

# STATS305A - Lecture 14

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## Contents

<b>1 Announcements</b>	<b>1</b>
<b>2 Recap</b>	<b>1</b>
2.1 Ridge regression . . . . .	1
2.2 PCA . . . . .	2
<b>3 Principal component regression</b>	<b>2</b>
<b>4 Feature selection methods</b>	<b>4</b>
4.1 All subsets . . . . .	4
4.2 Forward stepwise . . . . .	5
4.3 Backwards stepwise . . . . .	5
<b>5 Boosting</b>	<b>6</b>

## 1 Announcements

- HW 3 out now.
- An etude coming soon.
- There will be four homeworks and four etudes in total for the course.

## 2 Recap

### 2.1 Ridge regression

Last time, we started to discuss ridge regression. In ridge regression we use the estimator

$$\hat{\beta}_\lambda = \underset{\lambda}{\operatorname{argmin}} \left\{ \|Xb - Y\|_2^2 + \lambda \|b\|_2^2 \right\}.$$

We saw that if  $U\Gamma V^T$  is the SVD of  $X$ , then we have

$$\begin{aligned} H_\lambda &:= X(X^T X + \lambda I)^{-1} X^T \\ &= U \operatorname{diag} \left( \frac{\gamma_j^2}{\gamma_j^2 + \lambda} \right) U^T, \end{aligned}$$

and our predictions are given by

$$\widehat{Y}_\lambda = H_\lambda Y = \sum_{j=1}^d \frac{\gamma_j^2}{\gamma_j^2 + \lambda} u_j u_j^T Y.$$

We also saw that when doing ridge regression we often replace  $Y$  with  $Y - \bar{Y}\mathbf{1}$ .

## 2.2 PCA

We also started talking about principal component analysis (PCA). We wanted to project each  $x_i \in \mathbb{R}^d$  onto the directions in  $\mathbb{R}^d$  capturing the most variance in our data  $X$ . Equivalently we want to project onto the subspace closest to all the  $x'_i$ s.

## 3 Principal component regression

Roughly, if  $y$  is related to  $x$  we'd hope that the variance in  $y$  is explained by the most varying/important direction in  $X$ .

Recall that if  $X = U\Gamma V^T$  is the SVD of  $X$ , then the first  $k$  principal components are the first  $k$  vectors in  $V = [v_1, v_2, \dots, v_d]$ . The idea behind principal component regression (pcr) is to replace  $x_i \in \mathbb{R}^d$  with the projection of  $x_i$  onto  $\text{span}(V_k)$  where  $V_k = [v_1, \dots, v_k]$ . Define  $\tilde{x}_i = V_k V_k^T x_i$ . We will then run regression on

$$\tilde{X} = \begin{bmatrix} \tilde{x}_1^T \\ \vdots \\ \tilde{x}_n^T \end{bmatrix} \in \mathbb{R}^{n \times d}.$$

Thus our estimate of  $\beta$  is

$$\hat{\beta}_{\text{pcr}(k)} = \underset{b}{\operatorname{argmin}} \|Y - \tilde{X}b\|_2^2.$$

We can formulate this as doing regression in  $\mathbb{R}^k$ . Note that  $\tilde{X} = X V_k V_k^T$  thus we can let  $c = V_k^T b \in \mathbb{R}^k$  so that  $\tilde{X}b = X V_k c$ . We can this instead solve

$$\hat{c} = \underset{c \in \mathbb{R}^k}{\operatorname{argmin}} \|Y - X V_k c\| = \underset{c \in \mathbb{R}^k}{\operatorname{argmin}} \|Y - U\Gamma V^T V_k c\|.$$

Note that

$$V^T V_k = \begin{bmatrix} I_k \\ 0 \end{bmatrix},$$

and so

$$\hat{c} = \underset{c \in \mathbb{R}^k}{\operatorname{argmin}} \|Y - U\Gamma V^T V_k c\| = \underset{c \in \mathbb{R}^k}{\operatorname{argmin}} \|Y - U_k \Gamma_k c\|,$$

where  $U_k = [u_1, \dots, u_k] \in \mathbb{R}^{n \times k}$  and  $\Gamma_k = \text{diag}(\gamma_1, \dots, \gamma_k)$ . Thus

$$\hat{c} = \Gamma_k^{-1} U_k^T Y,$$

and

$$\hat{\beta}_{\text{pcr}(k)} = V_k \Gamma_k^{-1} U_k Y.$$

Thus in PCR we project  $X$  onto the span of the first  $k$  left singular vector  $X$  and then fit  $Y$  as well as possible within that span. The predictions from PCR are thus

$$\begin{aligned} \hat{Y}_{\text{pcr}(k)} &= X \hat{\beta}_{\text{pcr}(k)} \\ &= U\Gamma V^T V_k \Gamma^{-1} U_k^T Y \\ &= U_k U_k^T Y \\ &= H_k Y. \end{aligned}$$

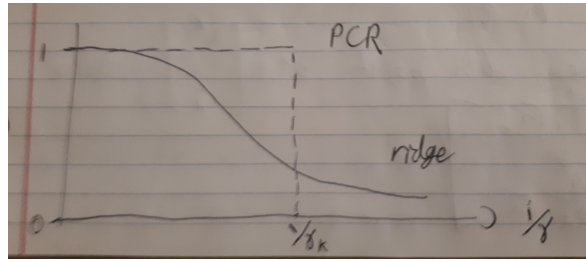
Thus

$$\hat{Y}_{\text{pcr}(k)} = H_k Y = \sum_{j=1}^k u_j u_j^T Y.$$

We can compare this to ridge regression

$$\hat{Y}_\lambda = \sum_{j=1}^d \frac{\gamma_j^2}{\gamma_j^2 + \lambda} u_j u_j^T Y.$$

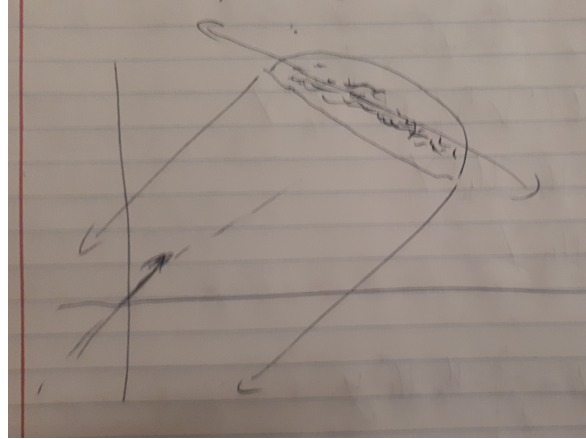
In ridge regression we apply a soft threshold to each of the singular direction  $u_j$  where we shrink by the factor  $\frac{\gamma_j^2}{\gamma_j^2 + \lambda}$  which increases with  $j$ . In PCR we have a hard threshold where we cut off the singular directions  $u_j$  for  $j > k$ . The thresholds look something like this:



**Remark 1.** A good question from the audience: what about the scale of the columns in

$$X = [x^{(1)}, \dots, x^{(d)}]?$$

In PCR we always standardize data so that  $\mathbb{E}[x^{(j)}] = 0$  and  $\text{Var}(x^{(j)}) = 0$ . This is to avoid situations like this



To do this we set  $\hat{\tau}_j^2 = \|x^{(j)}\|_2^2$  and replace  $x^{(j)}$  with  $\sqrt{n} \frac{x^{(j)}}{\hat{\tau}_j}$ . If we see a new data point  $x^* = [x_1^*, \dots, x_d^*]^T$ , to make predictions we have to first perform the transformation

$$x^* \mapsto \begin{bmatrix} \sqrt{n} \frac{x_1^*}{\hat{\tau}_1} \\ \vdots \\ \sqrt{n} \frac{x_d^*}{\hat{\tau}_d} \end{bmatrix}.$$

But sometimes we care about the scale of the variables and wouldn't want to standardize. This is a problem specific issue that can be hard. Boosting is a technique we will see later that can help. Boosting allows this process to be automated.

**Remark 2.** On the homework we are asked to compute the optimism/bias in PCR and ridge regressions. That is, what are the effective degrees of freedom in PCR and ridge?

## 4 Feature selection methods

Given  $X = [x^{(1)}, \dots, x^{(d)}] \in \mathbb{R}^{n \times d}$ , which features should we include? That is, which columns of  $X$  should we include in our regression?

A common approach is to pick different index subsets  $J \subseteq [d] = \{1, \dots, d\}$ . We then fit a model on  $X_J = [x^{(j)}]_{j \in J} \in \mathbb{R}^{n \times |J|}$ . We then ask if the fitted model is “good enough”. Our goal is to choose  $J$  to minimize errors on future data. There are two approaches for evaluating which subsets are “good enough”

- Hold out some data in a test set  $(x^{\text{test}}, y^{\text{test}})$  and evaluate the model on this test set. We will talk about this and variations such as permutation tests and cross validation in later lectures.
- Penalization approaches where we penalize the complexity of  $J$  using something like Mallows’s  $C_p$  statistic.

The notation we will use is

$$\widehat{\text{Risk}}(\beta) = \text{estimated error or risk on future data.}$$

The quantity  $\widehat{\text{Risk}}(\beta)$  will be the basis for which we select models. For example we may have

$$\widehat{\text{Risk}}(\beta) = \frac{1}{n} \|X\beta - Y\|_2^2 + \frac{2\hat{\sigma}^2 p}{n},$$

where  $p = \|\beta\|_0 = \# \text{ of non-zero entries}$ . Or we might have

$$\widehat{\text{Risk}}(\beta) = \frac{1}{n_{\text{test}}} \|X^{\text{test}}\beta - Y^{\text{test}}\|_2^2.$$

### 4.1 All subsets

In all subsets regression, we do the following:

- Look at all subsets  $J \subseteq [d]$ .
- Compute  $\widehat{\beta}_J = \arg\min_b \|X_J b - Y\|_2^2$ .
- Choose  $\widehat{\beta}$  to be the minimizer of  $\widehat{\text{Risk}}(\widehat{\beta}_J)$ .

There are some challenges to all subset regression.

- There are  $2^d$  such subsets which means it can be very expensive. (In practice, branch and bound methods can allow  $d \sim 1000$ ).
- Often the results are optimistic. Since we are searching over so many possible models, the models might overfit even with a good choice of  $\widehat{\text{Risk}}$ .
- The effectiveness of methods (especially when compared to convex optimization based approaches like ridge regression) depends a lot on  $\sigma^2 = \text{Var}(y|x)$ . If

$$\frac{\text{Var}(x^T \beta)}{\sigma^2},$$

is large (most of the variance is explained by  $x$ ), then all subsets is good. Otherwise all subsets is not so good.

- Recommended reading: this issue of Statistical Science on all subsets regression. See in particular the parts written by
  - Bertsimas,
  - Mazumdar,
  - Hastie/Tibshirani/Tibshirani.

## 4.2 Forward stepwise

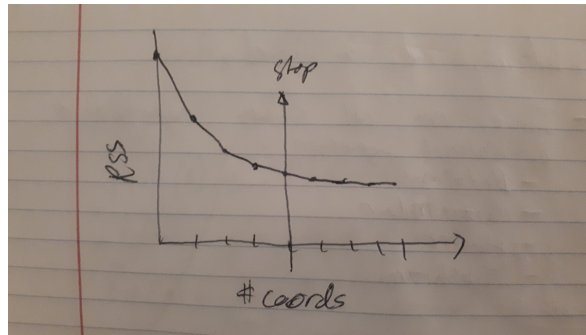
Consider the following greedy algorithm.

- Pick the feature  $j$  not in the model that makes the most progress. Add feature  $j$  to the model.
- Repeat.

For formally, Set  $J = \emptyset$ . Then for  $p = 1, 2, \dots$ :

- Define  $J_{+j} = J \cup \{j\}$  and  $r_j^2 = \min_b \|X_{J+j}b - y\|_2^2$  for  $j \notin J$ .
- Set  $j^* = \operatorname{argmin}_j r_j^2$ .
- Update  $J$  to  $J \cup \{j^*\}$ .

This is a reasonable approach and can be computed quickly. It is often effective even when  $d$  is large. The idea is that we would run this until the improvements get small. The plot of the residuals against  $|J|$  would look something like this:



When doing the forward stepwise algorithm we will need to make approximate  $d^2 \ll 2^d$  computations of  $r_j^2$ . Thus it is much better than a naive implementation of all subsets.

## 4.3 Backwards stepwise

Unfortunately the forward stepwise algorithm is a greedy algorithm and we might end up with some junk. This gives us the backwards stepwise algorithm.

- First fit using the full model (or the model produced by the forwards stepwise algorithm).
- Iteratively remove coordinates that increase RSS the least.

## 5 Boosting

This technique was originally developed in 1995 by Freund and Schapire in work on clustering.

It is designed to solve the challenge that we often do not know any of the following:

- What scales to use.
- What non-linear transformation to use.

Suppose we have a tool that can fit a small model well (a weak-learner). We then have the procedure:

- Start with raw data  $(x, y) \in \mathcal{X} \times \mathbb{R}$ .
- Iteratively
  - Find some feature mapping  $\phi : \mathcal{X} \rightarrow \mathbb{R}$  capturing some signal in  $x$  about  $y$ .
  - Add this feature to the model we're building.

More formally, for  $j = 1, 2, \dots$ , pick  $\phi^{(j)} : \mathcal{X} \rightarrow \mathbb{R}$  and add  $\phi^{(j)}$  to the model

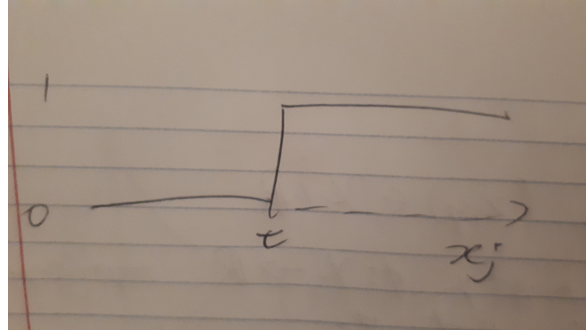
$$M(X) = \begin{bmatrix} \phi^{(1)}(x_1) & \phi^{(2)}(x_1) & \dots & \phi^{(j)}(x_1) \\ \phi^{(1)}(x_2) & \phi^{(2)}(x_2) & \dots & \phi^{(j)}(x_2) \\ \vdots & \vdots & \ddots & \vdots \\ \phi^{(1)}(x_n) & \phi^{(2)}(x_n) & \dots & \phi^{(j)}(x_n) \end{bmatrix} \in \mathbb{R}^{n \times d}.$$

Fit a model and repeat.

How do we actually do this? Roughly: coordinate descent. We first fix a family  $\Phi$  of feature mappings. For example we may use decision stumps for  $x \in \mathbb{R}^d$

$$\phi(x) = \begin{cases} 1 & \text{if } x_j \geq \tau, \\ 0 & \text{if } x_j < \tau. \end{cases}$$

That is  $\Phi$  consists of 0-1 thresholds on individual coordinates of raw data. The function  $\phi$  looks like this



The idea is that we start with  $M(X)$  and ask which feature improves the risk the most. That is we define

$$R(\delta, \phi; M) = \left\| [M, \phi] \begin{bmatrix} \beta \\ \delta \end{bmatrix} - y \right\|_2^2,$$

where  $\beta$  is the minimizer of  $\|Mb - y\|_2^2$ . There are then two approaches:

- Ask for

$$\hat{\delta} = \underset{\delta}{\operatorname{argmin}} R(\delta, \phi; M).$$

- Or ask what gives the most local progress

$$R'(\delta, \phi; M)|_{\delta=0}.$$

We will discuss this more on Thursday.