# Solution 2

#### STAT6306

### Introduction

A major issue with antiretroviral drugs is the mutation of the virus' genes. Because of its high rate of replication (10<sup>9</sup> to 10<sup>10</sup> virus per person per day) and error-prone polymerase<sup>1</sup>, HIV can easily develop mutations that alter susceptibility to antiretroviral drugs. The emergence of resistance to one or more antiretroviral drugs is one of the more common reasons for therapeutic failure in the treatment of HIV.

In the paper 'Genotypic predictors of human immunodeficiency virus type 1 drug resistance'<sup>2</sup>, a sample of in vitro<sup>3</sup> HIV viruses were grown and exposed to a particular antiretroviral therapy. The susceptibility of the virus to treatment and the number of genetic mutations of each virus were recorded.

### Question 1

```
load("hiv.rda")

X = hiv.train$x
Y = hiv.train$y

geneLabels = colnames(X)
```

(a)

What are n and p in this problem? What are the features in this problem? What are the observations? What is the supervisor? **Note:** Attempt to answer this question before moving on to the rest of the questions.

```
#SOLUTION
(n = nrow(X))

## [1] 704
(p = ncol(X))

## [1] 208
```

#### SOLUTION

There are 208 features (p) and 704 observations (n). The features are indicators for whether or not there was a mutation in a gene. The supervisor is the log(susceptibility) of the HIV virus to a particular drug therapy

<sup>&</sup>lt;sup>1</sup>An enzyme that 'stitches' back together DNA or RNA after replication

<sup>&</sup>lt;sup>2</sup>The entire paper is on the website. Try to see what you can get out of it if you have the time.

<sup>&</sup>lt;sup>3</sup>Latin for 'in glass', sometimes known colloquially as a test tube

### Question 2

Consider the feature matrix X. It is composed of 0's and 1's, with a 1 indicating a mutation in a particular gene. Look at the output for the following chunk of code.

table(X)

```
## X
## 0 1
## 135589 10843
```

What results do you see? What does this indicate?

### **SOLUTION**

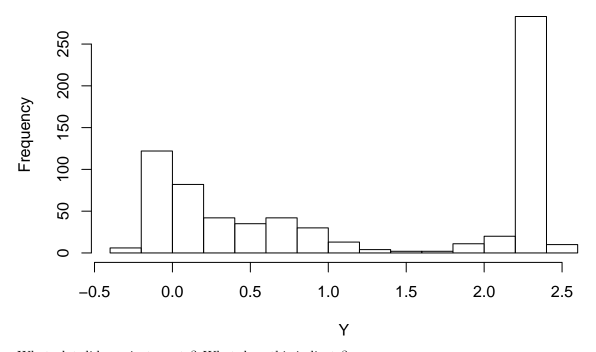
Based on the feature matrix X, we see that there are 135589 unmutated/"normal" genes and 10843 genes that have mutations.

## Question 3

The supervisor is the log transformed susceptibility of a virus to the considered treatment, with large values indicating the virus is relatively more resistant (that is, not susceptible). Run

hist(Y)

# **Histogram of Y**



What plot did you just create? What does this indicate?

#### **SOLUTION**

This gives us a histogram of the frequency of the susceptibility of a virus to the considered treatment. The marginal distribution of the supervisor Y is bimodal, with a peak around 0 (higher susceptible) and around 2.4 (lower susceptible)

### Question 4

We may have (at least) two goals with a data set such as this:

- inference: can we find some genes whose mutation seems to be most related to viral susceptibility?
- prediction: can we make a model that would predict whether this therapy would be efficacious, given a virus with a set of genetic mutations

(a)

Try to find the best subset solution for this problem. Discuss any problems or findings you discover. In particular, how many possible models are there?

#### **SOLUTION**

```
2^p
```

```
## [1] 4.113761e+62
```

There are  $2^p$  possible different solutions, which, given the size of p (208), gives us 4.1137614e+62 possible solutions, which is way too large to compute all subsets.

#### (b) Inference

(i)

Find the selected model for:

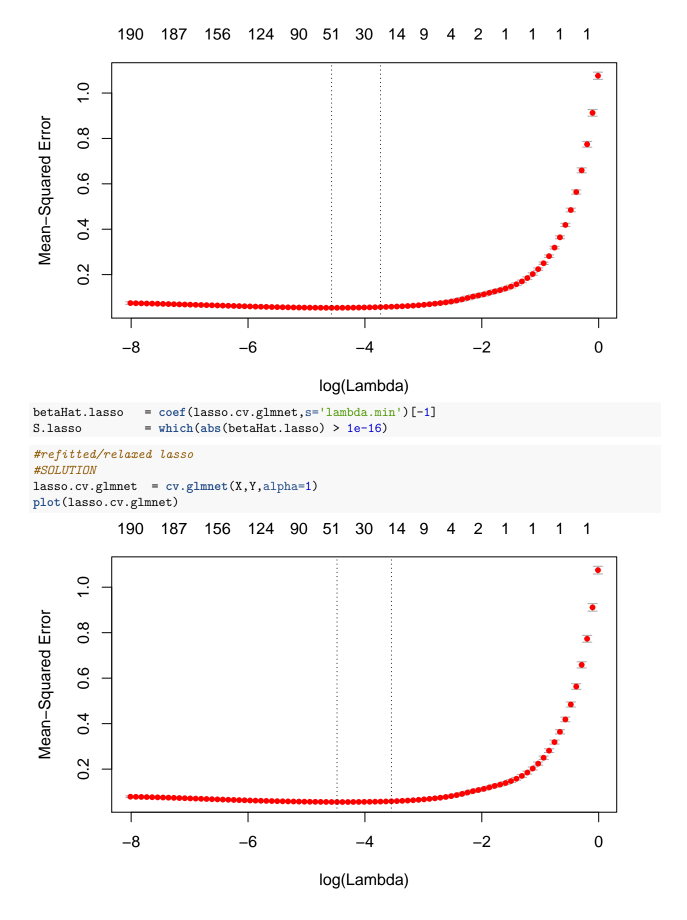
- forward selection using BIC as the criterion
- lasso
- refitted/relaxed lasso

```
#Forward selection
#SOLUTION
if(!require(leaps)){install.packages('leaps',repos='http://cran.us.r-project.org');require(leaps)}
## Loading required package: leaps
outForward = regsubsets(x=X,y=Y,nvmax=p,method='forward')

## Warning in leaps.setup(x, y, wt = weights, nbest = nbest, nvmax = nvmax, :
## 12 linear dependencies found

## Reordering variables and trying again:
## Warning in rval$lopt[] <- rval$vorder[rval$lopt]: number of items to
## replace is not a multiple of replacement length</pre>
```

```
# note this warning is that the feature matrix
# isn't full rank. This is, there are redundant
# columns in it:
cat('The rank is: ',qr(X)$rank,' while the # of features is: ',p,'\n')
## The rank is: 196 while the # of features is: 208
sumForward
              = summary(outForward)
model.forward = sumForward$which[which.min(sumForward$bic),]
               = model.forward[-1] #get rid of the intercept entry
S.forward
               = lm(Y~X[,S.forward]) #regsubsets only scores models, not fit them
betaHat.forward = coef(lm.forward)
betaHat.forward
          (Intercept) X[, S.forward]p33 X[, S.forward]p54
##
##
                              0.09489890
          0.07426093
                                                 0.60889375
## X[, S.forward]p58 X[, S.forward]p65 X[, S.forward]p67
##
         -0.66464456
                              0.66361310
                                                 0.12187768
## X[, S.forward]p69 X[, S.forward]p75 X[, S.forward]p90
##
          0.09158451
                              0.08995627
                                                 0.15145415
## X[, S.forward]p102 X[, S.forward]p115 X[, S.forward]p117
##
          0.07556800
                              0.29618456
                                                -0.45272186
## X[, S.forward]p151 X[, S.forward]p172 X[, S.forward]p184
          0.29833631
                              0.20732278
                                                 1.89449998
## X[, S.forward]p187 X[, S.forward]p210 X[, S.forward]p215
          -0.45850428
                              0.07844976
                                                 0.21895472
#lasso
if(!require(glmnet)){install.packages('glmnet',repos='http://cran.us.r-project.org');require(glmnet)}
## Loading required package: glmnet
## Loading required package: Matrix
## Loading required package: foreach
## Loaded glmnet 2.0-10
#SOLUTION
lasso.cv.glmnet = cv.glmnet(X,Y,alpha=1) #note: we are standardizing the features...
plot(lasso.cv.glmnet) #note that this output is random... why?
```



```
betaHat.temp = coef(lasso.cv.glmnet,s='lambda.1se')[-1]
S.refitted = which(abs(betaHat.temp) > 1e-16)
lm.refitted = lm(Y ~ X[,S.refitted])
betaHat.refitted = coef(lm.refitted)
```

(ii)

Comparing the selected models for each of the above methods

```
#SOLUTION
cat('The selected genes from forward selection + BIC are: \n',
    geneLabels[S.forward],'\n')
## The selected genes from forward selection + BIC are:
## p33 p54 p58 p65 p67 p69 p75 p90 p102 p115 p117 p151 p172 p184 p187 p210 p215
cat('The selected genes from lasso are: \n',
    geneLabels[S.lasso],'\n')
## The selected genes from lasso are:
## p21 p33 p41 p43 p54 p58 p60 p65 p66 p67 p69 p75 p77 p80 p83 p90 p102 p107 p110 p115 p116 p117 p118
cat('The selected genes from refitted lasso are: \n',
    geneLabels[S.refitted],'\n')
## The selected genes from refitted lasso are:
## p33 p41 p65 p67 p69 p75 p77 p90 p115 p118 p151 p181 p184 p190 p210 p215 p228
#Note that we can directly compare chosen models
geneLabels[S.forward] %in% geneLabels[S.refitted]
## [1]
        TRUE FALSE FALSE TRUE TRUE TRUE TRUE FALSE TRUE FALSE
        TRUE FALSE TRUE FALSE TRUE TRUE
## [12]
geneLabels[S.refitted] %in% geneLabels[S.forward]
## [1] TRUE FALSE TRUE TRUE TRUE TRUE FALSE TRUE TRUE FALSE TRUE
## [12] FALSE TRUE FALSE TRUE
                               TRUE FALSE
```

### (c) Prediction

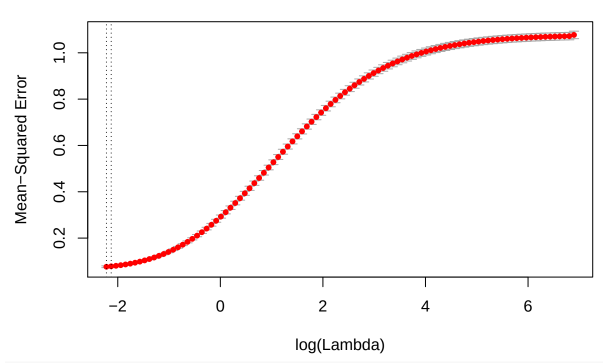
### (i) Ridge regression

Now that are looking at prediction, we can use ridge regression (which only addresses prediction). Using the package glmnet, plot the CV curve over the grid of  $\lambda$  values and indicate the minimum, and finally report the CV estimate of the prediction risk for  $\hat{\beta}_{\text{ridge}}(\hat{\lambda})$ 

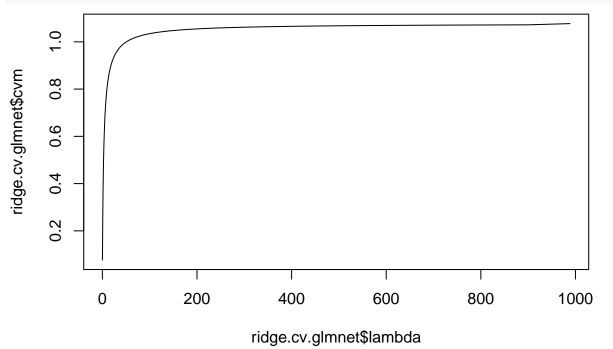
**Note:** There is no need to report the p coefficient estimates from the ridge solution. Also, glmnet has a grid problem. Make two plots, one that shows the problem and one that shows it being corrected.

```
#SOLUTION
ridge.cv.glmnet = cv.glmnet(X,Y,alpha=0)
plot(ridge.cv.glmnet)
```





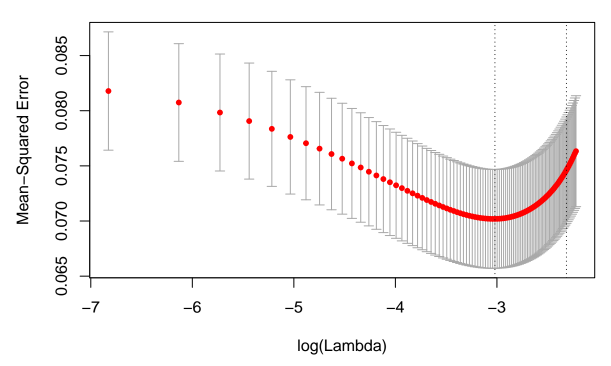
#or
plot(ridge.cv.glmnet\$lambda,ridge.cv.glmnet\$cvm,type='l')



```
#CV estimate of the prediction error:
min(ridge.cv.glmnet$cvm)
```

```
## [1] 0.07633416
min.lambda = min(ridge.cv.glmnet$lambda)
lambda.new = seq(min.lambda, min.lambda*0.01,length=100)
```

```
ridge.cv.glmnet = cv.glmnet(x = X, y = Y, alpha = 0,lambda = lambda.new)
plot(ridge.cv.glmnet) #now it is in middle
```



#### (ii) Prediction on a test set

Now, let's look at some predictions made by these methods. Use the following for the test set:

```
X_0 = hiv.test$x
Y_0 = hiv.test$y
```

Find an estimate of the risk using the test observations for

- forward selection using BIC as the criterion
- ridge
- lasso
- refitted/relaxed lasso

```
#### Get predictions on test set:
Yhat.test.forward = X_0[,S.forward] %*% betaHat.forward[-1] + betaHat.forward[1]
Yhat.test.ridge = predict(ridge.cv.glmnet,X_0,s='lambda.min')
Yhat.test.lasso = predict(lasso.cv.glmnet,X_0,s='lambda.min')
Yhat.test.refitted = X_0[,S.refitted] %*% betaHat.refitted[-1] + betaHat.refitted[1]

# Get estimate of prediction risk via the test set error
Yhat.test.forward = mean((Yhat.test.forward - Y_0)**2)
pred.error.ridge = mean((Yhat.test.ridge - Y_0)**2)
pred.error.lasso = mean((Yhat.test.lasso - Y_0)**2)
pred.error.refitted = mean((Yhat.test.refitted - Y_0)**2)
```

```
cat('The prediction error from forward selection + BIC are: \n',
    Yhat.test.forward, '\n')
## The prediction error from forward selection + BIC are:
## 0.07491952
cat('The prediction error from ridge is: \n',
     pred.error.ridge,'\n')
## The prediction error from ridge is:
## 0.09708477
cat('The prediction error from lasso is: \n',
     pred.error.lasso,'\n')
## The prediction error from lasso is:
## 0.06878929
cat('The prediction error from refitted lasso is: \n',
    pred.error.refitted,'\n')
## The prediction error from refitted lasso is:
## 0.06688864
(d)
Challenge Suppose we didn't have access to any test data. How could you provide an estimate of the risk?
What are the pros and cons of your proposal?
```

```
cat('The CV estimate of risk of ridge(lambdaHat) = ',min(ridge.cv.glmnet$cvm),'\n')
## The CV estimate of risk of ridge(lambdaHat) = 0.07019062
Compare this with the test set estimate:
```

```
cat('The test set estimate of risk from ridge is: \n',
    pred.error.ridge,'\n')
```

```
## The test set estimate of risk from ridge is:
## 0.09708477
```

So, by minimizing CV as a function of lambda, we have produced a reasonable, but overly optimistic estimate of the risk.

## Question 6

Using the lasso with CV minimum tuning parameter, which gene mutations are related to susceptibility?

```
#SOLUTION
geneLabels[S.lasso]
```

```
## [1] "p21" "p33" "p41" "p43" "p54" "p58" "p60" "p65" "p66" "p67" "## [11] "p69" "p75" "p77" "p80" "p83" "p90" "p102" "p107" "p110" "p115" "## [21] "p116" "p117" "p118" "p122" "p123" "p127" "p138" "p139" "p144" "p150" "## [31] "p151" "p162" "p171" "p172" "p173" "p176" "p181" "p184" "p187" "p188"
```

```
## [41] "p190" "p192" "p200" "p202" "p203" "p210" "p211" "p212" "p215" "p219" ## [51] "p228"
```

### Question 7

At which gene mutation sites are the presence of a mutation associated with a decrease in viral susceptibility to this particular drug? Hint: Consider the signs of the coefficients. What gene site has the largest estimated effect using  $\hat{\beta}_{lasso}(\hat{\lambda})$ ?

```
#SOI.UTTON
## Gene mutation sites related to a decrease in viral susceptibility:
   thought process: Y increase <-> decrease susceptibility (i.e. increase resistance)
                    so \hat{j} > 0 associated with decrease in susceptibility
geneLabels[betaHat.lasso > 1e-16]
               "p33"
                             "p43"
   [1] "p21"
                                    "p54"
                                           "p65"
                                                  "p66"
                      "p102" "p115" "p116" "p118" "p123" "p127" "p138" "p139"
## [11] "p77"
## [21] "p150" "p151" "p162" "p171" "p172" "p181" "p184" "p188" "p190" "p203"
## [31] "p210" "p211" "p215" "p219" "p228"
geneLabels[which.max(abs(betaHat.lasso))]
## [1] "p184"
```

## Additional challenge problems:

I don't want to overwhelm you with homework problems. However, there are additional topics that are relevant for an interested student. You don't need to do these/turn them in.

## Question 8

Derive, implement, and run both "batch" and "stochastic" gradient descent for this HIV data.

# Question 9

The LARS algorithm is quite similar to forward selection. Run LARS using the option forward stagewise and compare it to forward selection using Mallow's Cp.

# Question 10

Try and use a GIC-based method instead of K-fold CV for finding  $\hat{\lambda}$  using the HIV data.