Chapter 2

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What is this chapter about?

Problems with regression, and in particular, linear regression

A quick overview:

- 1. The truth is almost never linear.
- 2. Collinearity can cause difficulties for numerics and interpretation.
- 3. The estimator depends strongly on the marginal distribution of X.
- 4. Leaving out important variables is bad.
- 5. Noisy measurements of variables can be bad, but it may not matter.

Asymptotic notation

• The Taylor series expansion of the mean function $\mu(x)$ at some point u

$$\mu(x) = \mu(u) + (x - u)^{\top} \frac{\partial \mu(x)}{\partial x}|_{x=u} + O(\|x - u\|^2)$$

- The notation f(x) = O(g(x)) means that for any x there exists a constant C such that f(x)/g(x) < C.
- More intuitively, this notation means that the remainder (all the higher order terms) are about the size of the distance between x and u or smaller.
- So as long as we are looking at points u near by x, a linear approximation to $\mu(x) = \mathbb{E}[Y \mid X = x]$ is reasonably accurate.

What is bias?

- We need to be more specific about what we mean when we say bias.
- Bias is neither good nor bad in and of itself.
- A very simple example: let $Z_1, \ldots, Z_n \sim N(\mu, 1)$.
- We don't know μ , so we try to use the data (the Z_i 's) to estimate it.
- I propose 3 estimators:
 - 1. $\hat{\mu}_1 = 12$,
 - 2. $\hat{\mu}_2 = Z_6,$ 3. $\hat{\mu}_3 = \overline{Z}.$
- The bias (by definition) of my estimator is $\mathbb{E}[\hat{\mu}] \mu$.
- Calculate the bias and variance of each estimator.

Regression in general

• If I want to predict Y from X, it is almost always the case that

$$\mu(x) = \mathbb{E}\left[Y \mid X = x\right] \neq x^{\top}\beta$$

- There are always those errors $O(||x-u||)^2$, so the **bias** is not zero.
- We can include as many predictors as we like, but this doesn't change the fact that the world is non-linear.

Covariance between the prediction error and the predictors

• In theory, we have (if we know things about the state of nature)

$$\beta^* = \arg\min_{\beta} \mathbb{E}\left[\|Y - X\beta\|^2 \right] = \operatorname{Cov}\left[X, \ X\right]^{-1} \operatorname{Cov}\left[X, \ Y\right]$$

- Define $v^{-1} = \text{Cov}[X, X]^{-1}$.
- Using this optimal value β^* , what is $Cov[Y X\beta^*, X]$?

$$\begin{aligned} \operatorname{Cov}\left[Y-X\beta^{*},\ X\right]&=\operatorname{Cov}\left[Y,\ X\right]-\operatorname{Cov}\left[X\beta^{*},\ X\right] & \text{(Cov is linear)} \\ &=\operatorname{Cov}\left[Y,\ X\right]-\operatorname{Cov}\left[X(v^{-1}\operatorname{Cov}\left[X,\ Y\right]),\ X\right] & \text{(substitute the def. of }\beta^{*}) \\ &=\operatorname{Cov}\left[Y,\ X\right]-\operatorname{Cov}\left[X,\ X\right]v^{-1}\operatorname{Cov}\left[X,\ Y\right] & \text{(Cov is linear in the first arg)} \\ &=\operatorname{Cov}\left[Y,\ X\right]-\operatorname{Cov}\left[X,\ Y\right]=0. \end{aligned}$$

Bias and Collinearity

- Adding or dropping variables may impact the bias of a model
- Suppose $\mu(x) = \beta_0 + \beta_1 x_1$. It is linear. What is our estimator of β_0 ?
- If we instead estimate the model $y_i = \beta_0$, our estimator of β_0 will be biased. How biased?
- But now suppose that $x_1 = 12$ always. Then we don't need to include x_1 in the model. Why not?
- Form the matrix [1 x_1]. Are the columns collinear? What does this actually mean?

When two variables are collinear, a few things happen.

- 1. We cannot **numerically** calculate $(\mathbf{X}^{\top}\mathbf{X})^{-1}$. It is rank deficient.
- 2. We cannot **intellectually** separate the contributions of the two variables.
- 3. We can (and should) drop one of them. This will not change the bias of our estimator, but it will alter our interpretations.
- 4. Collinearity appears most frequently with many categorical variables.
- 5. In these cases, software **automatically** drops one of the levels resulting in the baseline case being in the intercept. Alternately, we could drop the intercept!
- 6. High-dimensional problems (where we have more predictors than data points) also lead to rank deficiencies.
- 7. There are methods (regularizing) which attempt to handle this issue (both the numerics and the interpretability). We may have time to cover them slightly.

White noise

White noise is a stronger assumption than Gaussian.

Consider a random vector ϵ .

- 1. $\epsilon \sim N(0, \Sigma)$.
- 2. $\epsilon_i \sim N(0, \sigma^2(x_i))$. 3. $\epsilon \sim N(0, \sigma^2 I)$.

The third is white noise. The ϵ are normal, their variance is constant for all i and independent of x_i , and they are independent.

Asymptotic efficiency

This and MLE are covered in 420.

There are many properties one can ask of estimators $\hat{\theta}$ of parameters θ

- 1. Unbiased: $\mathbb{E}\left[\hat{\theta}\right] \theta = 0$ 2. Consistent: $\hat{\theta} \xrightarrow{n \to \infty} \theta$ 3. Efficient: $\mathbb{V}\left[\hat{\theta}\right]$ is the smallest of all unbiased estimators
- 4. Asymptotically efficient: Maybe not efficient for every n, but in the limit, the variance is the smallest of all unbiased estimators.
- 5. Minimax: over all possible estimators in some class, this one has the smallest MSE for the worst problem.
- 6. ...

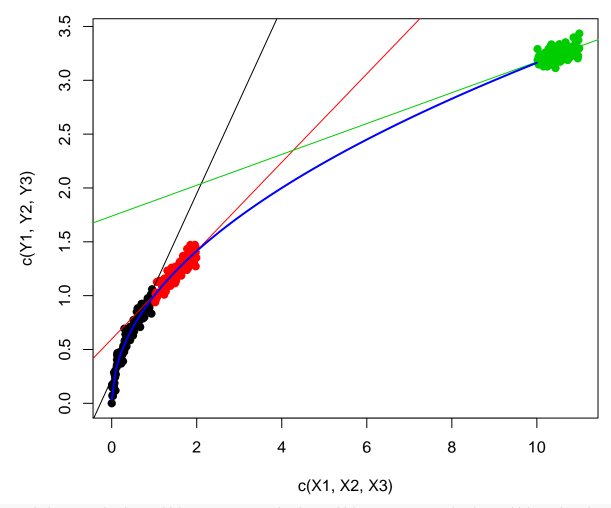
Problems with R-squared

$$R^{2} = 1 - \frac{SSE}{\sum_{i=1}^{n} (Y_{i} - \overline{Y})^{2}} = 1 - \frac{MSE}{\frac{1}{n} \sum_{i=1}^{n} (Y_{i} - \overline{Y})^{2}} = 1 - \frac{SSE}{SST}$$

- This gets spit out by software
- X and Y are both normal with (empirical) correlation r, then $R^2 = r^2$
- In this nice case, it measures how tightly grouped the data is about the regression line
- Data that is tightly grouped about the regression line can be predicted accurately by the regression line.
- Unfortunately, the implication does not go both ways.
- High R^2 can be achieved in many ways, same with low R^2
- You should just ignore it completely (and the adjusted version), and encourage your friends to do the same

High R-squared with non-linear relationship

```
genY <- function(X, sig) Y = sqrt(X)+sig*rnorm(length(X))</pre>
sig=0.05; n=100
X1 = runif(n,0,1); Y1 = genY(X1,sig)
X2 = runif(n,1,2); Y2 = genY(X2,sig)
X3 = runif(n,10,11); Y3 = genY(X3,sig)
plot(c(X1,X2,X3),c(Y1,Y2,Y3),pch=19,col=rep(1:3,each=n))
abline(lm(Y1~X1), col=1); abline(lm(Y2~X2), col=2); abline(lm(Y3~X3), col=3)
curve(sqrt(x),.001,10,col=4,lwd=2, add=TRUE)
```



round(c(summary(lm(Y1~X1))\$r.sq, summary(lm(Y2~X2))\$r.sq, summary(lm(Y3~X3))\$r.sq), 3)

[1] 0.907 0.826 0.371