Linear-Nonlinear Regression

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April 24, 2023

Linear-in-Parameters Models

We've often brushed over models which are non-linear in parameters. Why?

- Problems linear in parameters generally much easier to estimate; criterion functions often quadratic; normal equations linear.
- Perhaps more important: linear models can nevertheless be very effective for estimating non-linear phenomena.

Estimating non-linear phenomena with linear models

Our basic linear regression model is:

$$y = X\beta + u.$$

But the linearity that's important for estimation here is the linearity in parameters. We can just as well have

$$y = f(X) + u$$
, with $f(X) = \hat{f}(X; \alpha) + \epsilon$;

where

$$\hat{f}(X;\alpha) = \sum_{k=1}^{K} \alpha_k \phi_k(X);$$

The (ϕ_k) are basis functions with which we can try to approximate f. Note linearity in parameters $\alpha!$

Stepwise Basis Functions

For a function f defined over an interval (0,1) define:

K	$\phi_1(x)$	$\phi_2(x)$	$\phi_3(x)$	$\phi_4(x)$
2	$\mathbb{1}\{x \le \frac{1}{2}\}$	$1\{x > \frac{1}{2}\}$		
3	$1\{x \leq \frac{1}{3}\}$	$1\left\{\frac{1}{3} > x \leq \frac{2}{3}\right\}$	$1\{x>\frac{2}{3}\}$	
4	$1\{x \le \frac{1}{4}\}$	$1\left\{\frac{1}{4} > x \le \frac{1}{2}\right\}$	$ 1{x > \frac{2}{3}} 1{\frac{1}{2} < x \le \frac{3}{4}} $	$\mathbb{1}\{x > \frac{3}{4}\}$
:	:	:	:	:

Radial Basis Functions

If $x \in \mathbb{R}^n$, define a set of centers $c_k \in \mathbb{R}^n$, and let

$$\phi_k(x) = K(x, c_k) = e^{-\frac{1}{2}||x - c_k||^2}.$$

Gram Matrix

This may seem as though we then need to choose a bunch of non-linear parameters, but consider letting $c_k=x_k$, where x_k is the kth observation in a dataset; then we have:

$$\boldsymbol{K} = \begin{bmatrix} K(x_1, x_1) & K(x_1, x_2) & \cdots & K(x_1, x_N) \\ K(x_2, x_1) & K(x_2, x_2) & \cdots & K(x_2, x_N) \\ \vdots & \vdots & \ddots & \vdots \\ K(x_N, x_1) & K(x_N, x_2) & \cdots & K(x_N, x_N) \end{bmatrix}$$

Other Basis Functions

- Polynomials
- Splines
- Piecewise linear functions
- Periodic functions...

"Over-fitting" & MSE

We can fit any given dataset better by increasing the number of functions in the basis. However, at some point improving the fit for a *given* sample makes the fit worse for a *different* sample.

Mean Squared Error

In a given sample, large deviations from true f are evidence of either a large bias or large variance.

In this case we can compute the squared bias of this particular estimated function \hat{f} by

$$MSE(\hat{f}) = \int \left(f(x) - \hat{f}(x) \right)^2 dF(x),$$

where F(x) is the CDF of X.

Leave-one-out estimators

Whatever estimator we have of $\hat{y}(X)$, we presume that can be estimated using observations $j=1,\ldots,N$.

Estimator

An old idea for improving out-of-sample predictive power is the "leave-one-out" estimator. In this case for each observation j we construct a prediction function using every observation $except\ j;$ i.e.,

$$\hat{y}_{-j}(X) = \hat{f}(X|y_{-j}, X_{-j})$$
 $j = 1, \dots, N.$

That is, \hat{y}_{-j} is estimated using data (y_{-j}, X_{-j}) , but can then be evaluated at any X. Then the usual leave-one-out estimator is

$$\hat{y}(X) = \frac{1}{N} \sum_{j=1}^{N} \hat{y}_{-j}(X).$$

Note that every single $\hat{y}_{-j}(X_j)$ is an out-of-sample prediction, since (y_i, X_j) weren't used to construct \hat{y}_{-j} .

Cross-validation criterion

Let $e_{-j}=y_j-\hat{y}_{-j}(X_j)$. Call this the "leave-one-out error".

Criterion

Define the cross-validation criterion as

$$CV = \frac{1}{N} \sum_{j=1}^{N} e_{-j}^{2}.$$

Given the iid sampling assumption it's easy to see that this is an unbiased estimator of the expected squared out-of-sample prediction error.

General Approach

If we have multiple possible models or estimators of the relationship between y and X, say $(\hat{y}^1(X),\dots,\hat{y}^q(X))$ we select a model by choosing the one that minimizes the CV criterion.

Parameter Estimation

Rather than selecting among a finite number of discrete "models", we can also use the CV criterion as the basis for selecting a vector of continuous parameters.

K-fold Cross-Validation

A practical problem is that as N grows large our proposal to choose models by minimizing CV become impractical; typically the computational cost of implementing the estimator becomes $O(N^2)$ or worse. Enter K-fold cross-validation.

- ① Divide sample of N observations into K different "folds" $d_{(k)}$, each of roughly size N/K.
- ② Estimate a "leave N/K out" estimator that uses data from K-1 folds to estimate $\hat{y}^{(-k)}(X)$.
- Let

$$e_j^{(k)} = y_j - \hat{y}^{(-k)}(X_j)$$
 for $j \in d_{(k)}$.

The criteria

$$\mathsf{CV}^{(k)} = \frac{1}{K} \sum_{k=1}^K \frac{1}{N_k} \sum_{j \in d_{(k)}} (e_j^{(k)})^2$$

then approximates the CV criterion. (If K=N the two are identical).

Expected MSE

We can think of the expected MSE as the limit we'd reach taking averages in repeated samples. (This can be estimated in a given finite sample by our Cross-Validation measure). If \hat{f}_m is estimated using a sample $m=1,\ldots,M$, then

$$\mathsf{CV}_M = \frac{1}{M} \sum_{m=1}^M \mathsf{MSE}(\hat{f}_m) \overset{p}{\longrightarrow} \mathsf{EMSE}.$$

Various Penalizations

A variety of approaches to trying to encourage models with fewer parameters: goal is to achieve balance between bias and variance, say by minimizing EMSE.

- ullet Adjusted R^2 : $1-(1-R^2)\frac{N-1}{N-k-1}$
- Akaike Information Criterion: $N(1 + \log 2\pi \hat{\sigma}^2) + 2k$
- ullet Bayesian Information Criterion: $N(1+\log 2\pi\hat{\sigma}^2)+k\log N$

Loss-Penalty Form

A really wide variety of estimators can be written in so called "loss-penalty" form, where we try to choose a vector of parameters \boldsymbol{b} to solve

$$\min_{b \in B} L(b) + \lambda ||b||.$$

The first term is something like (minus) a log-likelihood, or the GMM criterion, or some other loss function. The second term is a "penalty" function, which induces a bias toward making the parameters b small (perhaps zero). The parameter $\lambda>0$ is a "tuning" parameter; larger values "penalize" large b more, increasing bias so as to reduce variance.

Effective Degrees of Freedom

Consider a regression linear in ${\cal K}$ parameters; then the model can be represented as

$$y = Xa + e$$
.

Let $V = \frac{1}{N} X^{\top} X - \bar{X} \bar{X}^{\top}$ be the sample covariance matrix of X, and let d_k^2 be the eigenvalues of this matrix. Then the *effective degrees of freedom* for the regression in Loss-Penalty form is

$$df(\lambda) = \sum_{k=1}^{K} \frac{d_k^2}{d_k^2 + \lambda}.$$

The Lasso (Least Absolute Shrinkage and Selection Operator)

The Lasso takes the form

$$\min_{b \in B} \sum_{j=1}^{N} (y_j - X_j b)^2 + \lambda \sum_{k=1}^{K} |b_k|$$

The absolute value penalty $(L_1 \text{ norm})$ means that the method will try to set coefficients to zero where doing so doesn't compromise the fit too much. Thus, the larger λ the fewer non-zero coefficients we expect to see (think, the more parsimonious the regression specification).

Tuning

So, how should we choose λ ? Too small, and we increase bias; too big we increase variance. Note that in the Lasso case choosing *one* parameter can the the effect of introducing or eliminating *lots* of parameters.

Cross-Validation

The cross-validation tools we discussed last time have many uses, but one very simple and effective use case involves tuning just a single parameter to try and minimize MSE. Let

$$\mathsf{CV}(\lambda) = \frac{1}{N} \sum_{j=1}^{N} e_{-j}(\lambda)^{2};$$

Then choose

$$\lambda^* = \operatorname*{arg\,min}_{\lambda \in \mathbb{R}_+} \mathsf{CV}(\lambda).$$

