STAT406- Methods of Statistical Learning Lecture 4

Matias Salibian-Barrera

UBC - Sep / Dec 2017

Proper use of CV

 An example of the importance and relevance of what we discussed in our last class:

Ambroise, C. and McLachlan, G.J. Selection bias in gene extraction on the basis of microarray gene-expression data, PNAS, 2002, 99 (10), 6562-6566.

https://doi.org/10.1073/pnas.102102699

Discussion points

- Why? Why would anybody want to not use all available features?
- "Somewhat obvious": model parsimony, identify features that are relevant for the process under study.
- "Not so obvious?": does prediction suffer if we use fewer variables? how much variability is induced by the feature selection step?



• Simple example:

```
set.seed(123)
x1 <- rnorm(506)
x2 <- rnorm(506, mean=2, sd=1)
x3 <- rexp(506, rate=1)
x4 <- x2 + rnorm(506, sd=.1)
x5 <- x1 + rnorm(506, sd=.1)
x6 <- x1 - x2 + rnorm(506, sd=.1)
x7 <- x1 + x3 + rnorm(506, sd=.1)
y <- x1*3 + x2/3 + rnorm(506, sd=2.2)</pre>
```

 Variables X₁ and X₂ are clearly important. But they are also highly correlated to X₄, X₅, X₆ and X₇.

 However, nothing is significant? > summary(lm(v~., data=x)) Call: $lm(formula = v \sim ., data = x)$ Residuals: Min 10 Median 30 Max -6.882 -1.474 -0.033 1.415 5.823 Coefficients: Estimate Std. Error t value Pr(>|t|) (Intercept) 0.03457 0.23018 0.150 0.8807 x 1 3.22612 1.68088 1.919 0.0555 . x2 0.23867 1.39355 0.171 0.8641 -0.35926 0.98680 -0.364 0.7160 x3 x4 -0.69359 0.99025 -0.700 0.4840 0.09271 0.91162 0.102 0.9190 x 5 -0.73887 1.01114 -0.731 0.4653 x 6 x7 0.31651 0.98610 0.321 0.7484 Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1 Residual standard error: 2.148 on 498 degrees of freedom Multiple R-squared: 0.6353, Adjusted R-squared: 0.6302 F-statistic: 123.9 on 7 and 498 DF, p-value: < 2.2e-16

But...

```
> summary(lm(v \sim x1 + x2, data=x))
Call:
lm(formula = v \sim x1 + x2, data = x)
Residuals:
   Min 10 Median 30
                                  Max
-6.9303 -1.5736 -0.0068 1.3840 5.9567
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.00733 0.20900 0.035 0.97204
x1
            2.89168 0.09806 29.490 < 2e-16 ***
x2
            0.27903 0.09249 3.017 0.00268 **
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 2.141 on 503 degrees of freedom
Multiple R-squared: 0.6343, Adjusted R-squared: 0.6328
F-statistic: 436.2 on 2 and 503 DF, p-value: < 2.2e-16
```

Even worse...

```
> summary (lm(v \sim x1 + x2 + x4, data=x))
Call:
lm(formula = v \sim x1 + x2 + x4, data = x)
Residuals:
   Min 10 Median 30
                                 Max
-6.8064 -1.5229 -0.0308 1.4226 5.8861
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.0001127 0.2093588 0.001 1.000
       2.8964461 0.0983390 29.454 <2e-16 ***
×1
        0.9740807 0.9917783 0.982 0.326
x2
×4
          -0.6934442 0.9851714 -0.704 0.482
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 2.142 on 502 degrees of freedom
Multiple R-squared: 0.6347, Adjusted R-squared: 0.6325
F-statistic: 290.7 on 3 and 502 DF, p-value: < 2.2e-16
```

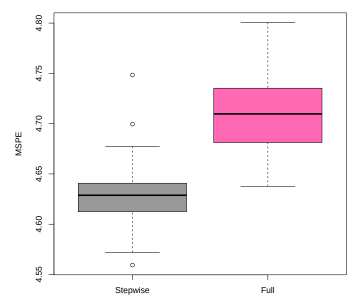
If we use AIC

```
> st <- stepAIC(null,
     scope=list(lower=null, upper=full))
> st
Call:
lm(formula = y \sim x1 + x6, data = x)
Coefficients:
(Intercept)
                       x1
                                    x6
  -0.000706
              3.175239
                            -0.282906
```

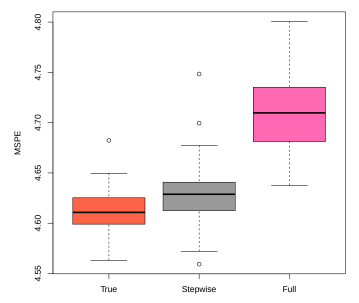
Discussion points

- Modeling problem (important variables may be missed)
- Prediction? Does stepwise give "the best" predicting model?

MSPEs



MSPEs



Discussion points

- Correlated covariates have become prevalent
- Researchers can (and do) collect data "blindly"
- Data are collected without a specific question in mind

Discussion points

- Correlated covariates:
- Mask each other when included simultaneously in a model
- May reduce prediction accuracy

One strategy:

- (1): Select models to be considered
- (2): Select a quantitative criterion to compare them (e.g. AIC, C_p , CV-based $\widehat{\mathsf{MSPE}}$)
- (3): Choose a strategy to explore the models under consideration

For example:

- (1): Consider all possible models
- (2): Use AIC to compare them
- (3): Best subset search (2^p fits!)
- (3'): Stepwise search

• Is this strategy prediction-based?

AIC?

Why not compare models using residual sum of squares, or R^2 ?

LS vs MLE

Note that, if we assume that the error distribution is Gaussian, then a least squares fit for a linear regression model is the same as the MLE fit

... or is it?

- Comparing likelihoods / residuals isn't very useful
- More complex models have higher likelihoods (smaller residuals)
- The Akaike Information Criterion provides a way to compare models with different number of parameters
- There are many different ways to motivate it

• We can measure the "distance" between the true distribution of the data $(f_0(y))$ and our model $f(y, \theta)$

$$d(\theta, f_0) = E_0 \left[-2 \ell(y, \theta) \right] =$$

$$\int -2 \ell(y, \theta) f_0(y) dy =$$

$$2 \left[\mathcal{K}(\theta, f_0) - \int \log(f_0(y)) f_0(y) dy \right]$$

• Given our estimator $\hat{\theta}_n$ we could use

$$d(\hat{\theta}_n, f_0) = E_0 \left[-2 \, \ell(y, \theta) \right]_{\theta = \hat{\theta}_n}$$

to see "how far" our model-based estimator is from the true distribution

- However, we can't compute $d(\hat{\theta}_n, f_0)$
- Can we use $-2 \ell(y, \hat{\theta}_n)$ to estimate $d(\hat{\theta}_n, f_0)$?

· Yes, but this estimator is biased

$$E_0\left[-2\,\ell(\mathbf{y},\hat{\theta}_n)\right] = \\ E_0\left\{E_0\left[-2\,\ell(\mathbf{y},\theta)\right]_{\theta=\hat{\theta}_n}\right\} - 2\,p + o(1)$$

In other words

$$E_0$$
 [AIC] $\approx E_0 \left[d(\hat{\theta}_n, f_0) \right]$

where

AIC =
$$-2\ell(y, \hat{\theta}_n) + 2p$$

For Gaussian errors we have

$$AIC = n \log \left(\frac{RSS}{n}\right) + 2p + constant$$

where

$$RSS = \sum_{i=1}^{n} r_i^2,$$

the **constant** depends on *n*, not on *p*

However, many times we find

AIC =
$$\frac{1}{n} \frac{1}{\hat{\sigma}^2} \left(RSS + 2 p \hat{\sigma}^2 \right) + constant$$

(e.g. [JWHT13])

Where does this expression come from?

Regularity assumptions are needed

- This is an asymptotic approximation, n should be large
- One of the models should include truth
- $\theta_1 \neq \theta_2 \Rightarrow f(y, \theta_1) \neq f(y, \theta_2)$
- Standard large-sample MLE assumptions to obtain asymptotic normality

- Air pollution example in R
- Synthetic data example

- AIC suggests a submodel
- Prediction-wise the full model is better
- AIC can be highly variable

"Smoother" model selection

- Ridge regression
- Can be thought as a type of feature selection
- It is a member of a larger class called "shrinkage methods"
- However, its origins are rather different

Without loss of generality...

• If covariates are centered, $\sum_{i=1}^{n} \mathbf{x}_{i} = \mathbf{0}$

$$\arg\min_{\alpha,\beta} \sum_{i=1}^{n} (y_i - \alpha - \beta' \mathbf{x}_i)^2$$

satisfies

$$\hat{\alpha}_n = \frac{1}{n} \sum_{i=1}^n y_i = \bar{y}_n,$$

and

$$\hat{\boldsymbol{\beta}}_{IS} = (\mathbf{X}'\mathbf{X})^{-1} \mathbf{X}'\mathbf{Y},$$

Without loss of generality...

• We can always assume that

$$\sum_{i=1}^{n} \mathbf{x}_{i} = \mathbf{0}$$

and hence

$$\hat{\alpha}_n = \frac{1}{n} \sum_{i=1}^n y_i = \bar{y}_n,$$

• In what follows, there is no intercept

Shrinkage methods

When covariates are correlated, LS estimators can be highly variable

$$\hat{eta}_{LS} = \left(\mathbf{X}'\mathbf{X}\right)^{-1}\,\mathbf{X}'\,\mathbf{Y}$$

$$\operatorname{var}\left(\hat{\boldsymbol{\beta}}_{n}\right) = \sigma^{2} \left(\mathbf{X}'\mathbf{X}\right)^{-1}$$

When X'X is close to singular...

Ridge Regression

 One way to "avoid" this problem is to add a "ridge" to X'X...

$$\hat{\boldsymbol{\beta}}_{RR} = \left(\mathbf{X}'\mathbf{X} + \lambda \, \mathbf{I}_{p}\right)^{-1} \, \mathbf{X}' \, \mathbf{Y}$$

where $\lambda > 0$ and

$$oldsymbol{I}_{
ho} = egin{pmatrix} 1 & 0 & \cdots & 0 \ 0 & 1 & \cdots & 0 \ 0 & \cdots & \ddots & 0 \ 0 & \cdots & \cdots & 1 \end{pmatrix}$$

Ridge Regression

• This is equivalent to solving

$$\min_{\boldsymbol{\beta}} \sum_{i=1}^{n} (y_i - \boldsymbol{\beta}' \mathbf{x}_i)^2 + \lambda \|\boldsymbol{\beta}\|_2^2$$

Ridge Regression

And also equivalent to solving

$$\min_{\beta} \sum_{i=1}^{n} (y_i - \beta' \mathbf{x}_i)^2$$

subject to

$$\sum_{j=1}^p \beta_j^2 \leq C$$

for some C > 0

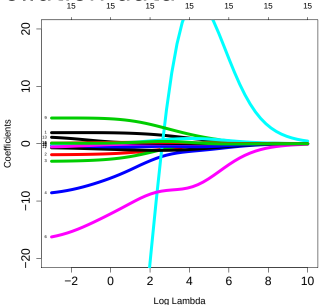
Bias / variance trade-off

- Ridge regression was originally proposed as a "hack" to "push" X'X away from singularity
- It can also be thought as a way of reducing the variance of $\hat{\beta}_n$
- This may increase the bias of the estimator, but if the variance is reduced even more, we might gain overall in expected squared error performance...

Ridge regression

- We now have a sequence ("path") of estimators (one for each $\lambda > 0$)
- $\mathbf{X}'\mathbf{X} + \lambda \mathbf{I}_p$ is always non-singular for $\lambda > 0$ (why?)
- Why are they called "shrinkage methods"?

Air pollution data



Questions

- What does λ measure?
- How do I choose one among these infinitely many "solutions"?

- How many "effective" parameters are we using?
- In linear regression, we have p parameters
- A more general definition is as follows. For a fitting method producing \hat{y}_1 , \hat{y}_2 , ..., \hat{y}_n ,

$$edf = \frac{1}{\sigma^2} \sum_{i=1}^{n} cov(\hat{y}_i, y_i)$$

 It is easy to see that for least squares predictors, we have

$$\hat{y} = Hy$$

with

$$\mathbf{H} = \mathbf{X} \left(\mathbf{X}' \mathbf{X} \right)^{-1} \mathbf{X}'$$

and

$$edf = \frac{1}{\sigma^2} \sum_{i=1}^{n} cov(\hat{y}_i, y_i) = trace(\mathbf{H}) = p$$

 More in general, for any linear predictor

$$\hat{\mathbf{y}} = \mathbf{S} \, \mathbf{y}$$

we have

$$edf = trace(\mathbf{S}) = \sum_{i=1}^{n} \mathbf{S}_{i,i}$$

The ridge regression fit satisfies

$$\hat{\mathbf{y}} = \mathbf{S}_{\lambda} \mathbf{y}$$

where

$$\mathbf{S}_{\lambda} = \mathbf{X} \left(\mathbf{X}' \mathbf{X} + \lambda \mathbf{I}_{p} \right)^{-1} \mathbf{X}'$$

$$trace(S) = ?$$

 Using the singular value decomposition (SVD) of X

$$X = U \Lambda V'$$

where $\mathbf{U} \in \mathbb{R}^{n \times p}$, $\mathbf{V} \in \mathbb{R}^{p \times p}$ with

$$\mathbf{U}'\mathbf{U} = \mathbf{I}_{p} = \mathbf{V}'\mathbf{V}$$

and

$$\Lambda = \operatorname{diag}(d_1,\ldots,d_p)$$
,

we have

trace (**S**) =
$$\sum_{i=1}^{p} \left(\frac{d_i^2}{d_i^2 + \lambda} \right)$$

 For example, in the Air Pollution data example, if we use

$$\lambda = \exp(6)$$

we get

$$edf = 9.9$$

How do we select λ ?

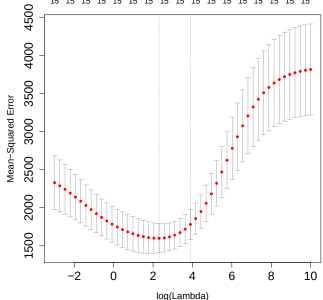
How can we select λ ?

How do we select λ ?

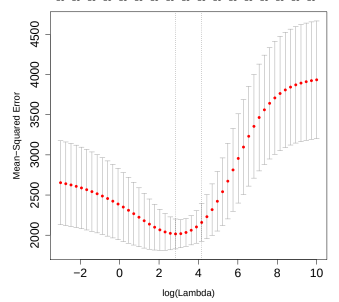
CV!

```
library(glmnet)
airp <- read.table('rutgers-lib-30861_CSV-1.csv'</pre>
      header=TRUE, sep=',')
v <- as.vector(airp$MORT)</pre>
xm < -as.matrix(airp[, -16])
lambdas \leftarrow exp( seq(-3, 10, length=50))
set.seed(123)
tmp <- cv.qlmnet(x=xm, y=y, lambda=lambdas,
      nfolds=5, alpha=0,
      family='gaussian')
```

Air pollution data



Air pollution data - another run



Questions

- How are the standard errors estimated?
- Can we use AIC to compare these models?
- Why or why not?
 - If the answer is yes, how?
 - If the answer is no, why not?

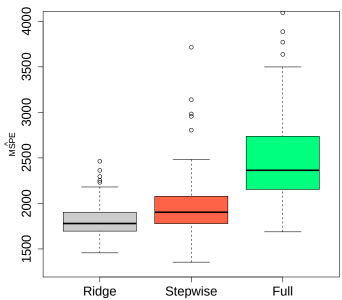
CV

Cross validation selects

$$\lambda_{\sf op} \approx \exp(3)$$
 edf ≈ 13

Stepwise selects

Air pollution - 100 5-fold CV runs



Sometimes...

- Selecting variables is not always necessary in terms of prediction accuracy.
- Doing so may in fact yield worse results.
- One such an example is discussed on Github.
- Read it carefully.