 3 genes:"

[,1] [,2]

- [98,] 6.400696 6.398118
- [99,] 5.950138 5.952305
- [100,] 6.378860 6.376469
- [1] "Mean % of nondiscriminatory genes: 0.7506"
- [1] "Mean ARI: 1"
- [1] "Mean sensitivity: 0.9968"
- [1] "Mean false positive rate: 0.000266666666666667"