Data Mining W4240 Section 001

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Outline

Reviewing Estimators

Cross Validation

Data Preprocessing

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Cross Validation

Data Preprocessing

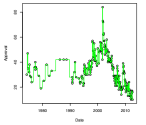
Some Estimators

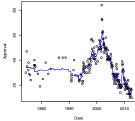
Recall:

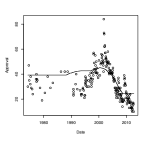
► kNN:

$$\hat{y} = \frac{1}{k} \sum_{i: x_i \in N_k(x)} y_i$$

▶ *k* controls the tradeoff between neighborhood size (bias) and estimator noise (variance)







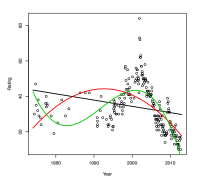
Some Estimators

Recall:

► Polynomial regression:

$$\hat{y} = \beta_0 + \beta_1 x + \beta_2 x^2 + \dots + \beta_m x^m$$

 Maximal degree m controls the tradeoff between estimator flexibility (bias) and estimator noise (variance)



Tunable Parameters

m and k are called **tunable parameters**

Their values affect how well a method works

- ▶ the "right" value should minimize the bias-variance tradeoff
- ...but the "right" value depends on the data

So how do we find the "right" value for a given dataset?

This problem is called model selection

Some More Estimators

Let's go back to kNN:

- suppose that I have chosen k and I now have an estimator
- ▶ I want to know how good this estimator is (i.e. what is the error on a new dataset?)

How would I estimate that?

This problem is called model assessment

Traditional Statistics



Ronald Aylmer Fisher (1890-1962)¹

- ▶ 1. study problem, 2. propose model, 3. fit model, 4. check assumptions, 5. go back to 2. if assumptions not met
- tunable parameters changed to meet assumptions (e.g. Gaussian residuals)
- ▶ if assumptions met, theoretical properties describe model behavior on new datasets (e.g. confidence intervals)

¹Photo credit: Wikipedia

Data Mining



Data Mining (Larry Page and Sergey Brin)²

- ▶ 1. look at data, 2. propose model, 3. select tunable parameters, 4. fit model, 5. assess model
- data most likely does not meet assumptions for parametric models
- usually care more about prediction than inference

²Photo credit: money.cnn.com

Generalization

Modeling for prediction:

- 1. get data
- 2. choose a model
- 3. fit the model
- 4. make predictions for new data

Generalization: making high quality predictions for new data

Expected Predictive Error

Tunable parameters α

Model with parameters α , $\hat{f}_{\alpha}(x)$

Goals for expected predictive error:

- ▶ **Model selection:** estimating the performance of different models in order to choose the best one (best α).
- ► Model assessment: having chosen a final model, estimating its prediction error (generalization error) on new data.

Expected Predictive Error

Training data: $\mathcal{T} = \{(x_1, y_1), \dots, (x_n, y_n)\}$

New data: X^0 , Y^0

Generalization error:

$$\operatorname{Err}_{\mathcal{T}} = \mathbb{E}_{X^0, Y^0} [L(Y^0, \hat{f}(X^0)) \mid \mathcal{T}]$$

Here, L(Y, f(X)) is a **loss function**

for regression: usually, the loss is the squared error

$$L(Y, f(X)) = (Y - f(X))^2$$

 for classification: usually, the loss is the Hamming distance (misclassification)

$$L(Y, f(X)) = \mathbf{1}_{\{Y \neq f(X)\}}$$

Outline

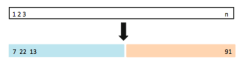
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Validation Sets

We could use a **validation set**³. Here n = 392 = 196 + 196



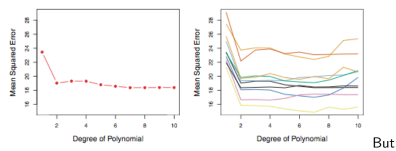
- randomly divide the data into a training set (blue) and a validation set (beige)
- ightharpoonup fit model on training set with differing values of lpha
- evaluate models performance on the validation set, and pick the best
- fit model on entire dataset

Doesn't use data twice!

³Some images from *An Introduction to Statistical Learning* by James, Witten, Hastie and Tibshirani

Validation Sets

Auto data set. Left: validation error for a single split. Right: validation error repeated 10 times (different random splits)



there are two problems:

- estimates depend heavily on the validation set (<u>high variance</u>)
- estimate of error is probably higher than error for full model

Both of these problems get worse with small n

What if we use the data twice?

- ▶ introduce a lot of bias if we fit model on (x_i, y_i) and then use $L(y_i, \hat{f}_{\alpha}(x_i))$ to estimate error
- ▶ introduce a *little* bias if we don't fit model on (x_i,y_i) , use $L(y_i,\hat{f}_{\alpha}(x_i))$ to select α , and then fit \hat{f}_{α} on training <u>and</u> validation sets

Let's try:

- use only **one** element as a validation set (all the rest as a training set)
- do this for all elements
- average results

Leave-One-Out-Cross-Validation

1 2 3		n
	1	
1 2 3		n
1 2 3		n
1 2 3		n
	:	
1 2 3		n

$$\operatorname{Err}_{\mathcal{T}} = \mathbb{E}_{X^{0}, Y^{0}}[L(Y^{0}, \hat{f}_{\alpha}(X^{0})) \mid \mathcal{T}]$$

$$\approx \frac{1}{n} \sum_{i=1}^{n} L(y_{i}, \hat{f}_{\alpha}^{(-i)}(x_{i}))$$

Leave-One-Out-Cross-Validation

Model Selection:

1. For all values of α , estimate generalization error with <u>LOOCV</u>:

$$\operatorname{Err}_{\alpha}^{LOOCV} = \frac{1}{n} \sum_{i=1}^{n} L(y_i, \hat{f}_{\alpha}^{(-i)}(x_i))$$

- 2. Choose α with lowest $\mathrm{Err}_{\alpha}^{LOOCV}$ (if many are similar, choose one in the middle of the range): let's call it α^*
- 3. Fit \hat{f}_{α^*} to $(x_1, y_1), \dots, (x_n, y_n)$

Model Assessment:

- 1. For a fixed value of α , estimate $L_{\alpha}^{(-i)} = L(y_i, \hat{f}_{\alpha}^{(-i)}(x_i))$
- 2. Can use $\{L_{\alpha}^{(-i)}\}_{i=1}^n$ to approximate distribution of predictive loss for given model

Generalized Cross-Validation

<u>Super fun fact</u>: for *ordinary least squares linear regression* we actually only need to fit the model once

$$\operatorname{Err}^{LOOCV} = \frac{1}{n} \sum_{i=1}^{n} \left(y_i - \hat{f}^{(-i)}(x_i) \right)^2$$
$$= \frac{1}{n} \sum_{i=1}^{n} \left(\frac{y_i - \hat{f}(x_i)}{1 - H_{ii}} \right)^2$$
$$\approx \frac{1}{n} \sum_{i=1}^{n} \left(\frac{y_i - \hat{f}(x_i)}{1 - \operatorname{trace}(\mathbf{H})/n} \right)^2$$

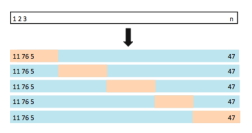
where $\hat{y} = \mathbf{H}y$, so $\mathbf{H} = X(X^TX)^{-1}X^T$. Note that $H_{ii} = h_i$, the leverage statistic.

Leave-One-Out-Cross-Validation

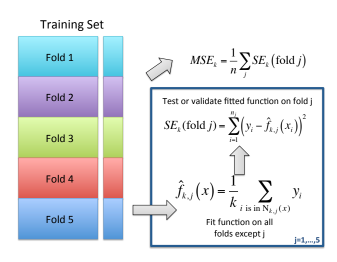
LOOCV sounds great! What could go wrong?!?

- suppose that I am training f on the entire Wikipedia corpus (> 3 million articles). I am a good coder, but it still takes about 30 hours to do one fit. Would LOOCV work? How could we fix it?
- ▶ suppose that $\alpha = (\alpha_1, \alpha_2, \dots, \alpha_m)$. How would LOOCV change? Would it work, and if not, how could we fix it?

k-Fold Cross-Validation (here k = 5)

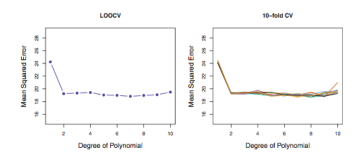


- 1. separate training set into K different, equally sized sets (folds)
- 2. for each tunable parameter value $\alpha = \alpha_1, \dots, \alpha_M$:
 - for k = 1, ..., K:
 - use all of the data except fold k as a training set to fit the function with parameter α
 - ightharpoonup use fold k as a testing set
 - ightharpoonup estimate squared error on fold k
 - average errors to approximate expected predictive error
- 3. compare error values; pick parameter with lowest error



k-Fold Cross-Validation

Auto data set: predicting mpg using polynomial functions of horsepower. Left: The LOOCV error curve (k=n). Right: 9 (slightly different) CV error curves, obtained with a 10-fold $(\underline{k=10})$ CV which was run 9 separate times, each with a different random split of the data into ten parts.



It turns out that K-fold CV is almost as good as LOOCV

k-Fold Cross-Validation

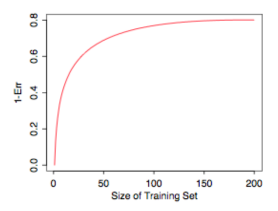
$$CV_{(n)} = \frac{1}{n} \sum_{i=1}^{n} MSE_{i}$$

$$CV_{(k)} = \frac{1}{k} \sum_{i=1}^{k} MSE_{i}$$

K-Fold Cross-Validation

How big should K be?

- lacktriangle need to do computation K times
- ▶ ...but larger K is more accurate estimator



K-Fold Cross-Validation

Why is 10-fold CV so close to LOOCV?

Intuition: Central Limit Theorem

- ▶ n iid random variables, $\mathbb{E}[X_i] = \mu$, $Var(X_i) = \sigma^2 < \infty$
- $\bar{\mu} = \frac{1}{n} \sum_{i=1}^{n} X_i$
- ▶ By CLT, $\bar{\mu} \sim N(\mu, \sigma^2/n)$
- So that means

$$|\bar{\mu} - \mu| \approx C \frac{1}{\sqrt{n}}$$

It turns out that $\frac{C}{\sqrt{n}}$ is the fastest we can expect an estimator to converge

- ▶ n = 500: Expected error for 5-fold CV is 0.05, for 10-fold CV is 0.047 and for LOOCV is 0.045
- ▶ n = 5,000: Expected error for 5-fold CV is 0.016, for 10-fold CV is 0.015 and for LOOCV is 0.014

Cross-Validation on Classification Problems

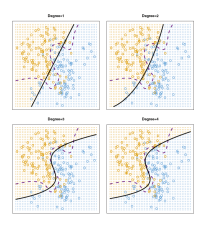
$$CV_{(n)} = \frac{1}{n} \sum_{i=1}^{n} Err_{i}$$

$$ERR_{(i)} = \mathbf{1}_{\{u_{i} \neq \hat{u}_{i}\}}$$

For example, fit a quadratic logistic regression model

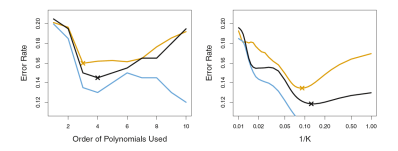
$$\log\left(\frac{p(X)}{1 - p(X)}\right) = \beta_0 + \beta_1 X_1 + \beta_2 X_1^2 + \beta_3 X_2 + \beta_4 X_2^2$$

Cross-Validation on Classification (simulated data)



Test errors rates: 0.2, 0.197, 0.160, 0.612. Bayes error rate: 0.133. In practice, for real data, the Bayes decision boundary and the test error rates are unknown. So how might we decide between the four logistic regression models? We can use cross-validation!

Cross-Validation on Classification Problems



True Test error (brown), training error (blue), and 10-fold CV error (black) on the two-dimensional classification data displayed before. Left: Logistic regression using polynomial functions of the predictors. Right: The $k{\rm NN}$ classifier with different values of k, the number of neighbors used in the KNN classifier.

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Data Preprocessing

Common data problems:

- missing data: not all data has all values
- high dimensional: p is too large to effectively use most methods

To deal with these problems, we often do preprocessing

- missing data: remove or impute values
- high dimensional: reduce dimensionality

Data Preprocessing

Ways to reduce dimensionality:

- select a set of covariates that are "highly predictive"
 - highly correlated with response
 - have large marginal information gains/variance reductions
- ▶ make some combination(s) of the covariates (like $0.7X_1 3.8X_2$) that is "highly predictive"

Finding a set of predictors before fitting an estimator is called **screening**.

The simplest method is selecting the K covariates that are the most correlated with Y.

Consider a problem with many predictors (ex: microarray data). A possible strategy:

- 1. Screen the predictors to find a "good" subset (e.g. choose the best subset based on data)
- 2. Using this set of predictors, build a multivariate classifier
- 3. Use cross-validation to select tunable parameters and estimate model error

Is this the right way to use cross-validation?

Consider this classification problem:

- n = 50, p = 5000
- $X \sim N_{5000}(0, I), Y \sim Bernoulli(0.5)$
- ▶ true error rate: 50% (labels independent from covariates)

Let's screen to select 100 most predictive covariates and then use 1-nn prediction.

What happens?

Right way to do cross-validation:

- 1. Divide samples into K cross-validation folds at random
- 2. For each fold $k = 1, \ldots, K$
 - a. Reserve fold k for test, use other folds for training
 - b. Find a subset of "good" predictors from training
 - Using this subset of predictors, build a multivariate classifier on training set
 - d. Use the classifier to predict labels on fold k

Your screening method is a part of your model!

See ESL, Ch. 7.10.2 on page 245

