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Shrinkage Methods for Regularized Regression

Shrinkage Methods - Regularized Regression

What to do if we have too many predictors?

- We have already discussed that too many predictors can create collinearity problems.
- Increasing the number of predictors might increase the prediction error.
- More predictors do not necessarily mean a better model, but more predictors would mean more information, and less bias.

We study three different methods to shrink the number of predictors in order to find a trade-off between model bias and prediction error.

- Principal Components Regression
- Ridge Regression
- Lasso Regression

Principal Components Regression

What to do when we have too many predictors?:

- Perform dimensionality reduction in the predictor space.
- Be aware that predictors might be highly correlated.
- Take matrix X of predictors and center the columns of X to have zero mean. Consider X with no intercept column.
- Find directions of greater variation in the data.

Principal Component Analysis:

The steps to find directions of greater variation in matrix X:

- Find \mathbf{u}_1 to maximize variance of $\mathbf{X}\mathbf{u}_1$ subject to $\mathbf{u}_1^{\mathsf{T}}\mathbf{u}_1=1$.
- Find ${\bf u}_2$ to maximize variance of ${\bf X}{\bf u}_2$ subject to ${\bf u}_1^\top {\bf u}_2 = 0$ and ${\bf u}_2^\top {\bf u}_2 = 1$.
- Continue looking for directions of greatest variation in the data which are orthogonal to the previous ones.
- Continue until the total number of dimensions is exhausted.

The principal components are given by the columns of matrix \mathbf{Z} , where

$$\mathbf{Z} = \mathbf{X} [\mathbf{u}_1 \mathbf{u}_2 \cdots \mathbf{u}_m] = \mathbf{X} \mathbf{U}$$

 \mathbf{z}_i and \mathbf{u}_i are the columns of \mathbf{Z} and \mathbf{U} respectively. \mathbf{U} is called the rotation matrix. \mathbf{Z} is a version of the data rotated in such a way that the resulting principal components are orthogonal.

• Each Principal Component is a linear combination of the original variables $\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_m$ with weights given by each column \mathbf{u}_i of matrix \mathbf{U} :

$$\mathbf{z}_i = u_{1i}\mathbf{x}_1 + u_{2i}\mathbf{x}_2 + \ldots + u_{mi}\mathbf{x}_m$$

- PCs are very sensitive to outliers.
- The Mahalanobis distance can be used to measure the distance of a point to the data mean, after adjusting for correlation in the data.
- Under the multivariate normality assumption in m dimensions, the Mahalanobis distance $d_i = \sqrt{(\mathbf{x}_i \mu)^\top \Sigma^{-1}(\mathbf{x}_i \mu)}$ can be estimated using the sample estimators for μ and Σ and the quantity d_i^2 follows a χ_m^2 distribution. This can be used to detect outliers in higher dimensions.

In Principal Components Regression (PCR):

- ullet Replace model $Y \sim X$ by the model $Y \sim Z$
- ullet Only need to use the first few columns of Z as predictors
- Interpretation of the PCAs as predictors might be challenging. We need to use the values of \mathbf{u}_i in the rotation matrix (also called the loadings) for interpretation.
- ullet Sometimes we can make better predictions with a small number of PCs in ${f Z}$ than with a large number of predictors in ${f X}$

How many Principal Components?

- The trace of the sample variance-covariance S of \mathbf{X} (total sample variance) is equal to the sum if its eigenvalues: $trace(S) = s_1^2 + s_2^2 + \ldots + s_m^2 = \lambda_1 + \lambda_2 + \ldots + \lambda_m$.
- Often most of the total variance of a data set is concentrated in the first principal components.
- Make a plot of the PCs standard deviations $(\sqrt{\lambda_i})$ vs. the PC index i. This is called the scree plot.
- Look for the PC index *i* where there is a big change in slope (the elbow) in the scree plot.
- Another way is to calculate the cumulative variance explained by the first PCs, and retain the number of PCs explaining between 70% to 90% of the total variation.
- Another option is to discard PCs such that $\lambda_i < \bar{\lambda}$.

PCR Example

- meatspec data set from the faraway library.
- The goal is to predict fat content (response) using 100-channel spectrum of absorbances (predictors) in 215 samples of finely chopped meat.
- Partition the data into training sample and testing sample to test model performance.
- Fit model with all predictors using the training data set and make a prediction on the testing set.
- Select few PCs representing the 100 predictors and repeat the process (shrinkage effect)
- What about if we select the number of PCs to minimize the prediction error instead?. We can use *cross-validation*.

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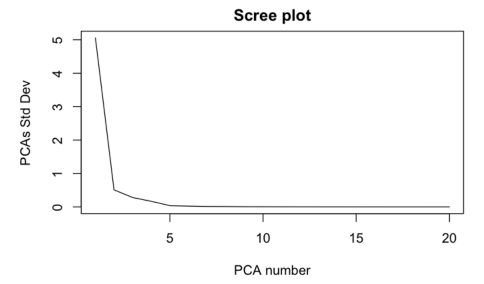
PCR Example: meatspec data set

Function **prcomp** can be used to calculate the PCs and extract the λ 's squared-roots (sdev) and *eigenvectors* (rotation) of the variance-covariance matrix:

```
data(meatspec,package="faraway")
trainmeat<-meatspec[1:172,]
testmeat<-meatspec[173:215,]
modl<-lm(fat~.,trainmeat)
meatpca<-prcomp(trainmeat[,-101])
round(meatpca$sdev,3)[1:50]</pre>
```

```
## [1] 5.055 0.511 0.282 0.168 0.038 0.025 0.014 0.011 0.005 0.003 0.002 0.002 ## [13] 0.001 0.001 0.001 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 ## [25] 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 ## [37] 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 ## [49] 0.000 0.000 0.000 0.000
```

The first two PC's explains around 90% of the total variation of meat fat content.



According to the scree plot, 4 PC's (elbow at 5th PC) seem adequate to represent the data.

The **pcr** function (principal component regression) from the *pls* package has useful features for prediction and cross-validation. We can easily calculate the RMSE for the training set and the testing set.

```
modpcr<-pcr(fat ~ ., data=trainmeat,ncomp=50)
#summary(modpcr)
#RMSE with 4 PCAs
rmse(predict(modpcr,ncomp=4),trainmeat$fat)

## [1] 4.064745

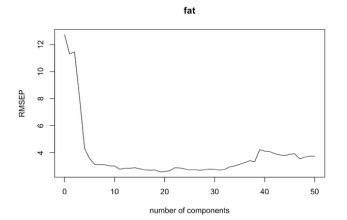
rmse(predict(modpcr,testmeat,ncomp=4),testmeat$fat)

## [1] 4.533982</pre>
```

As expected the testing error is larger than the training error. We can do better by selecting the number of PC's that minimize the CV error.

You can use the function **RMSEP** instead, to select the number of PC's that minimize the 10-fold Cross-Validation error. The resulting Cross-Validation error is $<2.5\,$

```
#Use 10-fold Cross-Validation
set.seed(123)
modpcrev<-pcr(fat-.,data=trainmeat,validation="CV",ncomp=50)
pcrCV<-RMSEP(modpcrev,estimate="CV")
plot(pcrCV)</pre>
```



Ridge Regression

- Although the aim of PCR is to reduce dimensionality in the number of predictors, you still have to measure all the predictors since each PC is a linear combination of all predictors.
- Ridge regression assumes that after normalization, some of the regression coefficients should not be very large.
- Ridge regression is very useful when you have collinearity and the LS regression coefficients are unstable.
- The method uses a penalized regression since the LS minimization problem has a penalty term:

$$\mathsf{minimize}(y - X\boldsymbol{\beta})^\top (y - X\boldsymbol{\beta}) + \lambda \sum_{i} \boldsymbol{\beta}_j^2$$

for some $\lambda \geq 0$. The penalty term is $\sum_{j} \beta_{j}^{2}$

- ullet Usually predictors are standardized first (centered by their means and scaled by their standard deviations) and the response y is centered.
- The ridge regression estimates are:

$$\hat{\boldsymbol{\beta}} = (X^{\top}X + \lambda I)^{-1}X^{\top}y$$

(Note the extra term λI or ridge in the $X^{\top}X$ matrix)

ullet The difference with standard LS is that the problem solution $oldsymbol{eta}$ minimizes:

$$(y - X\boldsymbol{\beta})^{\top}(y - X\boldsymbol{\beta})$$
 subject to $\sum_{j=1}^{p} \boldsymbol{\beta}_{j}^{2} \leq t^{2}$

• The parameter λ (or t) should be chosen to have stable estimates of β . Can use function *Im.ridge* from the *MASS* library.

- Note that when $\lambda=0$ the ridge regression estimation problem reduces to the standard LS problem, while when $\lambda\to\infty$, $\hat{\pmb{\beta}}\to {\bf 0}.$
- It is useful to plot the values of $\hat{\beta}_i$ as a function of λ .
- The value of λ can be also chosen using automated methods as Generalized Cross-Validation (GCV) (similar to Cross-Validation).
- Ridge regression coefficient estimates are biased.

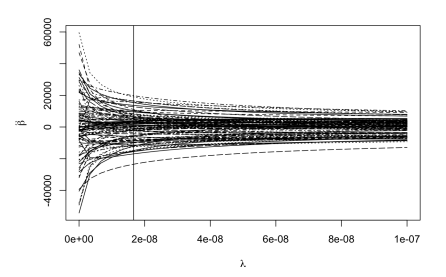
Example: meatspec data set

The value of λ is selected by minimizing the GCV. Some experimentation is required to get the initial λ range.

```
library(MASS)
library(faraway)
data(fat)
trainmeat<-meatspec[1:172,]
testmeat<-meatspec[173:215,]
ridgemod<-lm.ridge(fat-.,trainmeat,lambda=seq(0,0.0000001,len=31))
# Predictors are centered and scaled.
# The response is centered
matplot(ridgemod$lambda,coef(ridgemod),type="1",xlab=expression(lambda),ylab=expression(hat(beta)),col=1)
# Look for the value of lambda that minimizes GCV
which.min(ridgemod$GCV)</pre>
```

```
## 1.666667e-08
## 6
```

```
abline(v=1.666667e-08)
```



Trace plot for the *meatspec* data. The vertical line is the value of λ that minimizes the GCV.

Lasso Regression

• In this case the estimated $\hat{\beta}$ minimizes:

$$\mathsf{minimize}(y - X\boldsymbol{\beta})^\top (y - X\boldsymbol{\beta}) + \lambda \sum_j |\beta_j|$$

for some $\lambda \geq 0$. The penalty term is $\sum_{i} |\beta_{i}| (L_{1} \text{ constraint})$.

- In two-dimensions the constraint defines a square. In higher dimensions it defines a polytope.
- Lasso is useful when the response can be explained by few predictors with zero effect on the remaining predictors (Lasso is similar to a variable selection method).
- When $\beta_j=0$ the corresponding predictor is eliminated. This is not the case for ridge regression.

Lasso regression

- Use Lasso when the effect of predictors is *sparse*. This means that only few predictors will have an effect on the response (e.g. gene expression data) or when number of predictors is large (p > n)
- Use the *lars R package* for Lasso
- Select t in the constraint $\sum_{j=1}^{p} |\beta|_j \le t$ by using Cross-Validation (CV).
- As t increases, the number of predictors increases.

Example: state data set

We have data from 50 states:

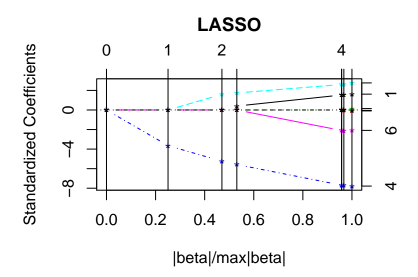
- Population: population estimate as of July 1, 1975
- Income: per capita income (1974)
- Illiteracy: illiteracy (1970, percent of population)
- Life Exp: life expectancy in years (1969–71)
- Murder: murder and non-negligent manslaughter rate per 100,000 population (1976)
- HS Grad: percent high-school graduates (1970)
- Frost: mean number of days with minimum temperature below freezing (1931–1960) in capital or large city
- Area: land area in square miles

We want to predict *Life Expectancy*

Example: *state* data set

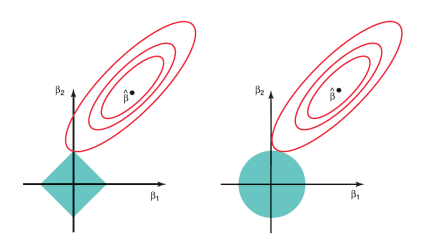
```
Use function lars from library lars
library(lars)
## Loaded lars 1.2
data(state)
statedata<-data.frame(state.x77,row.names = state.abb)</pre>
names(statedata)[1:4]
## [1] "Population" "Income" "Illiteracy" "Life.Exp"
names(statedata)[5:8]
## [1] "Murder" "HS.Grad" "Frost" "Area"
#No model equation in function lars
modlasso<-lars(as.matrix(statedata[,-4]),
                          statedata$Life.Exp)
```

plot(modlasso)



- ullet The x-axis in the graph is the scaled value of $t=\sum_{j=1}^p |eta_j|$
- This value has been scaled to its maximum value which is the least-square solution.
- As t increases more predictors enter into the model.
- Variable 4 (murder rate) enters first, followed by HS graduation, population and days of frost (*Frost*). The remaining variables enter when t is large (close to the LS solutions).
- t can be selected by cross-validation (lars uses 10-fold CV by default) using function *cvlars*

Comparing Ridge Regression and Lasso



Comparing Ridge Regression and Lasso

- Lasso selects a sub-set of predictors (some coefficients equal to zero).
- Ridge regression performs better when the response is a function of many predictors with coefficients around the same size.
- Lasso will perform better when a relatively small number of predictors have large coefficients and the rest are very small or equal to zero.
- Since the number of predictors is never known a priori, cross-validation can be used to decide which approach is better for a particular data set.