Supervised Learning: Regression, Part II

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Linear Models in High Dimensions

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Linear Models in High Dimensions

- ► When *p* is large, least squares regression will lead to very low training error but terrible test error.
- ▶ We will now see some approaches for fitting linear models in high dimensions, $p \gg n$.
- ▶ These approaches also work well when $p \approx n$ or n > p.

Motivating example

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- ► For instance, these biomarkers could be:
 - ▶ the expression levels of genes measured using a microarray.
 - protein levels.
 - mutations in genes potentially implicated in breast cancer.
- ► How can we develop a model with low test error in this setting?

Remember

- ► We have *n* training observations.
- Our goal is to get a model that will perform well on future test observations.
- ▶ We'll incur some bias in order to reduce variance.

Forward Stepwise Regression
Ridge Regression
Lasso Regression
Principal Components Regression

Variable Pre-Selection

The simplest approach for fitting a model in high dimensions:

- 1. Choose a small set of variables, say the q variables that are most correlated with the response, where q < n and q < p.
- 2. Use least squares to fit a model predicting *y* using only these *q* variables.

This approach is simple and straightforward.

Forward Stepwise Regression Ridge Regression Lasso Regression Principal Components Regression

Variable Pre-Selection in R

```
xtr <- matrix(rnorm(100*100),ncol=100)
beta <- c(rep(1,10),rep(0,90))
ytr <- xtr%*%beta + rnorm(100)
cors <- cor(xtr,ytr)
whichers <- which(abs(cors)>.2)
mod <- lm(ytr~xtr[,whichers])
print(summary(mod))</pre>
```

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- ▶ We want *q* that minimizes the test error.
- ► For a range of values of *q*, we can perform the validation set approach, leave-one-out cross-validation, or *K*-fold cross-validation in order to estimate the test error.

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- ► For a range of values of *q*, we can perform the validation set approach, leave-one-out cross-validation, or *K*-fold cross-validation in order to estimate the test error.
- ► Then choose the value of *q* for which the estimated test error is smallest.

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Estimating the Test Error For a Given q

This is the right way to estimate the test error using the validation set approach:

- 1. Split the observations into a training set and a validation set.
- 2. Using the training set only:
 - a. Identify the q variables most associated with the response.
 - b. Use least squares to fit a model predicting *y* using those *q* variables.
 - c. Let $\hat{\beta}_1, \dots, \hat{\beta}_q$ denote the resulting coefficient estimates.
- 3. Use $\hat{\beta}_1, \dots, \hat{\beta}_q$ obtained on training set to predict response on validation set, and compute the validation set MSE.

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Estimating the Test Error For a Given q

This is the wrong way to estimate the test error using the validation set approach:

- 1. Identify the *q* variables most associated with the response on the full data set.
- 2. Split the observations into a training set and a validation set.
- 3. Using the training set only:
 - a. Use least squares to fit a model predicting y using those q variables.
 - b. Let $\hat{\beta}_1, \dots, \hat{\beta}_q$ denote the resulting coefficient estimates.
- 4. Use $\hat{\beta}_1, \dots, \hat{\beta}_q$ obtained on training set to predict response on validation set, and compute the validation set MSE.

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Frequently Asked Questions

▶ **Q:** Does it really matter how you estimate the test error?

A: Yes.

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▶ **Q:** Does it really matter how you estimate the test error?

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▶ **Q:** Would anyone make such a silly mistake?

A: Yes.

A Better Approach

► The variable pre-selection approach is simple and easy to implement — all you need is a way to calculate correlations, and software to fit a linear model using least squares.

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A Better Approach

- ► The variable pre-selection approach is simple and easy to implement all you need is a way to calculate correlations, and software to fit a linear model using least squares.
- ▶ But it might not work well: just because a bunch of variables are correlated with the response doesn't mean that when used together in a linear model, they will predict the response well.
- ▶ What we really want to do: pick the *q* variables that best predict the response.

Best Subset Selection

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- ▶ Unfortunately, this is computationally intractable:
 - ▶ When p = 3, $2^p = 8$.
 - ▶ When p = 6, $2^p = 64$.
 - When p=250, there are $2^{250}\approx 10^{80}$ possible models. According to www.universetoday.com, this is around the number of atoms in the known universe.
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 - When p = 250, there are $2^{250} \approx 10^{80}$ possible models. According to www.universetoday.com, this is around the number of atoms in the known universe.
 - ▶ Not feasible to consider so many models!
- ► Need an efficient way to sift through all of these models: forward stepwise regression.

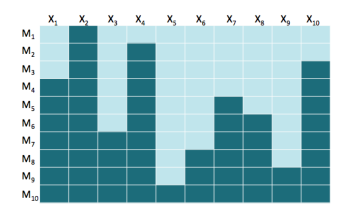
Forward Stepwise Regression

- Use least squares to fit p univariate regression models, and select the predictor corresponding to the best model (according to e.g. training set MSE).
- 2. Use least squares to fit p-1 models containing that one predictor, and each of the p-1 other predictors. Select the predictors in the best two-variable model.
- 3. Now use least squares to fit p-2 models containing those two predictors, and each of the p-2 other predictors. Select the predictors in the best three-variable model.
- 4. And so on....

This gives us a nested set of models, containing the predictors

$$\mathcal{M}_1 \subseteq \mathcal{M}_2 \subseteq \mathcal{M}_3 \subseteq \dots$$

Forward Stepwise Regression With p = 10



Example in R

```
xtr <- matrix(rnorm(100*100),ncol=100)
beta <- c(rep(1,10),rep(0,90))
ytr <- xtr%*%beta + rnorm(100)
library(leaps)
out <- regsubsets(xtr,ytr,nvmax=30,method="forward")
print(summary(out))
print(coef(out,1:10))</pre>
```

Which Value of *q* is Best?

- ► This procedure traces out a set of models, containing between 1 and *p* variables.
- ▶ The qth model contains q variables, given by the set \mathcal{M}_q .
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- ▶ The qth model contains q variables, given by the set \mathcal{M}_q .
- ▶ Q: Which value of q is best?A: The one that minimizes the test error!
- ► We can select the value of *q* using cross-validation or the validation set approach.

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- ▶ For instance, suppose that the best model with one variable is

$$y = \beta_3 X_3 + \epsilon$$

and the best model with two variables is

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▶ **Q:** Does this really happen in practice?

A: Yes.

How To Do Forward Stepwise?

Wrong: Split the data into a training set and a validation set. Perform forward stepwise on the training set, and identify the model with best performance on the validation set. Then, refit the model (using those q variables) on the full data set.

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Bottom Line: We estimate the test error in order to choose the correct level of **model complexity**. Then we refit the model on the full data set.

Let's Try It Out in R!

Chapter 6 R Lab, Part 1 www.statlearning.com

Ridge Regression and the Lasso

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- ► This is known as regularization or penalization.
- ▶ Hot area in statistical machine learning today.

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- ▶ When p > n, some of the variables are highly correlated.
- ► Why does correlation matter?
 - ▶ Suppose that X_1 and X_2 are highly correlated with each other... assume $X_1 = X_2$ for the sake of argument.
 - And suppose that the least squares model is

$$\hat{y} = X_1 - 2X_2 + 3X_3.$$

► Then this is also a least squares model:

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- ▶ Bottom Line: When there are too many variables, the least squares coefficients can get crazy!
- ► This craziness is directly responsible for poor test error.
- ▶ It amounts to too much model complexity.

A Solution: Don't Let the Coefficients Get Too Crazy

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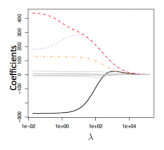
Ridge Regression

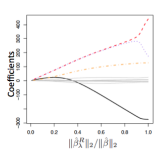
► Ridge regression coefficient estimates minimize

$$\|\mathbf{y} - \mathbf{X}\boldsymbol{\beta}\|^2 + \lambda \sum_j \beta_j^2.$$

- Here λ is a nonnegative tuning parameter that shrinks the coefficient estimates.
- ▶ When $\lambda = 0$, then ridge regression is just the same as least squares.
- ► As λ increases, then $\sum_{j=1}^{p} (\hat{\beta}_{\lambda,j}^{R})^2$ decreases i.e. coefficients become shrunken towards zero.
- When $\lambda = \infty$, $\hat{\boldsymbol{\beta}}_{\lambda}^{R} = 0$.

Ridge Regression As λ Varies





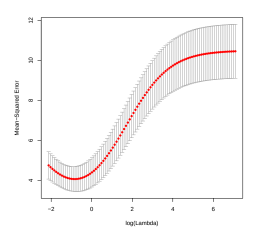
Ridge Regression In Practice

- ▶ Perform ridge regression for a very fine grid of λ values.
- Use cross-validation or the validation set approach to select the optimal value of λ – that is, the best level of model complexity.
- ▶ Perform ridge on the full data set, using that value of λ .

Example in R

```
xtr <- matrix(rnorm(100*100),ncol=100)
beta \leftarrow c(rep(1,10), rep(0,90))
ytr <- xtr%*%beta + rnorm(100)
library(glmnet)
cv.out <- cv.glmnet(xtr,ytr,alpha=0,nfolds=5)</pre>
print(cv.out$cvm)
plot(cv.out)
cat("CV Errors", cv.out$cvm,fill=TRUE)
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R Output



Drawbacks of Ridge

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- ► The lasso involves performing a little tweak to ridge regression so that the resulting model contains mostly zeros.
- ► In other words, the resulting model is sparse. We say that the lasso performs feature selection.
- ► The lasso is a very active area of research interest in the statistical community!

The Lasso

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subject to the constraint that

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▶ So lasso is just like ridge, except that β_j^2 has been replaced with $|\beta_i|$.

The Lasso

► Lasso is a lot like ridge:

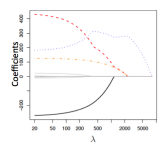
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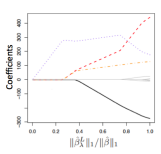
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- ▶ But unlike ridge, lasso will give some coefficients exactly equal to zero for intermediate values of λ !

Lasso As λ Varies





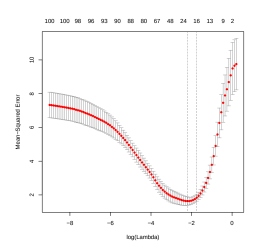
Lasso In Practice

- ▶ Perform lasso for a very fine grid of λ values.
- ► Use cross-validation or the validation set approach to select the optimal value of \(\lambda\) – that is, the best level of model complexity.
- ▶ Perform the lasso on the full data set, using that value of λ .

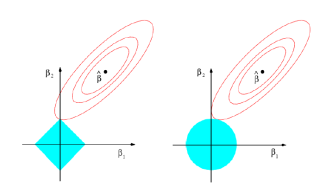
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R Output



Ridge and Lasso: A Geometric Interpretation



Let's Try It Out in R!

Chapter 6 R Lab, Part 2 www.statlearning.com

Review

- ► So far we have seen two approaches that select subsets of the features and fit a least squares model:
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Review

- ► So far we have seen two approaches that select subsets of the features and fit a least squares model:
 - ▶ Variable Pre-Selection
 - Forward Stepwise Selection
- ► And we have seen two approaches that fit a shrunken model instead of using least squares:
 - ► Ridge regression
 - ▶ Lasso
- ► Now we see one final approach, principal components regression, that first finds a low-dimensional subspace of the data and then fits a model on that low-dimensional subspace, using least squares.

Principal Components Regression

- ▶ Our data consist of *n* observations in a *p*-dimensional space.
- ► However, not all of those p dimensions are equally useful, especially when $p \gg n$.
- ► Many are either completely redundant (correlated features) or uninformative (noise features).
- ► Can we find a low-dimensional representation of the variables that captures most of the variability in the data?
- ► This is a dimension reduction approach.

PCR

► Let $Z_1, Z_2, ..., Z_M$ represent M < p linear combinations of the p predictors:

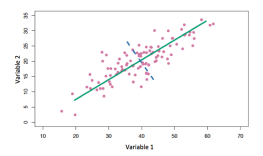
$$Z_m = \sum_{j=1}^p \phi_{mj} X_j.$$

▶ Use least squares to fit the model

$$y_i = \theta_0 + \sum_{m=1}^{M} \theta_m Z_{im} + \epsilon_i, \quad i = 1, \dots, n.$$

- ▶ In other words, we perform least squares using M new predictors, Z_1, \ldots, Z_M .
- $ightharpoonup Z_1, \ldots, Z_M$ chosen to be the principal components of the data.

Principal Components, Conceptually



- ► PCs are the linear combinations of the variables that contain as much as possible of the variability in the features.
- ▶ Will be discussed further in SISBID Module 5 Unsupervised Learning.

PCR

Our final model is linear in the original predictors:

$$y_{i} = \theta_{0} + \sum_{m=1}^{M} \theta_{m} Z_{im} + \epsilon_{i}$$

$$= \theta_{0} + \sum_{m=1}^{M} \theta_{m} \sum_{j=1}^{p} \phi_{mj} X_{ij} + \epsilon_{i}$$

$$= \theta_{0} + \sum_{j=1}^{p} \left(\sum_{m=1}^{M} \theta_{m} \phi_{mj} \right) X_{ij} + \epsilon_{i}$$

More on PCR

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- ► Shortcoming of PCR: the first *M* principal components are guaranteed to explain a lot of the variation in the features, but that doesn't mean that they are predictive of the response!
- ► In SISBID Module 5, will see how principal components can be used for unsupervised learning.

Example in R

```
xtr <- matrix(rnorm(100*100),ncol=100)
beta <- c(rep(1,10),rep(0,90))
ytr <- xtr%*%beta + rnorm(100)
library(pls)
out <- pcr(ytr~xtr,scale=TRUE,validation="CV")
summary(out)
validationplot(out,val.type="MSEP")</pre>
```

Let's Try It Out in R!

Chapter 6 R Lab, Part 3 www.statlearning.com

Pros/Cons of Each Approach

Approach	Simplicity?*	Sparsity?**	Predictions?***
Pre-Selection	Good	Yes	So-So
Forward Stepwise	Good	Yes	So-So
Ridge	Medium	No	Great
Lasso	Bad	Yes	Great
PCR	Medium	No	Great

^{*} How simple is this model-fitting procedure? If you were stranded on a desert island with pretty limited statistical software, could you fit this model?

^{**} Does this approach perform feature selection, i.e. is the resulting model sparse?

^{***} How good are the predictions resulting from this model?

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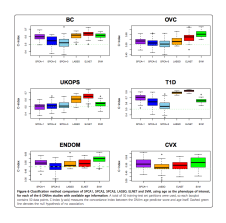
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 - ► Ridge will do better if all of the features are associated with the response.
- ► If somebody tells you that one approach is "best"... then they are mistaken. Politely contradict them.
- ► While no approach is "best", some approaches are wrong (e.g.: there is a wrong way to do cross-validation)!

Predicting Age Using DNA Methylation Data

- ► Comparison on 6 data sets
- ► SPC: Like principal components regression, but using a subset of features most associated with response. Between 1 and 3 principal components were used.
- ► Elastic Net: A hybrid between ridge and lasso.
- ► SVM: We'll see it next lecture in the classification context.
- ► Citation: Zhuang et al., BMC Bioinformatics, 2012

Didn't I Tell You? No Best Method!



High C-index indicates a low test error.

Bottom Line

Much more important than what model you fit is how you fit it.

- ▶ Was cross-validation performed properly?
- ► Did you select a model (or level of model complexity) based on an estimate of test error?