Classical Statistics Biological Big Data Supervised and Unsupervised Learning

High-Dimensional Statistical Learning: Introduction

Jean Feng & Ali Shojaie

November 15, 2020 Sixth Seattle Symposium in Biostatistics

1/24

Classical Statistics Biological Big Data Supervised and Unsupervised Learning

A Simple Example

- Suppose we have n = 400 people with diabetes for whom we have p = 3 serum-level measurements (LDL, HDL, GLU).
- We wish to predict these peoples' disease progression after 1 year.





























Want to develop a model of the form

$$y_i = \beta_0 + \beta_1 X_{i1} + \cdots + \beta_p X_{ip} + \epsilon_i.$$

- Here ϵ_i is a noise term associated with the *i*th observation.
- Must estimate $\beta_0, \beta_1, \ldots, \beta_p$ i.e. we must fit the model.



$$y_i = \beta_0 + \beta_1 X_{i1} + \cdots + \beta_p X_{ip} + \epsilon_i.$$

Most common approach in classical statistics is least squares:

$$\underset{\beta}{\text{minimize}} \sum_{i=1}^{n} (y_i - (\beta_1 X_{i1} + \dots + \beta_p X_{ip}))^2$$

• We are looking for β_1, \ldots, β_p such that

$$\sum_{i=1}^{n}(y_i-(\beta_1X_{i1}+\cdots+\beta_pX_{ip}))^2$$

is as small as possible, or in other words, such that

$$\sum_{i=1}^n (y_i - \hat{y}_i)^2$$

is as small as possible, where \hat{y}_i is the *i*th predicted value.

Least Squares Regression ▶ When we fit a model, we use a training set of observations. • We get coefficient estimates $\hat{\beta}_1, \ldots, \hat{\beta}_n$. ▶ We also get predictions using our model, of the form $\hat{\mathbf{y}}_i = \hat{\beta}_1 X_{i1} + \dots + \hat{\beta}_n X_{in}.$ ▶ We can evaluate the training error, i.e. the extent to which the model fits the observations used to train it. One way to quantify the training error is using the mean squared error (MSE): $MSE = \frac{1}{n}\sum_{i=1}^{n}(y_i - \hat{y}_i)^2 = \frac{1}{n}\sum_{i=1}^{n}(y_i - (\hat{\beta}_1 X_{i1} + \dots + \hat{\beta}_p X_{ip}))^2.$ • The training error is closely related to the R^2 for a linear model — that is, the proportion of variance explained. • Big $R^2 \Leftrightarrow$ Small Training Error. 7 / 40 Least Squares as More Variables are Included in the Model • Training error and R^2 are not good ways to evaluate a model's performance, because they will always improve as more variables are added into the model. ▶ The problem? Training error and R^2 evaluate the model's performance on the training observations.

- If I had an unlimited number of features to use in developing a model, then I could surely come up with a regression model that fits the training data perfectly! Unfortunately, this model wouldn't capture the true signal in the data.
- We really care about the model's performance on test observations — observations not used to fit the model.







Bias and Variance

- As model complexity increases, the bias of β̂ the average difference between β and β̂, if we were to repeat the experiment a huge number of times will decrease.
- But as complexity increases, the variance of $\hat{\beta}$ the amount by which the $\hat{\beta}$'s will differ across experiments will increase.
- ► The test error depends on both the bias and variance:

Test Error =
$$Bias^2 + Variance$$
.

There is a bias-variance trade-off. We want a model that is sufficiently complex as to have not too much bias, but not so complex that it has too much variance.











^{22 / 40}



Leave-One-Out Cross-Validation

For a given model, we perform the following procedure:

- 1. For i = 1, ..., n:
 - a. Fit the model using observations $1, \ldots, i 1, i + 1, \ldots, n$. Let $\hat{\beta}_{(i)}$ denote the regression coefficient estimates.
 - b. Compute the test error, $e_i = (y_i x_i^T \hat{\beta}_{(i)})^2$.
- 2. Calculate $\sum_{i=1}^{n} e_i$, the total CV error.

Out of a set of candidate models, the "best" one is the one for which the total error is smallest.

5-Fold Cross-Validation

Split the observations into 5 sets. Repeatedly train the model on 4 sets and evaluate its performance on the 5th.



K-fold cross-validation

A generalization of leave-one-out cross-validation. For a given model, we perform the following procedure:

- 1. Split the n observations into K equally-sized folds.
- 2. For k = 1, ..., K:
 - a. Fit the model using the observations not in the kth fold.
 - b. Let e_k denote the test error for the observations in the *k*th fold.
- 3. Calculate $\sum_{k=1}^{K} e_k$, the total CV error.

Out of a set of candidate models, the "best" one is the one for which the total error is smallest.

After Estimating the Test Error on the Training Set...

After we estimate the test error using the training set, we refit the "best" model on all of the available training observations. We then evaluate this model on the test set.







Why All the Bother?

Q: Why do I need to have a separate test set? Why can't I just estimate test error using cross-validation or a validation set approach using all the observations, and then be done with it?

A: In general, we are choosing between a whole bunch of models, and we will use the cross-validation or validation set approach to pick between these models. If we use the resulting estimate as a final estimate of test error, then this could be an extreme underestimate, because one model might give a lower estimated test error than others by chance. To avoid having an extreme underestimate of test error, we need to evaluate the "best" model obtained on an independent test set. *This is particularly important in high dimensions!!*

Regression in High Dimensions

- We usually cannot perform least squares regression to fit a model in the omics setting, because we will get zero training error but a terrible test error.
- Instead, we must fit a less complex model, e.g. a model with fewer variables.

If you

- fit your model carelessly;
- do not properly estimate the test error;
- or select a model based on training set rather than test set performance;

then you will woefully overfit your training data, leading to a model that looks good on training data but will perform atrociously on future observations.

Our intuition breaks down in high dimensions, and so rigorous model-fitting is crucial.



The Curse of Dimensionality

 ${\bf Q}$: A data set with more variables is better than a data set with fewer variables, right?

A: Not necessarily!

Noise variables – such as genes whose expression levels are not truly associated with the response being studied – will simply increase the risk of overfitting, and the difficulty of developing an effective model that will perform well on future observations.

On the other hand, more signal variables – variables that are truly associated with the response being studied – are always useful!

Wise	Words
VV15C	110103

In high-dimensional data analysis, common mistakes are simple, and simple mistakes are common.

- Keith Baggerly

Before You're Done Your Analysis Estimate the test error. Do a "sanity check" whenever possible. "Spot-check" the variables that have the largest coefficients in the model. Rewrite your code from scratch. Do you get the same answer again? Fitting models in high-dimensions: one mistake away from disaster!



Variable Pre-Selection Subset Selection		
Ridge Regression		
Lasso Regression		
Motivating example		
We would like to build a model to predict survival time for		
breast cancer patients using a number of clinical		
measurements (tumor stage, tumor grade, tumor size, patient		
age, etc.) as well as some high-dimensional biomarkers.		
For instance, these biomarkers could be		
 Tor instance, these biomarkers could be. The surgrassion levels of neuros 		
the expression levels of genes		
protein levels.		
mutations in genes potentially implicated in breast cancer.		
	3/33	
Variable Pre-Selection	3/33	
Variable Pre-Selection Subset Selection Ridge Regression	3/33	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression	3/33	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression	3/33	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	^{3/33} a?	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	a?	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	a?	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	^{3/33} a?	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	^{3/33} a?	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	^{3/33} a?	
Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression What are our options for analyzing high-dimensional data	^{3/33} a?	
 Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection 	^{3/33} a?	
 Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression 	^{3/33} a?	
Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression	^{3/33} a?	
 Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	
 Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	
 Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	
 Variable Pre-Selection Subset Selection Lasso Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	
 Variable Pre-Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	
 Variable Pre-Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	
 Variable Pre-Selection Subset Selection Ridge Regression What are our options for analyzing high-dimensional data Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression 	^{3/33} a?	

	Variable Pre-Selection	
	Subset Selection Ridge Regression	
	Lasso Regression	
Variable Pre-9	Selection	
A simple app	roach:	
1. Choose a	a small set of variables, say the q variables that are	
most cor	related with the response, where $q < n$.	
2. Use least	squares to fit a model predicting γ using only these	
q variabl	es.	
		5 / 33
	Variable Pre-Selection Subset Selection	
	Ridge Regression Lasso Regression	
Variable Pre-S	election: Bias-Variance Trade-off	
Variable pre-s	election tries to find the right trade-off between the	
bias and varia	ince.	
	High Bias Low Bias	
or	Low Variance High Variance	
E		
	q=4	
ion]	Least Squares	
ediction]	q=4 Least T Squares Test Sample Image: Control of the second seco	
Prediction]	q=4 Test Sample	
Prediction]	q=4 Least Test Sample	
Prediction]	Training Sample	
Prediction 1	Training Sample	
Prediction 1	Low High	


Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression

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 $\hat{\beta}_1, \ldots, \hat{\beta}_q$ denote the resulting coefficient estimates.
- 4. Use $\hat{\beta}_1, \ldots, \hat{\beta}_q$ obtained on training set to predict response on validation set, and compute the validation set MSE.

Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression

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 $\hat{\beta}_1, \ldots, \hat{\beta}_q$ denote the resulting coefficient estimates.
- 3. Use $\hat{\beta}_1, \ldots, \hat{\beta}_q$ obtained on training set to predict response on validation set, and compute the validation set MSE.







$$\hat{\boldsymbol{\beta}}_{\lambda}^{R} = \underset{\beta}{\operatorname{argmin}} \sum_{i=1}^{n} (y_{i} - (\beta_{1}X_{i1} + \dots + \beta_{p}X_{ip}))^{2} + \lambda \|\beta\|_{2}^{2}.$$

Here λ is a nonnegative tuning parameter that shrinks the coefficient estimates.

- When λ = 0, then ridge regression is just the same as least squares.
- As λ increases, coefficients shrink towards zero.
- When $\lambda = \infty$, $\hat{\boldsymbol{\beta}}_{\lambda}^{R} = 0$.







- When λ is very large, we get $\hat{\beta}_{\lambda}^{L} = 0$.
- But unlike ridge, lasso will give some coefficients exactly equal to zero for intermediate values of λ!







Ridge Regression Lasso Regression

Modeling non-linear relationships

What if the relationship isn't linear?

$$y = 3\sin(x) + \epsilon$$
$$y = 2e^{x} + \epsilon$$
$$y = 3x^{2} + 2x + 1 + \epsilon$$

If we know the functional form we can still use "linear regression"

Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression

Linear regression for non-linear functions

$$y = 3\sin(x) + \epsilon$$
:
 $\left(x\right) \rightarrow \left(\sin(x)\right)$

$$y = 3x^2 + 2x + 1 + \epsilon$$

$$\left(x\right) \rightarrow \left(x \mid x^2\right)$$

29 / 33

Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression

Linear regression for non-linear functions

What if we don't know the right functional form?

Use a flexible basis expansion h_1, h_2, \ldots, h_k :

$$\begin{pmatrix} x \\ \end{pmatrix} \rightarrow \begin{pmatrix} h_1(x) \\ h_2(x) \end{pmatrix} \cdots \begin{pmatrix} h_k(x) \end{pmatrix}$$

For example, the polynomial basis:

$\begin{pmatrix} x \end{pmatrix}$	\rightarrow	×	<i>x</i> ²	• • •	$\left(x^{k}\right)$



Variable Pre-Selection Subset Selection Ridge Regression Lasso Regression	
Summary	
You can use variable pre-selection or subset selection to limit the number of variables you feed into ordinary least squares.	
 Ridge regression protects against overfitting by shrinking model coefficients to zero. 	
 Lasso regression protects against overfitting by setting some model coefficients to exactly zero. 	
 To model non-linear relationships, expand the features using a basis expansion. 	a
	33 / 33

Classification Performance Assessment Batch Effects

High-Dimensional Statistical Learning: Classification

Jean Feng & Ali Shojaie

November 15, 2020 Sixth Seattle Symposium in Biostatistics



Classification Performance Assessment Batch Effects	
Classification	
 There are many approaches out there for performing classification. 	
We will discuss two, logistic regression and support vector machines	
machines.	
	3 / 30
	3/30
Classification	
Performance Assessment Batch Effects	
Performance Assessment Batch Effects Logistic Regression	
 Performance Assessment Batch Effects Logistic Regression ► Logistic regression is the straightforward extension of linear regression to the classification setting. 	
 Performance Assessment Batch Effects Logistic Regression Logistic regression is the straightforward extension of linear regression to the classification setting. For simplicity, suppose y ∈ {0,1}: a two-class classification problem 	
 Performance Assessment Batch Effects Logistic Regression Logistic regression is the straightforward extension of linear regression to the classification setting. For simplicity, suppose y ∈ {0,1}: a two-class classification problem. The logistic regression model is defined as 	
Performance Assessment Batch Effects Logistic Regression • Logistic regression is the straightforward extension of linear regression to the classification setting. • For simplicity, suppose $y \in \{0, 1\}$: a two-class classification problem. • The logistic regression model is defined as $f(X) = \frac{\exp(X^T \beta)}{1 + \exp(X^T \beta)}.$	
Performance Assessment Batch Effects Logistic Regression • Logistic regression is the straightforward extension of linear regression to the classification setting. • For simplicity, suppose $y \in \{0, 1\}$: a two-class classification problem. • The logistic regression model is defined as $f(X) = \frac{\exp(X^T \beta)}{1 + \exp(X^T \beta)}.$	
Performance Assessment Batch Effects Logistic Regression • Logistic regression is the straightforward extension of linear regression to the classification setting. • For simplicity, suppose $y \in \{0, 1\}$: a two-class classification problem. • The logistic regression model is defined as $f(X) = \frac{\exp(X^T \beta)}{1 + \exp(X^T \beta)}.$ • Logistic regression solves the following problem:	
Performance Assessment Batch Effects Logistic Regression • Logistic regression is the straightforward extension of linear regression to the classification setting. • For simplicity, suppose $y \in \{0, 1\}$: a two-class classification problem. • The logistic regression model is defined as $f(X) = \frac{\exp(X^T\beta)}{1 + \exp(X^T\beta)}.$ • Logistic regression solves the following problem: $\min_{\beta} \sum_{i=1}^{n} \log(1 + \exp(-f_{\beta}(x_i)y_i))$	



























Decision Trees Bagging Random Forest Boosting

High-Dimensional Statistical Learning: Tree-Based Methods

Jean Feng & Ali Shojaie

November 15, 2020 Sixth Seattle Symposium in Biostatistics















6.549

6.407

RBI < 80.5

7 007

7.289

Years < 6.5

6.459

Runs < 47.5

5 571

6 015

RBI < 60.5

5 183

Years < 3.5

5.394

Puto

5 487



Decision Trees Bagging Random Forest Boosting Regression vs Classification Trees Regression Classification • Predict mean • Predict mode • Evaluate split by improvement in mean squared error: how different are the outcomes from the mean for that split • Predict mode • Imaging Classification • Predict mode • Classification • Predict mode • Classification • Classification							
		15 / 30					
Decision Trees Bagging Random Forest Boosting							
Classification Tree for the Heart Data							
9 9 9 9 9 9 10 10 10 10 10 10 10 10 10 10	Theia Image: Carrier of the strain bc MaxtHR < 161.5	16 / 30					
















Machine Learning for Biomarker Development

High-Dimensional Statistical Learning: Biomarkers

Jean Feng & Ali Shojaie

November 15, 2020 Sixth Seattle Symposium in Biostatistics

1 / 18

Machine Learning for Biomarker Development

The Data

On each of *n* patients measure

 y_i – outcome

(eg. tumor growth, treatment response, survival time)

 $x_i - p$ -vector of features

(eg. SNPs, gene expression values)

2 / 18



1.0

0.8

0.6

0.0

0.2

0.4

biomark



1.0

0.8

0.2

0.4

biom

0.6

0.0



5 / 18

Machine Learning for Biomarker Development

Prognostic Biomarkers

How do we characterize E[y|x]?

Statistical Machine Learning!!

Everything developed so far aims at this problem.

6 / 18



Machine Learning for Biomarker Development

Predictive Biomarkers — Cox

For survival data, using Cox Model, need to be a bit more careful.

Assumes hazard factors as

$$H(time, x) = h(time)f(x)$$

- Decisions are generally made based on f(x)
- shared baseline hazard, h(time), often considered a nuisance



Predictive Biomarkers — Cox

Instead estimate a single $h_{shared}(time)$:

$$\begin{bmatrix} H_{\mathsf{T}}(time, x) \\ H_{\mathsf{C}}(time, x) \end{bmatrix} \longrightarrow \begin{bmatrix} \hat{h}_{\mathsf{shared}}(time) \hat{f}_{\mathsf{T}}(x) \\ \hat{h}_{\mathsf{shared}}(time) \hat{f}_{\mathsf{C}}(x) \end{bmatrix}$$

and make inference on

 $\hat{f}_{T}(x)$ vs $\hat{f}_{C}(x)$

10 / 18









Cross Validation

- 1. Split data into folds: $fold_1, \ldots, fold_K$.
- 2. Cycle through the folds; for each k train and evaluate:

$$folds_{-k} \longrightarrow \begin{pmatrix} \hat{\mathsf{E}}_{\mathcal{T}} [y|x] \\ \hat{\mathsf{E}}_{\mathcal{C}} [y|x] \end{pmatrix}$$
apply $\begin{pmatrix} \hat{\mathsf{E}}_{\mathcal{T}} [y|x] \\ \hat{\mathsf{E}}_{\mathcal{C}} [y|x] \end{pmatrix}$ to $folds_k \longrightarrow Z_{1,k}, \dots, Z_{n_k,k}$

- 3. Have a cross-validated Z_i for each observation
- 4. For obs with $Z_i > 0$, compare treatment to control.



Permutation Test

- 1. Run CV-procedure and calculate a *t*-statistic, T, for obs with $Z_i > 0$
- 2. For b = 1, ..., B:
 - 2.1 Permute the treatment labels
 - 2.2 Run the CV procedure on permuted data to get permuted scores, $Z_i^{(b)}$
 - 2.3 Calculate a *t*-statistic, T_b , comparing permuted treatment to control for those with $Z_i^{(b)} > 0$
- 3. Compare T to empirical distribution of $\{T_b\}_{b=1}^B$

In clinical trials, known as "cross-validated adaptive signature design"