# Lecture 5

### DJM

#### 23 October 2018

### Model selection and tuning parameters

- Often "model selection" means "choosing a set of predictors" E.g. Lasso performs model selection by setting many  $\widehat{\beta} = 0$
- I define "model selection" more broadly
- I mean "making any necessary decisions to arrive at a final model"
- Sometimes this means "choosing predictors"
- It could also mean "selecting a tuning parameter"
- Or "deciding whether to use LASSO or Ridge" (and picking tuning parameters)
- Recall Lecture 2: "A statistical model  $\mathcal{P}$  is a collection of probability
- Model selection means "choose  $\mathcal{P}$ " distributions or densities."

### My pet peeve

- Often people talk about "using LASSO" or "using an SVM"
- This isn't quite right.
- LASSO is a regularized procedure that depends on  $\lambda$
- To "use LASSO", you must pick a particular  $\lambda$
- Different ways to pick  $\lambda$  (today's topic) produce different final estimators
- Thus we should say "I used LASSO+CV" or "I used Ridge+GCV"
- Probably also indicate "how" (I used the CV minimum.)

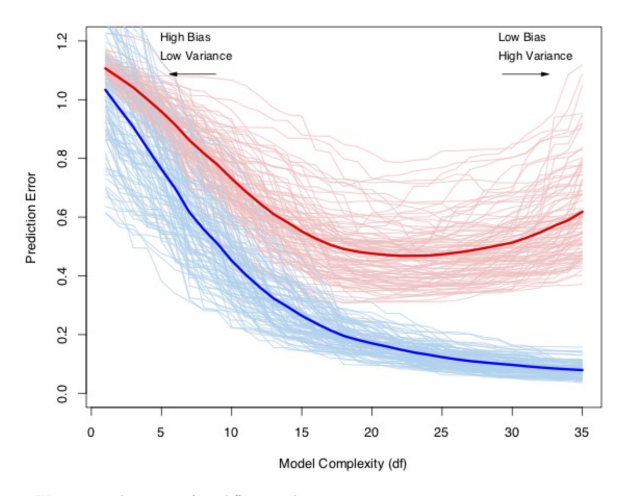
#### Bias and variance

Recall that  $\mathcal{D}$  is the training data.

$$R_n(f) := \mathbb{E}\left[L(Y, f(X))\right] = \mathbb{E}\left[\mathbb{E}\left[L(Y, f(X)) \mid \mathcal{D}\right]\right]$$

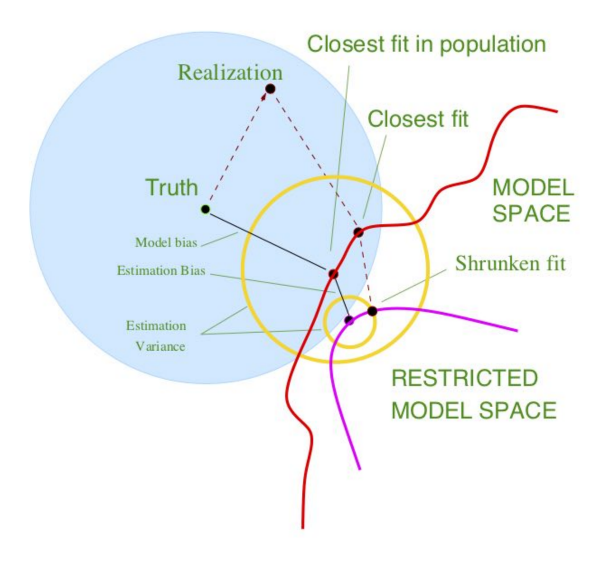
- The book calls  $R_n(f) = \text{Err and } \mathbb{E}[L(Y, f(X)) \mid \mathcal{D}] = \text{Err}_{\mathcal{D}}$
- If you use  $\mathcal{D}$  to choose f, then these are different.
- If you use  $\mathcal{D}$  to choose f, then both depend on how much data you have seen.

### Risk estimates



- We can use risk estimates for 2 different goals
- 1. Choosing between different potential models.
- 2. Characterizing the out-of-sample performance of the chosen model.
- I am not generally aware of other methods of accomplishing (1).
- You could avoid making a choice (Chapter 8), or you could use a procedure that makes the choice "automatically"
- The method you choose to estimate risk will have large implications for both 1 and 2.

### A model selection picture



### Why?

We want to do model selection for at least three reasons:

- Prediction accuracy: Can essentially always be improved by introducing some bias
- Interpretation: A large number of features can sometimes be distilled into a smaller number that comprise the "big (little?) picture"
- Computation: A large p can create a huge computational bottleneck.

### Things you shouldn't do

- Estimate  $R_n$  with  $\widehat{R}_n(f) = \sum_{i=1}^n L(Y_i, \widehat{f}(X_i))$ . Throw away variables with small p-values.
- Use F-tests
- Compare the log-likelihood between different models

(These last two can occasionally be ok, but aren't in general. You should investigate the assumptions that are implicit in them.)

### Risk estimators

#### Unbiased risk estimation

- It is very hard (impossible?) to estimate  $R_n$ .
- Instead we focus on

$$\overline{R}_n(f) = \mathbb{E}_{Y_1, \dots, Y_n} \left[ \mathbb{E}_{Y^0} \left[ \frac{1}{n} \sum_{i=1}^n L(Y_i^0, \widehat{f}(x_i)) \mid \mathcal{D} \right] \right].$$

- The difference is that  $\overline{R}_n(f)$  averages over the observed  $x_i$  rather than taking the expected value over the distribution of X.
- In the "fixed design" setting, these are equal.

For many L and some predictor  $\hat{f}$ , one can show

$$\overline{R}_n(\widehat{f}) = \mathbb{E}\left[\widehat{R}_n(\widehat{f})\right] + \frac{2}{n} \sum_{i=1}^n \operatorname{Cov}\left[Y_i, \ \widehat{f}(x_i)\right].$$

This suggests estimating  $\overline{R}_n(\widehat{f})$  with

$$\widehat{R}_{gic} := \widehat{R}_n(\widehat{f}) + \text{pen.}$$

If  $\mathbb{E}\left[\text{pen}\right] = \frac{2}{n} \sum_{i=1}^{n} \text{Cov}\left[Y_{i}, \ \widehat{f}(x_{i})\right]$ , we have an unbiased estimator of  $\overline{R}_{n}(\widehat{f})$ .

## Example: Normal means

#### Normal means model

Suppose we observe the following data:

$$Y_i = \beta_i + \epsilon_i, \qquad i = 1, \dots, n$$

where  $\epsilon_i \stackrel{iid}{\sim} N(0,1)$ .

We want to estimate

$$\boldsymbol{\beta} = (\beta_1, \dots, \beta_n).$$

The usual estimator (MLE) is

$$\widehat{\boldsymbol{\beta}}^{MLE} = (Y_1, \dots, Y_n).$$

This estimator has lots of nice properties: consistent, unbiased, UMVUE, (asymptotic) normality...

#### MLEs are bad

But, the standard estimator STINKS! It's a bad estimator.

It has no bias, but big variance.

$$R_n(\widehat{\boldsymbol{\beta}}^{MLE}) = \text{bias}^2 + \text{var} = 0 + n \cdot 1 = n$$

What if we use a biased estimator?

Consider the following estimator instead:

$$\widehat{\beta}_i^S = \begin{cases} Y_i & i \in S \\ 0 & \text{else.} \end{cases}$$

Here  $S \subseteq \{1, \ldots, n\}$ .

### Biased normal means

What is the risk of this estimator?

$$R_n(\widehat{\boldsymbol{\beta}}^S) = \sum_{i \notin S} \beta_i^2 + |S|.$$

In other words, if some  $|\beta_i| < 1$ , then don't bother estimating them! In general, introduced parameters like S will be called **tuning parameters**.

Of course we don't know which  $|\beta_i| < 1$ .

But we could try to estimate  $R_n(\widehat{\boldsymbol{\beta}}^S)$ , and choose S to minimize our estimate.

#### Estimating the risk

By definition, for any estimator  $\widehat{\beta}$ ,

$$R_n(\widehat{\boldsymbol{\beta}}) = \mathbb{E}\left[\sum_{i=1}^n (\widehat{\beta}_i - \beta_i)^2\right]$$

An intuitive estimator of  $R_n$  is

$$\widehat{R}_n(\widehat{\boldsymbol{\beta}}) = \sum_{i=1}^n (\widehat{\beta}_i - Y_i)^2.$$

This is known as the **training error** and it can be shown that

$$\widehat{R}_n(\widehat{\boldsymbol{\beta}}) \approx R_n(\widehat{\boldsymbol{\beta}}).$$

Also,

$$\widehat{\boldsymbol{\beta}}^{MLE} = \arg\min_{\boldsymbol{\beta}} \widehat{R}_n(\widehat{\boldsymbol{\beta}}^{MLE}).$$

What could possibly go wrong?

### Dangers of using the training error

Although

$$\widehat{R}_n(\widehat{\boldsymbol{\beta}}) \approx R_n(\widehat{\boldsymbol{\beta}}),$$

this approximation can be very bad. In fact:

Training Error:  $\widehat{R}_n(\widehat{\boldsymbol{\beta}}^{MLE}) = 0$ 

Risk:  $R_n(\widehat{\boldsymbol{\beta}}^{MLE}) = n$ 

In this case, the **optimism** of the training error is n.

### Normal means

What about  $\widehat{\boldsymbol{\beta}}^{S}$ ?

$$\widehat{R}_n(\widehat{\boldsymbol{\beta}}^S) = \sum_{i=1}^n (\widehat{\beta}_i - Y_i)^2 = \sum_{i \notin S} Y_i^2$$

Well

$$\mathbb{E}\left[\widehat{R}_n(\widehat{\boldsymbol{\beta}}^S)\right] = R_n(\widehat{\boldsymbol{\beta}}^S) - 2|S| + n.$$

So I can choose S by minimizing  $\widehat{R}_n(\widehat{\boldsymbol{\beta}}^S) + 2|S|$ .

Estimate of Risk = training error + penalty.

The penalty term corrects for the optimism.

#### pen in the nice cases

#### Result:

Suppose  $\widehat{f}(x_i) = HY$  for some matrix H, and  $Y_i$ 's are IID. Then

$$\frac{2}{n}\sum_{i=1}^{n}\operatorname{Cov}\left[Y_{i},\ \widehat{f}(x_{i})\right] = \frac{2}{n}\sum_{i=1}^{n}H_{ii}\operatorname{Cov}\left[Y_{i},\ Y_{i}\right] = \frac{2\mathbb{V}\left[Y\right]}{n}\operatorname{tr}(H).$$

- Such estimators are called "linear smoothers".
- Obvious extension to the heteroskedastic case.
- We call  $\frac{1}{\mathbb{V}[Y]} \sum_{i=1}^{n} \text{Cov} \left[ Y_i, \ \widehat{f}(x_i) \right]$  the **degrees of freedom** of  $\widehat{f}$ .
- Linear smoothers are ubiquitous
- Examples: OLS, ridge regression, KNN, dictionary regression, smoothing splines, kernel regression, etc.

### Examples of DF

• OLS

$$H = X^\top (X^\top X)^{-1} X^\top \Rightarrow \operatorname{tr}(H) = \operatorname{rank}(X) = p$$

• Ridge (decompose  $X = UDV^{\top}$ )

$$H = X^{\top} (X^{\top} X + \lambda I_p)^{-1} X^{\top} \Rightarrow \operatorname{tr}(H) = \sum_{i=1}^{p} \frac{d_i^2}{d_i^2 + \lambda} < \min\{p, n\}$$

• KNN df = n/K (each point is it's own nearest neighbor, it gets weight 1/K)

### Finding risk estimators

This isn't the way everyone introduces/conceptualizes prediction risk.

For me, thinking of  $\widehat{R}_n$  as overly optimistic and correcting for that optimism is conceptually appealing We need to also discuss **information criteria**.

In this case one forms a (pseudo)-metric on probability measures.

## Comparing probability measures

#### Kullback-Leibler

Suppose we have data Y that comes from the probability density function f.

What happens if we use the probability density function g instead?

#### Example:

Suppose  $Y \sim N(\mu, \sigma^2) =: f$ . We want to predict a new  $Y_*$ , but we model it as  $Y_* \sim N(\mu_*, \sigma^2) =: g$ 

How far away are we? We can either compare  $\mu$  to  $\mu_*$  or Y to  $Y^*$ .

Or, we can compute how far f is from g.

We need a notion of distance.

#### Kullback-Leibler

One central idea is Kullback-Leibler divergence (or discrepancy)

$$KL(f,g) = \int \log \left(\frac{f(y)}{g(y)}\right) f(y) dy$$

$$\propto -\int \log(g(y)) f(y) dy \qquad \text{(ignore term without } g)$$

$$= -\mathbb{E}_f[\log(g(Y))]$$

This gives us a sense of the **loss** incurred by using g instead of f.

• KL is not symmetric:  $KL(f,g) \neq KL(g,f)$ , so it's not a distance, but it is non-negative and satisfies the triangle inequality.

Usually, g will depend on some parameters, call them  $\theta$ 

#### KL example

- In regression, we can specify  $f = N(X^{\top}\beta_*, \sigma^2)$
- for a fixed (true)  $\beta_*$ ,
- let  $g_{\theta} = N(X^{\top}\beta, \sigma^2)$  over all  $\theta = (\beta, \sigma^2) \in \mathbb{R}^p \times \mathbb{R}^+$
- $KL(f, g_{\theta}) = -\mathbb{E}_f[\log(g_{\theta}(Y))]$ , we want to minimize this over  $\theta$ .
- But f is unknown, so we minimize  $-\log(g_{\theta}(Y))$  instead.

• This is the maximum likelihood value

$$\widehat{\theta}_{ML} = \arg\max_{\theta} g_{\theta}(Y)$$

We don't actually need to assume things about a true model nor have it be nested in the alternative
models to make this work.

### Operationalizing

• Now, to get an operational characterization of the KL divergence at the ML solution

$$-\mathbb{E}_f[\log(g_{\widehat{\theta}_{ML}}(Y))]$$

we need an approximation (don't know f, still).

#### Result:

If you maximize the likelihood for a finite dimensional parameter vector  $\theta$  of length p, then as  $n \to \infty$ ,

$$-\mathbb{E}_f[\log(g_\theta(Y))] = -\log(g_\theta(Y)) + p.$$

- This is AIC (originally "an information criterion", now "Akaike's information criterion").
- Choose the model with smallest AIC
- Often multiplied by 2 "for historical reasons". Ocassionally, given as the negative of this "to be extra annoying".
- Your estimator for  $\theta$  needs to be the MLE. p includes all estimated parameters.

### Back to the OLS example

• Suppose Y comes from the standard normal linear regression model with known variance  $\sigma^2$ .

$$-\log(g_{\widehat{\theta}}) \propto \log(\sigma^2) + \frac{1}{2\sigma^2} \sum_{i=1}^n (y_i - x_i^{\top} \widehat{\beta}_{MLE})^2$$
$$\Rightarrow AIC = 2 \frac{n}{2\sigma^2} \widehat{R}_n + 2p = \widehat{R}_n + \frac{2\sigma^2}{n} p.$$

• Suppose Y comes from the standard normal linear regression model with unknown variance  $\sigma^2$ . Note that  $\widehat{\sigma}_{MLE}^2 = \frac{1}{n} \sum_{i=1}^n (y_i - x_i^{\mathsf{T}} \widehat{\beta}_{MLE})^2$ .

$$-\log(g_{\widehat{\theta}}) \propto \frac{n}{2} \log(\widehat{\sigma}^2) + \frac{1}{2\widehat{\sigma}^2} \sum_{i=1}^n (y_i - x_i^{\top} \widehat{\beta}_{MLE})^2$$
$$\Rightarrow AIC \propto 2n \log(\widehat{\sigma}^2)/2 + 2(p+1) \propto \log(\widehat{R}_n) + \frac{2(p+1)}{n}.$$

## Related quantities

#### Mallow's Cp

- Defined for linear regression.
- No likelihood assumptions.
- Variance is known

$$C_p = \widehat{R}_n + 2\sigma^2 \frac{\mathrm{df}}{n} = AIC$$

### Bayes factor

For Bayesian Analysis, we want the posterior. Suppose we have two models A and B.

$$P(B \mid \mathcal{D}) = \frac{P(\mathcal{D} \mid B)P(B)}{P(\mathcal{D})} \propto P(\mathcal{D} \mid B)P(B)$$

$$P(A \mid \mathcal{D}) = \frac{P(\mathcal{D} \mid A)P(A)}{P(\mathcal{D})} \propto P(\mathcal{D} \mid A)P(A)$$

We assume that P(A) = P(B). Then to compare,

$$\frac{P(B \mid \mathcal{D})}{P(A \mid \mathcal{D})} = \frac{P(\mathcal{D} \mid B)}{P(\mathcal{D} \mid A)}.$$

- Called the Bayes Factor.
- This is the ratio of marginal likelihoods under the different models.
- Not easy to calculate generally.
- Use the Laplace approximation, some simplifications, assumptions:

$$\log P(\mathcal{D} \mid B) = \log P(\mathcal{D} \mid \widehat{\theta}, B) - \frac{p \log(n)}{2} + O(1)$$

• Multiply through by -2:

$$BIC = -\log(g_{\theta}(Y)) + p\log(n) = \log(\widehat{R}_n) + \frac{p\log(n)}{n}$$

• Also called Schwarz IC. Compare to AIC (variance unknown case)

#### SURE

$$\widehat{R}_{gic} := \widehat{R}_n(\widehat{f}) + \text{pen.}$$

If  $\mathbb{E}[\text{pen}] = \frac{2}{n} \sum_{i=1}^{n} \text{Cov}[Y_i, \ \widehat{f}(x_i)]$ , we have an unbiased estimator of  $\overline{R}_n(\widehat{f})$ .

#### Result: (Stein's Lemma)

Suppose  $Y_i \sim N(\mu_i, \sigma^2)$  and suppose f is weakly differentiable. Then

$$\frac{1}{\sigma^2} \sum_{i=1}^n \operatorname{Cov} \left[ Y_i, \ \widehat{f}_i(Y) \right] = \mathbb{E} \left[ \sum_{i=1}^n \frac{\partial f_i}{\partial y_i} \widehat{f}(Y) \right].$$

- Note: Here I'm writing  $\widehat{f}$  as a function of Y rather than x.
- This gives "Stein's Unbiased Risk Estimator"

$$SURE = \widehat{R}_n(\widehat{f}) + 2\sigma^2 \sum_{i=1}^n \frac{\partial f_i}{\partial y_i} \widehat{f}(Y) - n\sigma^2.$$

- If f(Y) = HY is linear, we're back to AIC (variance known case)
- If  $\sigma^2$  is unknown, may not be unbiased anymore. May not care.

### CV

#### What is Cross Validation

- Cross validation
- This is another way or estimating the prediction risk.
- Why?

To recap:

$$R_n(\widehat{f}) = \mathbb{E}[\ell(Y, \widehat{f}(X))]$$

where the expectation is taken over the new data point (Y, X) and  $\mathcal{D}_n$  (everything that is random).

We saw one estimator of  $R_n$ :

$$\widehat{R}_n(\widehat{f}) = \sum_{i=1}^n \ell(Y_i, \widehat{f}(X_i)).$$

This is the training error. It is a **BAD** estimator because it is often optimistic.

#### Intuition for CV

- One reason that  $\widehat{R}_n(\widehat{f})$  is bad is that we are using the same data to pick  $\widehat{f}$  **AND** to estimate  $R_n$ .
- Notice that  $R_n$  is an expected value over a **NEW** observation (Y, X).
- We don't have new data.

#### Wait a minute...

 $\dots$  or do we?

- What if we set aside one observation, say the first one  $(Y_1, X_1)$ .
- We estimate  $\hat{f}^{(1)}$  without using the first observation.
- Then we test our prediction:

$$\widetilde{R}_1(\widehat{f}^{(1)}) = \ell(Y_1, \widehat{f}^{(1)}(X_1)).$$

- But that was only one data point  $(Y_1, X_1)$ . Why stop there?
- Do the same with  $(Y_2, X_2)!$  Get an estimate  $\widehat{f}^{(2)}$  without using it, then

$$\widetilde{R}_2(\widehat{f}^{(2)}) = \ell(Y_2, \widehat{f}^{(2)}(X_2)).$$

### Keep going

- We can keep doing this until we try it for every data point.
- And then average them! (Averages are good)
- In the end we get

LOO-CV = 
$$\sum_{i=1}^{n} \widetilde{R}_{i}(\widehat{f}^{(i)}) = \sum_{i=1}^{n} \ell(Y_{i} - \widehat{f}^{(i)}(X_{i}))$$

• This is leave-one-out cross validation

### Problems with LOO-CV

- 1. Each held out set is small (n = 1). Therefore, the variance of my predictions is high.
- 2. Since each held out set is small, the training sets overlap. This is bad.
  - Usually, averaging reduces variance:

$$\mathbb{V}\left[\overline{X}\right] = \frac{1}{n^2} \sum_{i=1}^{n} \mathbb{V}\left[X_i\right] = \frac{1}{n} \mathbb{V}\left[X_1\right].$$

• But only if the variables are independent. If not, then

$$\mathbb{V}\left[\overline{X}\right] = \frac{1}{n^2} \mathbb{V}\left[\sum_{i=1}^n X_i\right]$$
$$= \frac{1}{n} \mathbb{V}\left[X_1\right] + \frac{1}{n^2} \sum_{i \neq j} \operatorname{Cov}\left[X_i, X_j\right].$$

- Since the training sets overlap a lot, that covariance can be pretty big.
- 3. We have to estimate this model n times.
  - There is an exception to this one. More on that in a minute.

#### Generic Cross Validation

Let  $\mathcal{N} = \{1, \dots, n\}$  be the index set for  $\mathcal{D}$ 

Define a distribution  $\mathcal{V}$  over  $\mathcal{N}$   $(v \sim \mathcal{V} \subseteq \mathcal{N})$ 

Then, we can form a general *cross-validation* estimator as

$$\mathrm{CV}_{\mathcal{V}}(\widehat{f}) = \mathbb{E}\left[\frac{1}{|v|} \sum_{i \in v} L\left(Y_i, \widehat{f}^{(v)}(X_i)\right) \mid \mathcal{V}\right]$$

More general cross-validation schemes: Examples

$$\mathrm{CV}_{\mathcal{V}}(\widehat{f}) = \mathbb{E}\left[\frac{1}{|v|} \sum_{i \in v} L\left(Y_i, \widehat{f}^{(v)}(X_i)\right) \mid \mathcal{V}\right]$$

• K-fold:

Fix  $V = \{v_1, \dots, v_K\}$  such that  $v_j \cap v_k = \emptyset$  and  $\bigcup_j v_j = \mathcal{N}$ 

$$CV_K(\hat{f}) = \frac{1}{K} \sum_{v \in V} \frac{1}{|v|} \sum_{i \in v} (Y_i - \hat{f}^{(v)}(X_i))^2$$

• Bootstrap:

Let  $\mathcal{V}$  be given by the bootstrap distribution over  $\mathcal{N}$  (that is, sampling B indices randomly with replacement many times)

• Factorial

Let V be given by all subsets (or a subset of all subsets) of  $\mathcal{N}$  (that is, putting mass  $1/(2^n-2)$  on each subset)

### More general cross-validation schemes: A comparison

- $CV_K$  gets more computationally demanding as  $K \to n$
- The bias of  $CV_K$  goes down, but the variance increases as  $K \to n$
- The factorial version isn't commonly used except when doing a 'real' data example for a methods paper
- There are many other flavors of CV. One of them, called "consistent cross validation" is a recent addition that is designed to work with sparsifying algorithms
- K-fold is most common (like K = 10 or K = 5)

#### K-fold CV

- 1. Divide the data into K groups.
- 2. Leave a group out and estimate with the rest.
- 3. Test on the held-out group. Calculate an average risk over these  $\sim n/K$  data.
- 4. Repeat for all K groups.
- 5. Average the average risks.

### Why K-fold better?

- 1. Less overlap, smaller covariance.
- 2. Larger hold-out sets, smaller variance.
- 3. Less computations (only need to estimate K times)

### Why might it be worse?

- 1. LOO-CV is (nearly) unbiased.
- 2. The risk depends on how much data you use to estimate the model.
- 3. LOO-CV uses almost the same amount of data.

### A picture

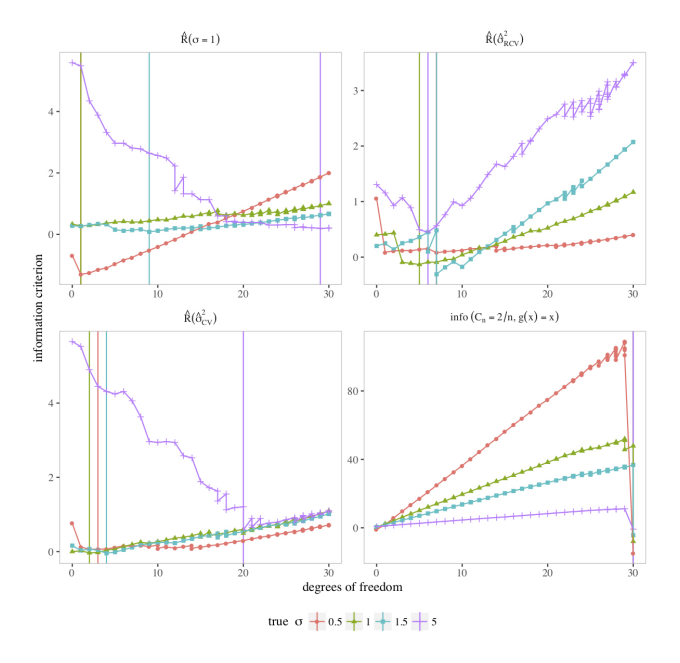


### Comparison

- LOO-CV and AIC are asymptotically equivalent  $(n \to \infty, p < n)$ , (Stone 1977)
- Properties of AIC/BIC in high dimensions are not well understood.
- In low dimensions, AIC is minimax optimal for the prediction risk (Yang and Barron 1998)
- CV is consistent for the prediction risk (Dudoit and Laan 2005)
- Both tend to over-select predictors (unproven, except empirically)
- BIC asymptotically selects the correct linear model in low dimensions (Shao 1997) and in high dimensions (Wang, Li, and Leng 2009)
- In linear regression, it is impossible for a model selection criterion to be minimax optimal and select the correct model asymptotically (Yang 2005)
- In high dimensions, if the variance is unknown, the "known" variance form of AIC/BIC is disastrous.
- Conclusion: your choice of risk estimator impacts results. Thus,
  - 1. If you want to select models, you might pick BIC
  - 2. If you want good predictions, you might use CV
  - 3. It's possible LASSO+CV(1se) picks models better than LASSO+CV(min)

### AIC/BIC disaster

```
n = 30; p = 150
sigma = c(.5, 1, 1.5, 5)
beta = c(1, 0, ..., 0)
Y = X %*% beta + sigma * rnorm(n)
```



# (Brief) foray into model averaging

### What if we don't want to choose?

- 1. Choose a risk estimator  $\widehat{R}$
- 2. Calculate weights  $p_i = \exp{-\widehat{R}(\text{Model}_i)}$ 3. Create final estimator  $\widehat{f} = \sum_{\text{models}} \frac{p_i}{\sum p_i} \widehat{f}_i$ .
- If  $\widehat{R}$  is BIC, this is (poor-man's) Bayesian Model Averaging. Real BMA integrates over the models:

$$P(f \mid \mathcal{D}) = \int P(f \mid M_i, \mathcal{D}) P(M_i \mid \mathcal{D}) dM$$

- Averaging + Sparsity is pretty hard.
- Interesting open problem: how can we combine LASSO models over the path?
- Issue with MA:  $e^{-BIC}$  can be tiny for all but a few models. You're not averaging anymore.

#### Selected references

Dudoit, Sandrine, and Mark J. van der Laan. 2005. "Asymptotics of Cross-Validation Risk Estimation in Estimator Selection and Performance Assessment." *Statistical Methodology*, 131–54.

Shao, Jun. 1997. "An Asymptotic Theory for Linear Model Selection." Statistica Sinica 7 (2): 221-42.

Stone, M. 1977. "Asymptotics for and Against Cross-Validation." Biometrika 64 (1): 29–35.

Wang, Hansheng, Bo Li, and Chenlei Leng. 2009. "Shrinkage Tuning Parameter Selection with a Diverging Number of Parameters." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)* 71 (3): 671–83.

Yang, Yuhong. 2005. "Can the Strengths of Aic and Bic Be Shared? A Conflict Between Model Indentification and Regression Estimation." *Biometrika* 92 (4): 937–50.

Yang, Yuhong, and Andrew Barron. 1998. "An Asymptotic Property of Model Selection Criteria." *IEEE Transactions on Information Theory* 44: 95–116.