# STAT 331 - Applied Linear Models

Cameron Roopnarine

Last updated: March 8, 2021

# **Contents**

Co	Contents 1				
1	Simple Linear Regression         1.1 Introduction          1.2 Inference          1.3 Prediction          1.4 Application          1.4.1 R Demo	4 6 8 10 11			
2	2.2 Multivariate Normal Distribution 2.3 Inference 2.4 Application 2.5 Prediction 2.6 Categorical Predictors 2.6.1 R Demo 2.7 Analysis of Variance 2.8 Coefficient of Determination 2.9 General Linear Hypothesis Tests Based on F Distribution 2.10 ANOVA F Test 2.10.1 R Demo 2.11 Multicollinearity 2.12 Detection of Multicollinearity 2.12.1 R Demo 2.13 Model Selection Criteria 2.14 Model Selection Basic Strategies 2.14.1 R Demo 2.15 Residual Plots for Linear Regression Assumptions 2.15.1 R Demo 2.16 Addressing Problems With Regression Model Assumptions 2.16.1 R Demo 2.17 Effects of Individual Observations 2.17.1 R Demo 2.18 Prediction Error	15 18 18 21 24 24 32 32 34 35 50 51 58 61 62 66 67 75 76 88 90 97 98			
	2.19.1 R Demo 2.20 Combining Cross Validation With Model Selection 2.21 Iterative Conditional Minimization	99 102			

CONTENTS	2

3	Gen	eralized Linear Models	106
	3.1	Beyond STAT 331: Generalized Linear Models and Logistic Regression	106
	3.2	Last Lecture!	108

## Introduction

## LECTURE 1 | 2020-09-08

## **DEFINITION 0.0.1: Response variable**

A **response** (**dependent**) **variable** is the primary variable of interest, denoted by a capital roman letter Y.

## **DEFINITION 0.0.2: Explanatory Variable**

An **explanatory** (independent, predictor) variable are variables that impact the response, denoted by  $x_i$  for i = 1, ..., p.

#### **DEFINITION 0.0.3: Regression Model**

A **regression model** deals with modelling the functional relationship between a response variable and one or more explanatory variables.

#### **EXAMPLE 0.0.4: Alligators in Florida**

Let Y be the length in metres of an alligator and  $x_1 := \{0,1\}$  (male or female). The mass in an alligators stomach consists of fish  $(x_2)$ , invertebrates  $(x_3)$ , reptiles  $(x_4)$ , birds  $(x_5)$ , and other  $(x_6,\ldots,x_p)$ . We imagine we can explain Y in terms of  $(x_1,\ldots,x_p)$  using some function such that  $Y = f(x_1,\ldots,x_p)$ .

In this course, we will be looking at linear models.

#### **DEFINITION 0.0.5: Linear model**

A general **linear model** is defined as  $Y=\beta_0+\beta_1x_1+\cdots+\beta_px_p+\varepsilon$  where Y is the response variable,  $(x_1,\ldots,x_p)$  are the p explanatory variables,  $(\beta_0,\beta_1,\ldots,\beta_p)$  are the model parameters, and  $\varepsilon$  is the random error. We assume that  $(x_1,\ldots,x_p)$  are fixed constants,  $\beta_0$  is the intercept of Y,  $(\beta_1,\ldots,\beta_p)$  all quantify effect on  $x_j$  on Y, and  $\varepsilon\sim\mathcal{N}(0,\sigma^2)$ .

#### **REMARK 0.0.6**

In general, the model will not perfectly explain the data. "All models are wrong, but some are useful."

$$Y \sim \mathcal{N}(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p, \sigma^2) \text{ since } \mathbb{E}[Y] = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p \text{ and } \mathbb{V}(Y) = \mathbb{V}(\varepsilon) = \sigma^2.$$

## Chapter 1

## Simple Linear Regression

## LECTURE 2 | 2020-09-09

## 1.1 Introduction

## **DEFINITION 1.1.1: Simple linear regression**

A simple linear regression is a linear model that uses only one explanatory variable; that is,  $Y = \beta_0 + \beta_1 x + \varepsilon$ . The data in a simple linear regression consists of pairs  $(x_i, y_i)$  where i = 1, ..., n.

#### **REMARK 1.1.2**

Before fitting any model, we might want to make a scatter plot to visualize if there is a linear relationship between x and y, or calculate the *correlation*.

#### **DEFINITION 1.1.3: Correlation**

The **correlation** of random variables X and Y is  $\rho_{XY} = \frac{\operatorname{Cov}(X,Y)}{\operatorname{Sd}(X)\operatorname{Sd}(Y)}$ .

#### **DEFINITION 1.1.4: Sample correlation**

The **sample correlation** of all pairs  $(x_i, y_i)$  is

$$\begin{split} r &= \frac{\frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x}) (y_i - \bar{y})}{\sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})^2} \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (y_i - \bar{y})^2}} \\ &= \frac{\sum_{i=1}^{n} (x_i - \bar{x}) (y_i - \bar{y})}{\sqrt{\sum_{i=1}^{n} (x_i - \bar{x})^2} \sqrt{\sum_{i=1}^{n} (y_i - \bar{y})^2}} \\ &= \frac{S_{xy}}{\sqrt{S_{xx} S_{yy}}} \end{split}$$

#### **REMARK 1.1.5**

The sample correlation measures the strength and direction of the linear relationship between x and y. Note that  $-1\leqslant r\leqslant 1$ . If  $|r|\approx 1$ , then there is a strong linear relationship, and if  $|r|\approx 0$  then there is a lack of linear relationship. Also, if r>0, then there is a positive relationship, and if r<0 then there is a negative relationship. It does not tell us how to predict y from x. To do so, we need to estimate  $\beta_0$  and  $\beta_1$ .

#### **DEFINITION 1.1.6: Simple linear regression model**

For data  $(x_i, y_i)$  for i = 1, ..., n, the **simple linear regression model** is  $Y_i = \beta_0 + \beta_1 x_i + \varepsilon_i$  with the assumption that  $\varepsilon_i \stackrel{\text{iid}}{\sim} \mathcal{N}(0, \sigma^2)$ . Therefore,  $Y_i \sim \mathcal{N}(\mu_i = \beta_0 + \beta_1 x_i, \sigma^2)$ .

#### **DEFINITION 1.1.7: Least squares**

The method of estimating  $\beta_0$  and  $\beta_1$  by minimizing  $S(\beta_0,\beta_1)=\sum_{i=1}^n(y_i-(\beta_0+\beta_1x_i))^2$  is referred to as the **method of the least squares**.

#### **REMARK 1.1.8**

The least squares is equivalent to maximum likelihood estimate when  $\varepsilon_i \stackrel{\mathrm{iid}}{\sim} \mathcal{N}(0,\sigma^2)$ .

#### THEOREM 1.1.9: Least Square Estimates (LSEs) for SLR

Minimizing  $S(\beta_0, \beta_1)$ , gives the least square estimates

$$\hat{\beta}_0 = \bar{y} - \hat{\beta}_1 \bar{x} \quad \text{and} \quad \hat{\beta}_1 = \frac{\sum_{i=1}^n (x_i - \bar{x})(y_i - \bar{y})}{\sum_{i=1}^n (x_i - \bar{x})^2} = \frac{S_{xy}}{S_{xx}}$$

#### **Proof of: 1.1.9**

$$\begin{split} \frac{\partial S}{\partial \beta_0} &= 2 \sum_{i=1}^n \big[ y_i - (\beta_0 + \beta_1 x_i) \big] (-1) \text{ and } \frac{\partial S}{\partial \beta_1} = 2 \sum_{i=1}^n \big[ y_i - (\beta_0 + \beta_1 x_i) \big] (-x_i). \\ \text{Now,} \\ \frac{dS}{d\beta_0} &\coloneqq 0 \iff \sum_{i=1}^n y_i - n\beta_0 - \beta_1 \sum_{i=1}^n x_i = 0 \iff \beta_0 = \bar{y} - \beta_1 \bar{x} \\ \frac{dS}{d\beta_1} &\coloneqq 0 \iff \sum_{i=1}^n \big[ y_i - \bar{y} + \beta_1 \bar{x} - \beta_1 x_i \big] x_i = 0 \\ &\iff \sum_{i=1}^n x_i (y_i - \bar{y}) - \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) = 0 \\ &\iff \beta_1 = \frac{\sum_{i=1}^n x_i (y_i - \bar{y})}{\sum_{i=1}^n x_i (x_i - \bar{x})} = \frac{\sum_{i=1}^n (x_i - \bar{x}) (y_i - \bar{y})}{\sum_{i=1}^n (x_i - \bar{x})^2} = \frac{S_{xy}}{S_{xx}} \end{split}$$

#### **REMARK 1.1.10**

We use a hat on the  $\beta$ 's to show that they are estimates.

#### **DEFINITION 1.1.11: Fitted value, Residual**

The expression  $\hat{\mu}_i = \hat{\beta}_0 + \hat{\beta}_1 x_i$  is called the **fitted value** that corresponds to the  $i^{\text{th}}$  observation with  $x_i$  as the explanatory variable. The difference between  $y_i$  and  $\hat{\mu}_i$ , and  $e_i = y_i - \hat{\mu}_i$  is referred to as the **residual**. It is the vertical distance between the observation  $y_i$  and the estimated line  $\hat{\mu}_i$  evaluated at  $x_i$ .

## LECTURE 3 | 2020-09-14

## 1.2 Inference

For  $Y_i \sim \mathcal{N}(\beta_0 + \beta_1 x_i, \sigma^2)$ , the equation of fitted line is given by  $y = \hat{\beta}_0 + \hat{\beta}_1 x$ . Our interpretation of the parameters is as follows.

- $\hat{\beta}_0$  is the estimate of the expected response when x = 0 (but not always meaningful if outside range of  $x_i$ 's in data)
- $\hat{\beta}_1$  is the estimate of expected change in response for unit increase in x
- $\sigma^2$  is the "variability around the line" where  $\sigma^2=\mathbb{V}(\varepsilon_i)=\mathbb{V}(Y_i)$

Q: How should we estimate  $\sigma^2$ ?

$$\varepsilon_i = Y_i - (\beta_0 + \beta_1 x_i)$$
 and  $e_i = y_i - (\hat{\beta}_0 + \hat{\beta}_1 x_i)$ 

Our intuition tells us to use variability in the residuals to estimate  $\sigma^2$ , so we use

$$\hat{\sigma}^2 = \frac{\sum_{i=1}^n (e_i - \bar{e})^2}{n-2} = \frac{\sum_{i=1}^n e_i^2}{n-2}$$

where the first term looks like sample variance of  $e_i$ 's. The second equality follows since  $\bar{e} = \bar{y} - (\hat{\beta}_0 + \hat{\beta}_1 \bar{x}) = 0$  by definition of our  $\beta_0$  estimate.

#### **DEFINITION 1.2.1: Residual sum of squares**

 $SS(Res) = \sum_{i=1}^{n} (y_i - \hat{\mu}_i)^2 = \sum_{i=1}^{n} e_i^2$ , is known as the **residual (error) sum of squares**.

#### **REMARK 1.2.2**

The n-2 will be looked at in more detail later, but for now it suffices to say that the degrees of freedom is n-2 or equivalently, n- number of parameters estimated. It allows  $\hat{\sigma}^2$  to be an unbiased estimator for the true value of  $\sigma^2$ ; that is,  $\mathbb{E}[\hat{\sigma}^2] = \sigma^2$  whenever  $\hat{\sigma}^2$  is viewed as a random variable.

#### THEOREM 1.2.3: Linear Combination of Independent Normal Random Variables

If  $Y_i \sim \mathcal{N}(\mu_i, \sigma^2)$ , i = 1, ..., n independently, then

$$\sum_{i=1}^n a_i Y_i \sim N\left(\sum_{i=1}^n a_i \mu_i, \sum_{i=1}^n a_i^2 \sigma_i^2\right)$$

#### **Proof of: 1.2.3**

The proof is completed in STAT 330 with moment generating functions.

Viewing  $\hat{\beta}_1$  as a random variable:

$$\hat{\beta}_1 = \frac{\sum_{i=1}^n (x_i - \bar{x})(Y_i - \bar{Y})}{\sum_{i=1}^n (x_i - \bar{x})^2} = \frac{\sum_{i=1}^n (x_i - \bar{x})Y_i - \bar{Y} \sum_{i=1}^n (x_i - \bar{x})}{\sum_{i=1}^n (x_i - \bar{x})x_i - \bar{x} \sum_{i=1}^n (x_i - \bar{x})} = \frac{\sum_{i=1}^n (x_i - \bar{x})Y_i}{\sum_{i=1}^n (x_i - \bar{x})x_i} = \sum_{i=1}^n a_i Y_i$$

where  $a_i = \frac{x_i - \bar{x}}{\sum_{i=1}^n x_i (x_i - \bar{x})}$ . Therefore,

$$\mathbb{E}[\hat{\beta}_1] = \sum_{i=1}^n a_i \mathbb{E}[Y_i] = \frac{\sum_{i=1}^n (x_i - \bar{x})(\beta_0 + \beta_1 x_i)}{\sum_{i=1}^n x_i (x_i - \bar{x})} = \frac{\beta_0 \sum_{i=1}^n (x_i - \bar{x}) + \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x})}{\sum_{i=1}^n x_i (x_i - \bar{x})} = \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) + \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) = \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) + \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) + \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) = \beta_1 \sum_{i=1}^n x_i (x_i - \bar{x}) + \beta_1 \sum_{i=1}^n x_i (x_i -$$

Now, we calculate the variance of  $\hat{\beta}_1$ :

$$\mathbb{V}(\hat{\beta}_1) = \sum_{i=1}^n a_i^2 \mathbb{V}(Y_i) = \sigma^2 \frac{\sum_{i=1}^n (x_i - \bar{x})^2}{[\sum_{i=1}^n x_i (x_i - \bar{x})]^2} = \sigma^2 \frac{\sum_{i=1}^n (x_i - \bar{x})^2}{[\sum_{i=1}^n (x_i - \bar{x})^2]^2} = \frac{\sigma^2}{S_{xx}}$$

Using our calculations from  $\hat{\beta}_1$ , and viewing  $\hat{\beta}_0$  as a random variable:

$$\mathbb{E}[\hat{\beta}_0] = \mathbb{E}[\bar{Y}] - \bar{x}\mathbb{E}[\hat{\beta}_1] = \mathbb{E}\left[\frac{\sum_{i=1}^n Y_i}{n}\right] - \bar{x}\beta_1 = \frac{\sum_{i=1}^n (\beta_0 + \beta_1 x_i)}{n} - \beta_1 \bar{x} = \beta_0 + \beta_1 \bar{x} - \beta_1 \bar{x} = \beta_0$$

Now, we calculate the variance of  $\hat{\beta}_0$ :

$$\mathbb{V}(\hat{\beta}_1) = \mathbb{V}(\bar{Y} - \beta_1 \bar{x}) = \mathbb{V}(\bar{Y}) + (-\bar{x}^2)\mathbb{V}(\beta_1) = \mathbb{V}\left(\frac{\sum_{i=1}^n Y_i}{n}\right) + \bar{x}^2\bigg(\frac{\sigma^2}{S_{xx}}\bigg) = \frac{n\sigma^2}{n^2} + \frac{\sigma^2 x^2}{S_{xx}}$$

Also, since  $\hat{\beta}_1$  and  $\hat{\beta}_0$  are linear combination of Normal random variables, they follow a Normal distribution. Therefore, we get the following theorem.

#### **THEOREM 1.2.4: Distribution of LSEs**

The distribution of the least square estimates are given by

$$\hat{\beta}_1 \sim \mathcal{N} \bigg( \beta_1, \frac{\sigma^2}{S_{xx}} \bigg) \quad \text{ and } \quad \hat{\beta}_0 \sim \mathcal{N} \bigg( \beta_0, \sigma^2 \bigg( \frac{1}{n} + \frac{\bar{x}^2}{S_{xx}} \bigg) \bigg)$$

Since  $\mathbb{E}[\hat{\beta}_1] = \beta_1$ , we say  $\hat{\beta}_1$  is an unbiased estimator of  $\beta_1$ . This implies that as  $n \to \infty$ , the average of the estimates  $\hat{\beta}_1$ ; that is,  $\mathbb{E}[\hat{\beta}_1]$  coincides with the true value of  $\beta_1$ . A similar argument can be made for  $\beta_0$ .

Then, 
$$\frac{\hat{\beta}_1 - \beta_1}{\sigma/\sqrt{S_{xx}}} \sim \mathcal{N}(0,1)$$
, but  $\sigma$  is unknown, so need to use  $\hat{\sigma}$  to get  $\frac{\hat{\beta}_1 - \beta_1}{\hat{\sigma}/\sqrt{S_{xx}}} \sim t(n-2)$ .

## **DEFINITION 1.2.5: Standard deviation and standard error of** $\hat{\beta}_1$

The standard deviation of  $\hat{\beta}_1$  is defined as  $\mathrm{Sd}(\hat{\beta}_1) = \sigma/\sqrt{S_{xx}}$ . The estimated standard deviation of  $\hat{\beta}_1$  is also referred to as the standard error of the estimate  $\hat{\beta}_1$ , and we write  $\mathrm{Se}(\hat{\beta}_1) = \hat{\sigma}/\sqrt{S_{xx}}$ .

#### **DEFINITION 1.2.6: Student** *t* **distribution**

Suppose  $Z \sim \mathcal{N}(0,1)$  and  $U \sim \chi^2(\nu)$ , with Z and U independent. Then,  $T = Z/\sqrt{U/\nu}$  has a **Student** t **distribution** with  $\nu$  degrees of freedom.

#### **THEOREM 1.2.7**

For a simple linear regression model,

$$\frac{\hat{\sigma}^2(n-2)}{\sigma^2} = \frac{\mathit{SS}(\mathit{Res})}{\sigma^2} \sim \chi^2(n-2)$$

#### **Proof of: 1.2.7**

Too hard for sure.

Using the theorem stated, we justify the fact that replacing  $\sigma$  with  $\hat{\sigma}$  gives us a t(n-2) distribution.

$$\frac{\hat{\beta}_1 - \beta_1}{\hat{\sigma}/\sqrt{S_{xx}}} = \frac{\frac{\hat{\beta}_1 - \beta_1}{\sigma/\sqrt{S_{xx}}}}{\sqrt{\frac{\hat{\sigma}^2(n-2)}{\sigma^2} \left(\frac{1}{n-2}\right)}} = \frac{Z}{\sqrt{U/\nu}} = T \sim t(n-2)$$

where  $\frac{\hat{\sigma}^2(n-2)}{\sigma^2}=U$ ,  $\nu=n-2$ , and  $Z=\frac{\hat{eta}_1-eta_1}{\hat{\sigma}/\sqrt{S_{xx}}}$ . A (1-lpha) confidence interval for  $eta_1$  is

$$\hat{\beta}_1 \pm c \operatorname{Se}(\hat{\beta}_1)$$

where c is the  $1-\frac{\alpha}{2}$  quantile of t(n-2); that is,  $P(|T|\leqslant c)=1-\alpha$  or  $P(T\leqslant c)=1-\frac{\alpha}{2}$  where  $T\sim t(n-2)$ .

<u>Hypothesis test</u>:  $H_0$ :  $\beta=0$  versus  $H_A$ :  $\beta_1\neq 0$ . If  $H_0$  is true, then  $\hat{\beta}_1/\mathrm{Se}(\hat{\beta}_1)\sim t(n-2)$ , so calculate the **t statistic**  $t=\hat{\beta}_1/\mathrm{Se}(\hat{\beta}_1)$ , and reject  $H_0$  at level  $\alpha$  if |t|>c where c is  $1-\frac{\alpha}{2}$  quantile of t(n-2). Therefore, p-value  $=P(|T|\geqslant |t|)=2P(T\geqslant |t|)$ .

## LECTURE 4 | 2020-09-16

## 1.3 Prediction

Suppose we want to predict the response y for a new value of x, say  $x=x_0$ . Then, SLR model says  $Y_0 \sim \mathcal{N}(\beta_0+\beta_1x_0,\sigma^2)$  where  $Y_0$  is a random variable for response when  $x=x_0$ ; that is,  $\hat{Y}_0=\hat{\beta}_0+\hat{\beta}_1x_0$ . The fitted model predicts the value of y to be  $\hat{y}_0=\hat{\beta}_0+\hat{\beta}_1x_0$ .

Also,  $\mathbb{E}[\hat{Y}_0] = \mathbb{E}[\hat{\beta}_0] + x_0 \mathbb{E}[\hat{\beta}_1] = \beta_0 + \beta_1 x_0 = \mathbb{E}[Y_0]$ , since  $\hat{\beta}_i$  for i = 0, 1 are unbiased. Therefore, we can say that  $\hat{Y}_0$  is an unbiased estimate of the random variable for the mean of  $Y_0$ . For the variance of  $\hat{Y}_0$  we

write

$$\begin{split} \hat{Y}_0 &= \hat{\beta}_0 + \hat{\beta}_1 x_0 = \bar{Y} - \hat{\beta}_1 \bar{x} + \hat{\beta}_1 x_0 \\ &= \bar{Y} + \hat{\beta}_1 (x_0 - \bar{x}) \\ &= \sum_{i=1}^n \left[ \frac{Y_i}{n} + (x_0 - \bar{x}) \left( \frac{(x_i - \bar{x})(Y_i - \bar{Y})}{S_{xx}} \right) \right] \\ &= \sum_{i=1}^n \left[ \frac{Y_i}{n} + (x_0 - \bar{x}) \left( \frac{(x_i - \bar{x})Y_i}{S_{xx}} \right) \right] \\ &= \sum_{i=1}^n \left[ \frac{1}{n} + \frac{(x_0 - \bar{x})(x_i - \bar{x})}{S_{xx}} \right] Y_i \\ &= \sum_{i=1}^n a_i Y_i \end{split}$$

where  $a_i = \frac{1}{n} + \frac{(x_0 - \bar{x})(x_i - \bar{x})}{S_{xx}}$ . Therefore,

$$\begin{split} \mathbb{V}(Y_0) &= \sum_{i=1}^n \left[ \frac{1}{n} + \frac{(x_0 - \bar{x})(x_i - \bar{x})}{S_{xx}} \right]^2 \\ &= \sum_{i=1}^n \left[ \frac{1}{n^2} + \frac{2(x_0 - \bar{x})(x_i - \bar{x})}{nS_{xx}} + \frac{(x_0 - \bar{x})^2(x_i - \bar{x})^2}{(S_{xx})^2} \right] \\ &= \sum_{i=1}^n \left[ \frac{1}{n^2} \right] + \frac{2(x_0 - \bar{x})}{nS_{xx}} \sum_{i=1}^n (x_i - \bar{x}) + \frac{(x_0 - \bar{x})^2}{(S_{xx})^2} \sum_{i=1}^n (x_i - \bar{x})^2 \\ &= \frac{1}{n} + \frac{2(x_0 - \bar{x})}{S_{xx}} (0) + \frac{(x_0 - \bar{x})^2}{(S_{xx})^2} (S_{xx}) \\ &= \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \end{split}$$

We proved the following theorem.

#### **THEOREM 1.3.1: Distribution of Prediction**

The distribution of the prediction random variable is given by

$$\hat{Y}_0 \sim \mathcal{N} \bigg( \beta_0 + \beta_1 x_0, \sigma^2 \bigg( \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \bigg) \bigg)$$

#### **DEFINITION 1.3.2: Prediction error**

The random variable for **prediction error** is defined as  $Y_0 - \hat{Y}_0$  where  $Y_0$  and  $\hat{Y}_0$  are independent and  $\hat{Y}_0$  is a function of  $Y_1, \dots, Y_n$ .

$$\begin{split} \mathbb{E}[Y_0 - \hat{Y}_0] &= \mathbb{E}[Y_0] - \mathbb{E}[\hat{Y}_0] = 0 \\ \mathbb{V}(Y_0 - \hat{Y}_0) &= \mathbb{V}(Y_0) + (-1)^2 \mathbb{V}(\hat{Y}_0) = \sigma^2 + \sigma^2 \bigg(\frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}}\bigg) \end{split}$$

We proved the following theorem.

#### **THEOREM 1.3.3: Distribution of Prediction Error**

The distribution of the prediction error is given by

$$Y_0 - \hat{Y}_0 \sim \mathcal{N}\bigg(0, \sigma^2\bigg(1 + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}}\bigg)\bigg)$$

Since  $\sigma$  is unknown, we use  $\hat{\sigma}$  and get the following:

$$\frac{Y_0 - \hat{Y}_0}{\hat{\sigma} \sqrt{1 + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}}}} \sim t(n - 2)$$

Intuition for prediction error composed of 2 terms:

- $\mathbb{V}(Y_0)$ : random error of new observation
- $\mathbb{V}(\hat{Y}_0)$  (predictor): estimating  $\beta_0$  and  $\beta_1$

Those are 2 sources of uncertainty.

#### **REMARK 1.3.4**

Be careful that the prediction may not make sense if  $x_0$  is outside the range of the  $x_i$ 's in the data.

A  $(1-\alpha)$  prediction interval for the mean response  $y_0 = \beta_0 + \beta_1 x_0$  at  $x_0$  is

$$\hat{y}_0 \pm c\,\hat{\sigma}\sqrt{1+\frac{1}{n}+\frac{(x_0-\bar{x})^2}{S_{xx}}}$$

where c is the  $1-\frac{\alpha}{2}$  quantile of t(n-2).

#### **Application** 1.4

#### **EXAMPLE 1.4.1: Orange production 2018 in FL**

We are given the following information.

- *x*: acres
- *y*: # boxes of oranges (thousands)
- $(x_i, y_i)$  recorded for each of 25 FL counties
- r = 0.964
- $\bar{x} = 16133$
- $\bar{y} = 1798$
- $\begin{array}{l} \bullet \ \, S_{xx} = 1.245 \times 10^{10} \\ \bullet \ \, S_{xy} = 1.453 \times 10^9 \\ \end{array}$

Now,  $\beta_1 = S_{xy}/S_{xx} = 0.1167$  has a positive slope, therefore x and y are positively correlated. The expected number of boxes produced is estimated to be about 117 higher per an additional acre.

Computing  $\beta_0 = \bar{y} - \beta_1 \bar{x} = -85.3$ , we see that it is not meaningful to interpret, since it is the expected production if there were 0 acres (outside the range of  $x_i$ ) as no county has x = 0.

Now suppose  $SS(Res) = 1.31 \times 10^7$  the residuals are the differences between  $y_i$  and the fitted regression

• 
$$\hat{\sigma}^2 = \frac{\sum_{i=1}^n e_i^2}{n-2} = \frac{1.31 \times 10^7}{25-2} = 5.7 \times 10^5$$

• 
$$\operatorname{Se}(\hat{\beta}_1) = \frac{\hat{\sigma}}{\sqrt{S_{xx}}} = 0.00676$$

- To test  $H_0$ :  $\beta_1=0$ , calculate  $t=(\hat{\beta}_1-0)/\mathrm{Se}(\hat{\beta}_1)=0.1167/0.00676\approx 17.3$ , then elect the 0.975 quantile (for demonstration purposes) of t(23) which is 2.07.
- Note that 17.3 is very unlikely to see in t(23).

Since  $17.3 \gg 2.07$ , we reject  $H_0$  at  $\alpha = 0.05$  level, and conclude there's a significant linear relationship between acres and oranges produced.

The 95% confidence interval for  $\beta_1$  is given by  $0.1167 \pm 2.07(0.00676)$ , which does not contain 0.

$$p$$
-value =  $P(|t_{23}| \ge 17.3) = 2P(t_{23} \ge 17.3) \approx 1.2 \times 10^{-14}$ 

Predict the # of boxes in thousands produced if we had 10000 acres to grow oranges.

$$\hat{\beta}_0 + \hat{\beta}_1 x_0 = -85.3 + (0.1167)(10000) \approx 1082$$

The 95% prediction interval is given by

$$1082 \pm 2.07 \sqrt{5.69 \times 10^5} \sqrt{1 + \frac{1}{25} + \frac{(6133)^2}{1.245 \times 10^{10}}} = [-512.0407, 2675.595]$$

#### **REMARK 1.4.2**

We are **not** trying to establish causation.

#### 1.4.1 R Demo

```
# Read data from florange.csv and input it into the dat vector.
dat <- read.csv("csv/florange.csv")</pre>
# Done to make the predict function work well.
x <- dat$acres
y <- dat$boxes
# Output the first 6 rows in dat.
head(dat)
##
        county boxes acres
## 1
       Brevard
                   51
                        696
## 2 Charlotte
                  821 13447
## 3
       Collier 2088 29351
## 4
        DeSoto 7688 66365
## 5
        Glades
                 368 5396
## 6
        Hardee 5306 43126
# Draw a scatterplot with x-axis as 'acres' and y-axis as 'boxes'.
plot(x, y)
                                                                                  0
                                                                           0
                                                                                     0
     2000 4000 6000
                                                            0
                                                                                0
                                             0
                            0
                     0
                           0
                     10000
                                          30000
             0
                                20000
                                                      40000
                                                                 50000
                                                                            60000
                                                 Χ
# Compute some common variables with common functions.
r <- cor(x, y)
xbar <- mean(x)
ybar <- mean(y)</pre>
cat("r:", r, "xbar:", xbar, "ybar:", ybar)
## r: 0.9635098 xbar: 16132.64 ybar: 1797.56
Therefore, r = 0.9635098, \bar{x} = 16132.64, and \bar{y} = 1797.56.
# Compute some common variables manually.
Sxx \leftarrow sum((x - xbar)^2)
Sxy \leftarrow sum((x - xbar) * (y - ybar))
cat("Sxx: ", Sxx, "Sxy: ", Sxy)
```

## Sxx: 12450023404 Sxy: 1453128337

```
Therefore, S_{xx} = 12450023404 = 1.245 \times 10^{10} and S_{xy} = 1453128337 = 1.453 \times 10^{9}.
# R's lm function fits linear models
lm.1 < - lm(y ~ x)
summary(lm.1)
##
## Call:
## lm(formula = y \sim x)
## Residuals:
                    1Q
                          Median
##
         Min
                                         3Q
                                   106.46 1677.32
## -2470.81
                 -6.17
                           71.72
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept) -85.391989 186.178031 -0.459
## x
                   0.116717
                               0.006761 17.263 1.16e-14 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 754.4 on 23 degrees of freedom
## Multiple R-squared: 0.9284, Adjusted R-squared: 0.9252
## F-statistic: 298 on 1 and 23 DF, p-value: 1.164e-14
From the summary, we can see that \hat{\beta}_0 = -85.391989, \hat{\beta}_1 = 0.116717, Se(\hat{\beta}_1) = 0.006761, t = 17.263,
p-value = 1.64 \times 10^{-14}, and \hat{\sigma} = 754.4.
# Sum Squared Fitted Values
sum(lm.1$fitted.values ^ 2)
## [1] 250385207
# Sum Squared Residuals
sum(lm.1$residuals ^ 2)
## [1] 13089860
Therefore, SS(Res) = \sum_{i=1}^{n} e_i^2 = 13089860 = 1.31 \times 10^7.
# Manual calculation of sigma^2 estimate
sum(lm.1$residuals^2) / 23
## [1] 569124.3
Therefore, \hat{\sigma}^2 = 69124.3 = 5.7 \times 10^5.
# Manual calculation of sigma estimate
sqrt(sum(lm.1$residuals ^ 2) / 23)
## [1] 754.4033
Therefore, \hat{\sigma} = 754.4.
# t distribution values
qt(0.975, 23)
## [1] 2.068658
Therefore, c = 2.07.
```

```
# 95% confidence interval
confint(lm.1)

## 2.5 % 97.5 %

## (Intercept) -470.5305905 299.7466119

## x 0.1027305 0.1307034

# 95% prediction interval with predicted boxes if we had 10000 acres
predict(lm.1, data.frame(x = 10000), interval = "prediction")

## fit lwr upr
## 1 1081.777 -512.0407 2675.595
```

Q: Is  $\sigma$  the same for all values of y?

A: It appears to not in the sense that the variance appears to be higher with respect to higher acres. Sigma will be smaller when there's less acres. Later, this will be testing equal variance or homoscedastic assumption. Later, when we talk about variable transformations we can consider taking the logarithm.

Q: Are the error terms plausibly independent? In other words, does knowing one  $e_i$  (residual) help predict  $e_j$  (another residual) for a different county?

A: There's diagnostics for checking this. However, intuitively there could be some common factors at play when two counties are geographically close.

## Chapter 2

## **Multiple Linear Regression**

LECTURE 5 | 2020-09-21

## 2.1 Random Vectors

## **DEFINITION 2.1.1: Multiple linear regression**

A multiple linear regression (MLR) model is defined as

$$Y = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p + \varepsilon$$

which links a response variable y to several independent explanatory variables  $x_1, x_2, \dots, x_p$ .

#### **EXAMPLE 2.1.2: Rocket MLR**

- $x_1$ : nozzle area (large or small, 0 or 1)
- $x_2$ : mixture in propellant, ratio oxidized fuel
- Y: thrust

Want to develop linear relationship between response y and  $x_1, x_2$ ; that is, we want to develop a linear relationship between thrust and both nozzle area and mixture in propellant.

In multiple linear regression, there are n observations, where each consists of p response variables  $(y_i)$ , and p explanatory variables  $(x_{i1}, x_{i2}, \dots, x_{ip})$ . Then,

$$Y_i \sim N(\underbrace{\beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_p x_{ip}}_{\mathbb{E}[Y_i] = \mu_i}, \sigma^2)$$

or  $Y_i=\mu_i+\varepsilon_i$  where  $\varepsilon_i\stackrel{\mathrm{iid}}{\sim}\mathcal{N}(0,\sigma^2).$  We can write in vector/matrix form

$$\begin{bmatrix} Y_1 \\ Y_2 \\ \vdots \\ Y_n \end{bmatrix}_{n \times 1} = \begin{bmatrix} \beta_0 + \beta_1 x_{11} + \dots + \beta_p x_{1p} \\ \beta_0 + \beta_1 x_{21} + \dots + \beta_p x_{2p} \\ \vdots \\ \beta_0 + \beta_1 x_{n1} + \dots + \beta_p x_{np} \end{bmatrix} + \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1}$$

Which we can more commonly write as  $Y = X\beta + \varepsilon$  where

$$\boldsymbol{Y} = \begin{bmatrix} Y_1 \\ Y_2 \\ \vdots \\ Y_n \end{bmatrix}_{n \times 1} \boldsymbol{X} = \begin{bmatrix} 1 & x_{11} & x_{12} & \cdots & x_{1(p-1)} & x_{1p} \\ 1 & x_{21} & x_{22} & \cdots & x_{2(p-1)} & x_{2p} \\ \vdots & & \ddots & & \vdots \\ 1 & x_{(n-1)1} & x_{(n-1)2} & \cdots & x_{(n-1)(p-1)} & x_{(n-1)p} \\ 1 & x_{n1} & x_{n2} & \cdots & x_{n(p-1)} & x_{np} \end{bmatrix}_{n \times (p+1)} \boldsymbol{\beta} = \begin{bmatrix} \beta_0 \\ \vdots \\ \beta_p \end{bmatrix}_{(p+1) \times 1} \boldsymbol{\varepsilon} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} = \begin{bmatrix} \varepsilon_1 \\ \varepsilon_1 \\ \vdots \\ \varepsilon_n \end{bmatrix}_{n \times 1} \boldsymbol{\zeta} =$$

#### **DEFINITION 2.1.3: Random vector**

We call  $Y = (Y_1, Y_2, \dots, Y_n)^{\top}$  a random vector.

#### **DEFINITION 2.1.4: Mean vector**

The **mean vector** of Y is defined as  $\mathbb{E}[Y] = (\mathbb{E}[Y_1], \mathbb{E}[Y_2], \dots, \mathbb{E}[Y_n])^{\top}$ .

#### **DEFINITION 2.1.5: Covariance matrix**

The **covariance matrix** (or **variance-covariance matrix**) of *Y* is defined as

$$\mathbb{V}(\boldsymbol{Y}) = \begin{bmatrix} \mathbb{V}(Y_1) & \operatorname{Cov}(Y_1,Y_2) & \cdots & \operatorname{Cov}(Y_1,Y_{n-1}) & \operatorname{Cov}(Y_1,Y_n) \\ \operatorname{Cov}(Y_2,Y_1) & \mathbb{V}(Y_2) & \cdots & \operatorname{Cov}(Y_2,Y_{n-1}) & \operatorname{Cov}(Y_2,Y_n) \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ \operatorname{Cov}(Y_{n-1},Y_1) & \operatorname{Cov}(Y_{n-1},Y_2) & \cdots & \mathbb{V}(Y_{n-1}) & \operatorname{Cov}(Y_{n-1},Y_n) \\ \operatorname{Cov}(Y_n,Y_1) & \operatorname{Cov}(Y_n,Y_2) & \cdots & \operatorname{Cov}(Y_n,Y_{n-1}) & \mathbb{V}(Y_n) \end{bmatrix}_{n \times n}$$

#### **PROPOSITION 2.1.6: Properties of Covariance Matrix**

Let Y be a random vector and  $a \in \mathbb{R}^n$ , then the covariance matrix has the following properties.

- (1) Symmetric since  $Cov(Y_i, Y_i) = Cov(Y_i, Y_i)$ ; that is  $\mathbb{V}(\mathbf{Y})^{\top} = \mathbb{V}(\mathbf{Y})$ .
- (2) Positive semi-definite since  $a^{\top} \mathbb{V}(Y) a \geqslant 0$  for all  $a \in \mathbb{R}^n$ . (3)  $\mathbb{V}(Y) = \mathbb{E} \left[ (Y \mathbb{E}[Y])(Y \mathbb{E}[Y])^{\top} \right]$

#### **Proof of: 2.1.6**

Trivial.

#### PROPOSITION 2.1.7: Properties of Random Vector

Let a be a  $1 \times n$  matrix (row vector) of constants and A be an  $n \times n$  matrix of constants, then the random vector has the following properties.

- (1)  $\mathbb{E}[aY] = a\mathbb{E}[Y]$
- (2)  $\mathbb{E}[AY] = A\mathbb{E}[Y]$
- (3)  $V(\boldsymbol{a}\boldsymbol{Y}) = \boldsymbol{a}V(\boldsymbol{Y})\boldsymbol{a}^{\top}$
- (4)  $\mathbb{V}(A\mathbf{Y}) = A\mathbb{V}(\mathbf{Y})A^{\top}$

#### **Proof of: 2.1.7**

We prove property (4) only.

$$\begin{split} \mathbb{V}(A\boldsymbol{Y}) &= \mathbb{E}\Big[\big(A\boldsymbol{Y} - \mathbb{E}[A\boldsymbol{Y}]\big)\big(A\boldsymbol{Y} - \mathbb{E}[A\boldsymbol{Y}]\big)^{\top}\Big] \\ &= \mathbb{E}\Big[\big(A\boldsymbol{Y} - A\mathbb{E}[\boldsymbol{Y}]\big)\big(A\boldsymbol{Y} - A\mathbb{E}[\boldsymbol{Y}]\big)^{\top}\Big] \\ &= \mathbb{E}\Big[A(\boldsymbol{Y} - \mathbb{E}[\boldsymbol{Y}])\big(A(\boldsymbol{Y} - \mathbb{E}[\boldsymbol{Y}])\big)^{\top}\Big] \\ &= \mathbb{E}\Big[A(\boldsymbol{Y} - \mathbb{E}[\boldsymbol{Y}])\big(\boldsymbol{Y} - \mathbb{E}[\boldsymbol{Y}]\big)^{\top}A^{\top}\Big] \\ &= A\mathbb{E}\Big[(\boldsymbol{Y} - \mathbb{E}[\boldsymbol{Y}])\big(\boldsymbol{Y} - \mathbb{E}[\boldsymbol{Y}]\big)^{\top}\Big]A^{\top} \\ &= A\mathbb{V}(\boldsymbol{Y})A^{\top} \end{split}$$

#### **EXAMPLE 2.1.8: Calculations with MLR Varaibles**

Let 
$$Y = (Y_1, Y_2, Y_3)^{\top}$$
. Suppose  $\mathbb{E}[Y] = (3, 1, 2)^{\top}$ . Let  $\mathbb{V}(Y) = \begin{bmatrix} 4 & 1/2 & -2 \\ 1/2 & 1 & 0 \\ -2 & 0 & 3 \end{bmatrix}$  and  $\mathbf{a} = (1, -1, 2)$ 

and  $A = \begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}$ . Note that  $\boldsymbol{a}$  is a  $1 \times 3$  row vector. Compute the following.

- (i)  $\mathbb{E}[aY]$
- (ii)  $\mathbb{V}(aY)$
- (iii)  $\mathbb{E}[AY]$
- (iv)  $\mathbb{V}(A\mathbf{Y})$

**Solution.** We do the first two and leave the rest as an exercise.

(i) 
$$\mathbb{E}[aY] = a\mathbb{E}[Y] = \begin{bmatrix} 1 & -1 & 2 \end{bmatrix} \begin{bmatrix} 3 \\ 1 \\ 2 \end{bmatrix} = 1(3) - 1(1) + 2(2) = 6.$$

(ii)

$$\begin{split} \mathbb{V}(\boldsymbol{a}\boldsymbol{Y}) &= \boldsymbol{a}\mathbb{V}(\boldsymbol{Y})\boldsymbol{a}^{\top} \\ &= \begin{bmatrix} 1 & -1 & 2 \end{bmatrix} \begin{bmatrix} 4 & 1/2 & -2 \\ 1/2 & 1 & 0 \\ -2 & 0 & 3 \end{bmatrix} \begin{bmatrix} 1 \\ -1 \\ 2 \end{bmatrix} \\ &= \begin{bmatrix} 1 & -1 & 2 \end{bmatrix} \begin{bmatrix} 4(1) + (1/2)(-1) - 2(2) \\ (1/2)(1) + 1(-1) + 0(2) \\ -2(1) + 0(-1) + 3(2) \end{bmatrix} \\ &= \begin{bmatrix} 1 & -1 & 2 \end{bmatrix} \begin{bmatrix} -1/2 \\ -1/2 \\ 4 \end{bmatrix} \\ &= 1(-1/2) - 1(-1/2) + 2(4) \\ &= 8 \end{split}$$

## 2.2 Multivariate Normal Distribution

#### **DEFINITION 2.2.1: Multivariate normal distribution**

Let  $Y = (Y_1, \dots, Y_n)^{\top}$  be a random vector. We say that  $Y \sim \text{MVN}(\mu, \sigma)$ ; that is, Y follows a **multivariate normal distribution** (MVN) when

$$f(\boldsymbol{y};\boldsymbol{\mu},\boldsymbol{\varSigma}) = \frac{1}{(2\pi)^{n/2}|\boldsymbol{\varSigma}|^{1/2}} \mathrm{exp} \bigg\{ -\frac{1}{2} (\boldsymbol{y} - \boldsymbol{\mu})^{\top} \boldsymbol{\varSigma}^{-1} (\boldsymbol{y} - \boldsymbol{\mu}) \bigg\}$$

where  $\mu$  is defined as the **mean vector**, and  $\Sigma$  is defined as the **covariance matrix**. Note that  $\Sigma^{-1}$  is the inverse of the covariance matrix and  $|\Sigma|$  is the determinant of  $\Sigma$ .

## **THEOREM 2.2.2: Properties of Multivariate Normal Distribution**

Let  $\mathbf{Y} = (Y_1, Y_2, \dots, Y_n)^{\top} \sim MVN(\boldsymbol{\mu}, \boldsymbol{\Sigma}).$ 

(1) Any subset of  $Y_1, Y_2, \dots, Y_n$  also has a multivariate normal distribution and in particular

$$Y_i \sim \mathcal{N}(\mu_i, \Sigma_{ii})$$
  $i = 1, 2, \dots, n$ 

(2) Let  $\mathbf{a} = (a_1, a_2 \dots, a_n)$  be a non-zero row vector  $(1 \times n)$  of constants, then

$$aY \sim \mathcal{N}(a\mu, a\Sigma a^{\top})$$

(3) Let A be an  $n \times n$  matrix of rank n, then

$$AY \sim MVN(A\mu, A\Sigma A^{\top})$$

- (4) The conditional distribution of any subset of  $(Y_1, Y_2, \dots, Y_n)$  given any other coordinates that are not in the subset is a multivariate normal distribution.
- (5)  $Y_i$  and  $Y_j$  are independent random variables if and only if  $\Sigma_{ij} = \text{Cov}(Y_i, Y_j) = 0$ .

### LECTURE 6 | 2020-09-23

## 2.3 Inference

Recall that last lecture, for multiple linear regression, we have  $Y = XB + \varepsilon$  with the assumption that  $\varepsilon \stackrel{\text{iid}}{\sim} \mathcal{N}(0, \sigma^2)$ . Therefore, for a random vector  $\varepsilon$ , we have

$$\boldsymbol{\varepsilon} \sim \text{MVN} \left( \begin{bmatrix} \boldsymbol{0} \\ \boldsymbol{0} \\ \vdots \\ \boldsymbol{0} \\ \boldsymbol{0} \end{bmatrix}, \begin{bmatrix} \boldsymbol{\sigma}^2 & \boldsymbol{0} & \cdots & \boldsymbol{0} & \boldsymbol{0} \\ \boldsymbol{0} & \boldsymbol{\sigma}^2 & \cdots & \boldsymbol{0} & \boldsymbol{0} \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ \boldsymbol{0} & \boldsymbol{0} & \cdots & \boldsymbol{\sigma}^2 & \boldsymbol{0} \\ \boldsymbol{0} & \boldsymbol{0} & \cdots & \boldsymbol{0} & \boldsymbol{\sigma}^2 \end{bmatrix} \right) = \text{MVN}(\boldsymbol{0}_{n \times 1}, \boldsymbol{\sigma}^2 \boldsymbol{I}_{n \times n})$$

since  $Cov(\varepsilon_1, \varepsilon_2) = 0$  due to independence.

Thus,  $Y \sim MVN(XB, \sigma^2 I)$ .

#### **DEFINITION 2.3.1: Least squares for MLR**

We define the least squares for a multiple linear regression model as

$$S(\beta_0,\beta_1,\dots,\beta_p) = \sum_{i=1}^n (y_i - (\underline{\beta_0 + \beta_1 x_{i1} + \dots + \beta_p x_{ip}}))^2$$

#### THEOREM 2.3.2: Least Square Estimates (LSEs) for MLR

Minimizing  $S(\beta_0, \beta_1, \dots, \beta_p)$ , gives the least squares estimate  $\hat{\beta} = (X^T X)^{-1} X^T y$ .

#### Proof of: 2.3.2

The first partial is  $\frac{\partial S}{\partial \beta_0} = \sum_{i=1}^n 2(y_i - \mu_u)(-1)$ , and all other partials for  $j = 1, \dots, p$  are

$$\frac{\partial S}{\partial \boldsymbol{\beta}_j} = \sum_{i=1}^n 2(y_i - \boldsymbol{\mu}_i)(-x_{ij})$$

Set  $\frac{\partial S}{\partial \beta_0}=0$  and  $\frac{\partial S}{\partial \beta_j}=0$  for  $j=1,\dots,p$  to get

$$\begin{cases} \sum_{i=1}^n (y_i - \mu_i) = 0 \iff \mathbf{1}^\top (\boldsymbol{y} - \boldsymbol{\mu}) = 0 \\ \sum_{i=1}^n (y_i - \mu_i) x_{ij} = 0 \iff \boldsymbol{x}_j^\top (\boldsymbol{y} - \boldsymbol{\mu}) = 0 \quad j = 1, \dots, p \end{cases}$$

since we recall that

$$X = \begin{bmatrix} 1 & x_{11} & \cdots & x_{1(p-1)} & x_{1p} \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 1 & x_{n1} & \cdots & x_{n(p-1)} & x_{np} \end{bmatrix} = \begin{bmatrix} \mathbf{1} & \boldsymbol{x}_1 & \cdots & \boldsymbol{x}_{p-1} & \boldsymbol{x}_p \end{bmatrix}$$

Therefore,

$$X^{\top}(\boldsymbol{y} - X\boldsymbol{\beta}) = 0 \iff X^{\top}\boldsymbol{y} - X^{\top}X\boldsymbol{\beta} = 0 \iff X^{\top}X\boldsymbol{\beta} = X^{\top}\boldsymbol{y} \iff \boldsymbol{\beta} = (X^{\top}X)^{-1}X^{\top}\boldsymbol{y}$$

assuming  $X^{\top}X$  is invertible; that is,  $\operatorname{rank}(X^{\top}X) = p+1$ . So, the LS solution for  $\beta$  is given by  $\hat{\beta} = (X^{\top}X)^{-1}X^{\top}y$ .

#### **DEFINITION 2.3.3: Residuals for MLR**

The residuals for a multiple linear regression model is defined as

$$e_i = y_i - (\underline{\hat{\beta}_0 + \hat{\beta}_1 x_{i1} + \cdots \hat{\beta}_p x_{ip}})$$
 fitted value  $\mu_i$ 

or equivalently,  $\hat{\mu} = X\hat{B}$  and  $e = y - \hat{\mu}$ .

The estimate  $\sigma^2$  based on  $e_i$ 's is

$$\hat{\sigma}^2 = \frac{\text{SS}(\text{Res})}{n - (p + 1)} = \frac{\sum_{i=1}^n e_i^2}{n - p - 1} = \frac{e^\top e}{n - p - 1}$$

since d.f. is n – (number of estimated parameters). When viewed as a random variable,

$$\frac{(n-p-1)\hat{\sigma}^2}{\sigma^2} \sim \chi^2(n-p-1)$$

Inference for  $\hat{\pmb{\beta}}=(\hat{\beta}_0,\dots,\hat{\beta}_p)^\top=(X^\top X)^{-1}X^\top \pmb{Y}.$ 

Note that  $\hat{\beta}$  is a matrix of constants and Y is a random vector, and  $Y \sim \text{MVN}(X\beta, \sigma^2 I)$ , so

$$\begin{split} \mathbb{E}[\hat{\boldsymbol{\beta}}] &= \mathbb{E}[(X^{\top}X)^{-1}X^{\top}\boldsymbol{Y}] \\ &= (X^{\top}X)^{-1}X^{\top}\mathbb{E}[\boldsymbol{Y}] \\ &= (X^{\top}X)^{-1}(X^{\top}X)\boldsymbol{\beta} \\ &= \boldsymbol{\beta} \end{split}$$

That is,  $\mathbb{E}[\hat{\beta}_0],\dots,\mathbb{E}[\hat{\beta}_p]=\beta_p$  all unbiased.

$$\begin{split} \mathbb{V}((X^\top X)^{-1}X^\top \boldsymbol{Y}) &= (X^\top X)^{-1}X^\top \mathbb{V}(\boldsymbol{Y}) \big[ (X^\top X)^{-1}X^\top \big]^\top \\ &= (X^\top X)^{-1}X^\top \sigma^2 I(X^\top)^\top \big[ (X^\top X)^{-1} \big]^\top \\ &= \sigma^2 (X^\top X)^{-1} (X^\top X) (X^\top X)^{-1} \end{split} \qquad X^\top X \text{ symmetric}$$

Since  $\hat{\beta}$  is a linear transformation of Y we have  $\hat{\beta} \sim \text{MVN}(\beta, \sigma^2 \underbrace{(X^\top X)^{-1})}_V$ . We proved the following theorem.

## THEOREM 2.3.4: Distribution of $\hat{\beta}_i$

The distribution of a given  $\hat{\beta}_i$  is

$$\hat{\beta}_{i} \sim \mathcal{N}(\beta_{i}, \sigma^{2}V_{ij})$$
  $j = 0, 1, \dots, p$ 

$$Z = \frac{\hat{\beta}_j - \beta_j}{\sigma \sqrt{V_{ij}}} \sim \mathcal{N}(0,1) \quad \text{ and } \quad T = \frac{\hat{\beta}_j - \beta_j}{\hat{\sigma} \sqrt{V_{ij}}} \sim t(n-p-1) \qquad j = 0,1,\dots,p$$

## **DEFINITION 2.3.5: Standard error for** $\hat{\beta}_i$

We define the **standard error** of  $\hat{\beta}_i$  as

$$\mathrm{Se}(\hat{\beta}_j) = \hat{\sigma} \sqrt{V_{jj}} \qquad j = 0, 1, \dots, p$$

So, a  $(1-\alpha)$  confidence interval for  $\beta_j$  is

$$\hat{\beta}_i \pm c \operatorname{Se}(\hat{\beta}_i)$$

where c is  $(1-(\alpha/2))$  quantile of t(n-p-1).

To test  $H_0$ :  $\beta_j=0$  vs  $H_A$ :  $\beta_j\neq 0$ , calculate t-statistic  $t=\frac{\beta_j}{\mathrm{Se}(\hat{\beta}_j)}$  reject at level  $\alpha$  if |t|>c and p-value is  $2P(T\geqslant |t|)$  where  $T\sim t(n-p-1)$ .

Interpretation of  $\hat{\beta}$ : fitted linear regression model says  $\widehat{\mathbb{E}[Y]}$  (estimate of the expected response) is

$$\hat{\beta}_0 + \hat{\beta}_1 x_1 + \dots + \hat{\beta}_p x_p$$

- $\hat{eta}_0$  is the estimate of expected response when all explanatory variables are equal to 0.
- $\hat{\beta}_j$  is the estimated change in expected response for a unit increase in  $x_j$  when holding all other explanatory variables constant.

$$\hat{\beta}_0 + \hat{\beta}_1(x_1+1) + \dots + \hat{\beta}_p x_p - (\hat{\beta}_0 + \hat{\beta}_1 x_1 + \dots + \hat{\beta}_p x_p) = \hat{\beta}_1$$

#### **REMARK 2.3.6**

When it's written  $V_{jj}$ , that means the  $j+1^{\text{th}}$  column and  $j+1^{\text{th}}$  row since we start from index 0 for these matrices. Some unfortunate events may have happened on the quiz to me due to this.

## 2.4 Application

#### **EXAMPLE 2.4.1: Rocket MLR**

Let n=12,  $\hat{\pmb{\beta}}=(473.6,16.7,-1.09)^{\top}=(\hat{\beta}_0,\hat{\beta}_1,\hat{\beta}_2)^{\top}$ .

- $x_1$ : nozzle area (1 = L, 0 = S)
- $x_2$ : propellant ratio
- Y: thrust

$$\hat{\sigma} = \sqrt{\frac{\sum\limits_{i=1}^{12} e_i^2}{12 - 1 - 2}} = \sqrt{\frac{\pmb{e}^\top \pmb{e}}{9}} = 2.655$$

Interpretation of  $\hat{\beta}$ :

- $\hat{\beta}_1$  estimated change in expected thrust is 16.7 when changing small to large nozzle while holding other variables (propellant ratio) constant.
- $\hat{\beta}_2$  estimated thrust to decrease by 1.09 on average for a unit increase in propellant ratio while holding other variables (nozzle area) constant.

Given Se( $\hat{\beta}_2$ ) = 0.94, we compute the t-statistic for  $H_0$ :  $\beta_2 = 0$  vs  $H_A$ :  $\beta_2 \neq 0$  which is t = -1.09/0.94 = -1.16.

$$p\text{-value} = 2P(T \geqslant 1.16) = 0.275 \text{ from R where } T \sim t(9)$$

Do not reject  $H_0$  (e.g.,  $\alpha=0.05$ ), therefore propellant ratio does not significantly influence thrust.

## LECTURE 7 | 2020-09-28

## 2.5 Prediction

Recall that  $\mathbf{Y} = X\boldsymbol{\beta} + \boldsymbol{\varepsilon} \sim \text{MVN}(X\boldsymbol{\beta}, \sigma^2 I)$ , and

- Estimates:  $\hat{\boldsymbol{\beta}} = (X^{\top}X)^{-1}X^{\top}\boldsymbol{Y}$
- Fitted values:  $\hat{\mu} = X\hat{\beta}$
- Residuals:  $e = y \hat{\mu}$
- Constants:  $X = \begin{bmatrix} \mathbf{1} & \mathbf{x}_1 & \cdots & \mathbf{x}_p \end{bmatrix}_{n \times (p+1)}$
- Values of responses:  $\boldsymbol{y} = (y_1, y_2, \dots, y_n)^\top \in \mathbb{R}^n$

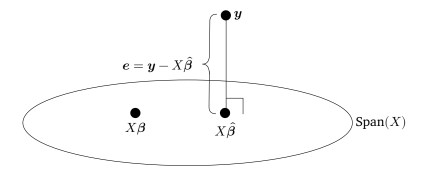
Author's Note: Geometric interpretation of data is omitted in these notes because I'm simply too lazy.

The span of X is  $\mathrm{Span}(X) = \{b_0\mathbf{1} + b_1\mathbf{x}_1 + \dots + b_p\mathbf{x}_p : b_0, \dots, b_p \in \mathbb{R}\} \subset \mathbb{R}^n$  which is all linear combinations of columns of X which is a subspace of  $\mathbb{R}^n$ , and by assumption we know  $\mathrm{rank}(X) = p + 1$ .

We can say  $\mathrm{Span}(X)$  represents all possible vector values  $X \boldsymbol{b}$  where  $\boldsymbol{b} = (b_0, b_1, \dots, b_p)^{\top}$ .

Generally,  $y \notin \operatorname{Span}(X)$ , so since the linear model is an approximation,  $\varepsilon$  variability not explained by model.

Intuitively, it makes sense to choose an estimate  $\hat{\beta}$  so that  $X\hat{\beta}$  is as close to y as possible.



Therefore, e must be orthogonal to Span $(X) \iff e$  is orthogonal to all columns of X.

$$\begin{split} \mathbf{1}^\top (\boldsymbol{y} - \hat{\boldsymbol{\mu}}) &= 0 \\ \boldsymbol{x}_1^\top (\boldsymbol{y} - \hat{\boldsymbol{\mu}}) &= 0 \\ &\vdots \\ \boldsymbol{x}_n^\top (\boldsymbol{y} - \hat{\boldsymbol{\mu}}) &= 0 \end{split}$$

which is the same as LS estimates. We also know  $\hat{\mu} = X\hat{\beta}$  and  $e = y - \hat{\mu}$ .

#### **DEFINITION 2.5.1: Hat matrix**

The **hat matrix** is defined as  $H = X(X^{T}X)^{-1}X^{T}$ .

## **PROPOSITION 2.5.2: Properties of Hat Matrix**

Let H be a hat matrix, then H has the following properties.

- (1) H is symmetric; that is,  $H = H^{\top}$ .
- (2) H is idempotent; that is,  $H^2 = HH = H$ .
- (3) I-H is symmetric idempotent; that is,  $(I-H)^2=(I-H)(I-H)=I-H$ .

#### **Proof of: 2.5.2**

We prove all three because it's easy.

- (1)  $H^{\top} = [X(X^{\top}X)^{-1}X^{\top}]^{\top} = X(X^{\top}X)^{-1}X^{\top} = H.$
- (2)  $HH = X(X^{\top}X)^{-1}(X^{\top}X)(X^{\top}X)^{-1}X^{\top} = H.$
- (3) (I-H)(I-H) = I(I-H) H(I-H) = II IH HI + HH = I 2H + HH = I 2H + H = I H.

Let's view  $\hat{\mu}$  and e as random vectors

$$\begin{split} \hat{\boldsymbol{\mu}} &= X \hat{\boldsymbol{\beta}} = X (X^\top X)^{-1} X^\top \boldsymbol{Y} = H \boldsymbol{Y} \\ \boldsymbol{e} &= \boldsymbol{Y} - \hat{\boldsymbol{\mu}} = I \boldsymbol{Y} - H \boldsymbol{Y} = (I - H) \boldsymbol{Y} \\ \mathbb{E}[\hat{\boldsymbol{\mu}}] &= \mathbb{E}[H \boldsymbol{Y}] = H \mathbb{E}[\boldsymbol{Y}] = X (X^\top X)^{-1} X^\top \underbrace{X \boldsymbol{\beta}}_{\mathbb{E}[\boldsymbol{Y}]} = X \boldsymbol{\beta} \\ \mathbb{V}(\hat{\boldsymbol{\mu}}) &= \mathbb{V}(H \boldsymbol{Y}) = H \mathbb{V}(\boldsymbol{Y}) H^\top = H \sigma^2 I H^\top = \sigma^2 (H H^\top) = \sigma^2 H \\ \mathbb{E}[\boldsymbol{e}] &= \mathbb{E}[(I - H) \boldsymbol{Y}] = \mathbb{E}[\boldsymbol{Y}] - \mathbb{E}[H \boldsymbol{Y}] = X \boldsymbol{\beta} - X \boldsymbol{\beta} = 0 \\ \mathbb{V}(\boldsymbol{e}) &= (I - H) \mathbb{V}(\boldsymbol{Y}) (I - H)^\top = \sigma^2 (I - H) (I - H)^\top = \sigma^2 (I - H) \end{split}$$

So since  $\hat{\mu}$  and e are linear transformations of Y we have proved the following theorem.

#### THEOREM 2.5.3: Distribution of $\hat{\mu}$ and e

 $\hat{\mu}$  and  $\hat{e}$  have the following distribution.

$$\hat{\boldsymbol{\mu}} \sim MVN(X\boldsymbol{\beta}, \sigma^2 H)$$

$$\hat{\boldsymbol{e}} \sim MVN(0, \sigma^2(I-H))$$

Suppose we want to predict response for  $x_0$  where the first 1 represents the intercept in the row vector.

$$\boldsymbol{x}_0 = \begin{bmatrix} 1 & x_{01} & x_{02} & \cdots & x_{0p} \end{bmatrix}_{1 \times (p+1)}$$

Let  $Y_0$  random variable representing the response associated with  $x_0$ . In multiple linear regression,

$$Y_0 \sim \mathcal{N}(\beta_0 + \beta_1 x_{01} + \dots + \beta_p x_{0p}, \sigma^2)$$

So we predict the value

$$\hat{y}_0 = \hat{\beta}_0 + \hat{\beta}_1 x_{01} + \dots + \hat{\beta}_p x_{0p} = \mathbf{x}_0 \hat{\beta}$$

which represents the estimated mean response given  $x_{01}, x_{02}, \dots, x_{0p}$ . Corresponding distribution has

$$\mathbb{E}[\hat{Y}_0] = oldsymbol{x}_0 \mathbb{E}[\hat{oldsymbol{eta}}] = oldsymbol{x}_0 oldsymbol{eta} = \mathbb{E}[Y_0]$$

$$\mathbb{V}(\hat{Y}_0) = \boldsymbol{x}_0 \mathbb{V}(\hat{\boldsymbol{\beta}}) \boldsymbol{x}_0^\top = \boldsymbol{x}_0 \sigma^2 (X^\top X)^{-1} \boldsymbol{x}_0^\top$$

We have proved the following theorem.

#### **THEOREM 2.5.4: Distribution of Predictor**

The distribution of  $\hat{Y}_0$  which is a function of  $Y_1, \dots, Y_n$  is

$$\hat{Y}_0 \sim \mathcal{N}(\boldsymbol{x}_0\boldsymbol{\beta}, \sigma^2\boldsymbol{x}_0(X^\top X)^{-1}\boldsymbol{x}_0^\top)$$

$$\frac{\hat{Y}_0 - \boldsymbol{x}_0 \boldsymbol{\beta}}{\sigma \sqrt{\boldsymbol{x}_0 (X^\top X)^{-1} \boldsymbol{x}_0^\top}} \sim \mathcal{N}(0, 1)$$

$$\frac{\hat{Y}_0 - \boldsymbol{x}_0 \boldsymbol{\beta}}{\hat{\sigma} \sqrt{\boldsymbol{x}_0 (X^\top X)^{-1} \boldsymbol{x}_0^\top}} \sim t(n - (p+1)) = t(n-p-1)$$

A  $(1-\alpha)$  confidence interval for the mean response  $y_0={m x}_0\hat{m \beta}$  given  ${m x}_0$  is

$$\hat{y}_0 \pm c \hat{\sigma} \sqrt{\boldsymbol{x}_0 (X^\top X)^{-1} \boldsymbol{x}_0^\top}$$

where c is the  $1 - \alpha/2$  quantile of t(n - p - 1).

Prediction error:  $Y_0 - \hat{Y}_0$  which are independent since  $Y_0$  is a random variable with variance  $\sigma^2$  and  $\hat{Y}_0$  is a function of  $Y_1, \dots, Y_n$ . Therefore,

$$\begin{split} \mathbb{E}[Y_0 - \hat{Y}_0] &= \boldsymbol{x}_0 \boldsymbol{\beta} - \boldsymbol{x}_0 \boldsymbol{\beta} = 0 \\ \mathbb{V}(Y_0 - \hat{Y}_0) &= \mathbb{V}(Y_0) + (-1)^2 \mathbb{V}(\hat{Y}_0) = \sigma^2 + \sigma^2 (\boldsymbol{x}_0 (\boldsymbol{X}^\top \boldsymbol{X})^{-1} \boldsymbol{x}_0^\top) \end{split}$$

We have proved the following theorem.

## **THEOREM 2.5.5: Distribution of Prediction Error**

The distribution of the prediction error is

$$Y_0 - \hat{Y}_0 \sim N(0, \sigma^2(1 + \pmb{x}_0(X^\top X)^{-1} \pmb{x}_0^\top)).$$

A  $(1-\alpha)$  prediction interval for the mean response  $y_0={m x}_0\hat{m \beta}$  given  ${m x}_0$  is

$$\hat{y}_0 \pm c \hat{\sigma} \sqrt{1 + \boldsymbol{x}_0 (X^\top X)^{-1} \boldsymbol{x}_0^\top}$$

where c is the  $1 - \alpha/2$  quantile of t(n - p - 1).

## **REMARK 2.5.6**

Our intuition tells us that the prediction interval is wider than the confidence interval for mean. In other words, estimating an average is "easier" than an individual response.

## LECTURE 8 | 2020-09-30

## 2.6 Categorical Predictors

## 2.6.1 R Demo

```
## NASA rocket data example

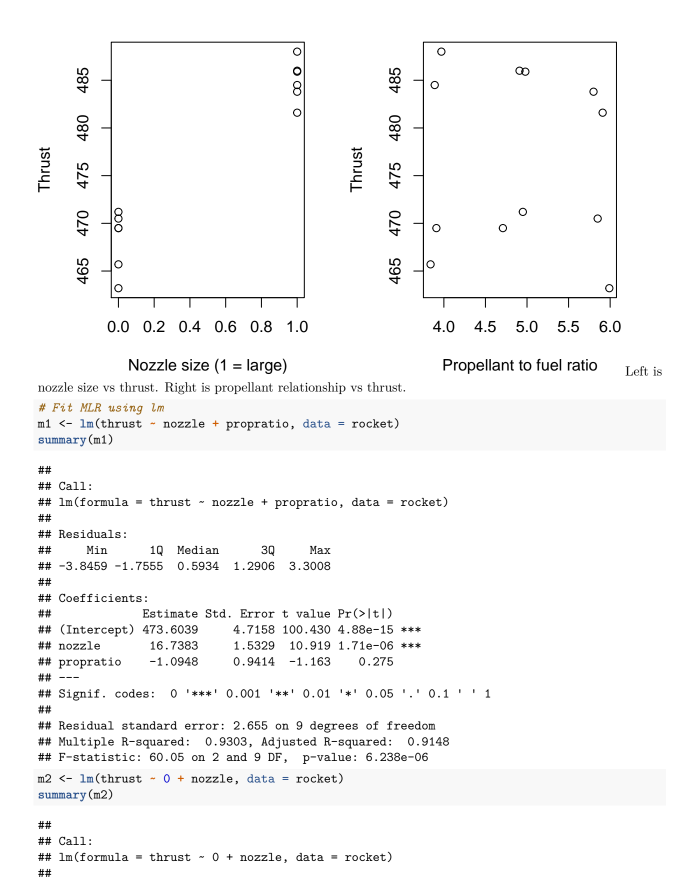
## From: R.S. Jankovsky, T.D. Smith, A.J. Pavli (1999). "High-Area-Ratio Rocket
## Nozzle at High Combustion Chamber Pressure-Experimental and Analytical
## Validation".

# setwd(...) first if your CSV file is somewhere else
rocket <- read.csv("csv/rocket.csv")
# output all data in rocket vector
rocket</pre>
```

```
##
     thrust nozzle propratio
## 1
      488.0
            1
                     3.97
## 2
     481.6
              1
                     5.91
## 3
     485.9
              1
                     4.98
## 4
     486.0
               1
                     4.91
## 5
     484.5
              1
                     3.89
## 6 483.8
                     5.80
              1
## 7
     463.2
              0
                     5.99
## 8
     471.2
               0
                     4.95
## 9
      469.5
               0
                     3.91
## 10 470.5
               0
                     5.85
## 11 469.5
               0
                     4.71
## 12 465.7
               0
                     3.84
```

Y (thrust) is the response variable, and there are two explanatory variables  $x_1, x_2$  (nozzle, propratio) where nozzle is coded as 1 if it's large.

```
# Scatter plots where mfrow is used to put multiple plots on one image
par(mfrow = c(1, 2))
plot(rocket$nozzle,
    rocket$thrust,
    ylab = "Thrust",
    xlab = "Nozzle size (1 = large)")
plot(rocket$propratio,
    rocket$thrust,
    ylab = "Thrust",
    xlab = "Propellant to fuel ratio")
```



## Residuals:

```
##
      Min
               1Q Median
                              3Q
                                    Max
   -3.37
            0.58 233.12 469.50 471.20
##
##
## Coefficients:
##
          Estimate Std. Error t value Pr(>|t|)
             485.0
                         141.2 3.435 0.00558 **
## nozzle
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 345.8 on 11 degrees of freedom
## Multiple R-squared: 0.5175, Adjusted R-squared: 0.4736
## F-statistic: 11.8 on 1 and 11 DF, p-value: 0.005575
anova(m1)
## Analysis of Variance Table
##
## Response: thrust
              Df Sum Sq Mean Sq F value
##
                                              Pr(>F)
## nozzle
               1 836.67 836.67 118.7377 1.743e-06 ***
## propratio 1
                   9.53
                            9.53
                                    1.3524
                                              0.2748
                            7.05
## Residuals 9 63.42
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
On the left it's Y (response variable) and on the right it's x_1, x_2 (explanatory variables). From summary, we
get the estimate vector \hat{\beta} = (473.6039, 16.7383, -1.0948)^{\top}.
# Manual beta estimates where rep is used to make the columns of 1s
X <- cbind(rep(1, 12), rocket$nozzle, rocket$propratio) # X matrix</pre>
y <- matrix(rocket$thrust, ncol = 1) # response vector
beta_hat <- solve(t(X) %*% X) %*% t(X) %*% y
beta_hat
##
               [,1]
## [1,] 473.603924
## [2,] 16.738319
## [3,]
         -1.094822
solve is used for the inverse. ** is used for matrix-matrix multiplication, and t(X) is used for transposing
# Manual sigma estimate
mu_hat <- X %*% beta_hat # fitted values</pre>
e <- y - mu_hat # residuals
sigma_hat <- sqrt((t(e) %*% e) / 9) # Note n-p-1 = 12-2-1 = 9
sigma_hat
           [,1]
## [1,] 2.6545
sigma_hat <- sqrt(sum(e ^ 2) / 9) # equivalent
sigma_hat
## [1] 2.6545
  • \hat{\boldsymbol{\mu}} = X\hat{\boldsymbol{\beta}}
  • e = y - \hat{\mu}
```

```
• \hat{\sigma} = \sqrt{\left(\sum_{i=1}^{n} e_i^2\right)/9} = 2.6545, or
   • \hat{\sigma} = \sqrt{(e^{\top}e)/9} = 2.6545
# Covariance matrix of beta_hat
vcov(m1)
                 (Intercept)
                                    nozzle
                                             propratio
## (Intercept) 22.238325 -1.02316688 -4.32080608
## nozzle
                   -1.023167 2.34987593 -0.03102117
                   -4.320806 -0.03102117 0.88631920
## propratio
sqrt(diag(vcov(m1))) # SEs of individual betas
## (Intercept)
                                 propratio
                       nozzle
     4.7157528
                   1.5329305
                                 0.9414453
# Manual
se_beta <- sigma_hat * sqrt(diag(solve(t(X) %*% X)))</pre>
se_beta
## [1] 4.7157528 1.5329305 0.9414453
   • Se(\hat{\beta}) = \hat{\sigma}\sqrt{(X^{\top}X)^{-1}} = (4.71, 1.53, 0.94)^{\top}
# Estimate the mean response for units with small nozzle and propellant ratio 5.5
# include a 95% CI
predict(
  object = m1,
  newdata = data.frame(nozzle = 0, propratio = 5.5),
  interval = "confidence",
  level = 0.95
)
##
           fit
## 1 467.5824 464.7929 470.3719
Therefore, \hat{y}_0 = 467.58. The 95% confidence interval for the mean response given x_0 is [464.7929, 470.3719].
# Manual calculation
x0 \leftarrow matrix(c(1, 0, 5.5), nrow = 1)
y0 hat <- x0 %*% beta hat
y0_hat
              [,1]
## [1,] 467.5824
# mu0 is also known as \hat{Y}_0
se_mu0 <- sigma_hat * sqrt(x0 %*% solve(t(X) %*% X) %*% t(x0))</pre>
se mu0
##
              [,1]
## [1,] 1.233132
crit val <-qt(0.975, 9)
ci_lo <- y0_hat - crit_val * se_mu0
ci_hi <- y0_hat + crit_val * se_mu0</pre>
c(y0_hat, ci_lo, ci_hi)
```

## [1] 467.5824 464.7929 470.3719

```
• x_0 = \begin{bmatrix} 1 & 0 & 5.5 \end{bmatrix}
   • \hat{y}_0 = x_0 \hat{\beta} = 467.5824
   • Se(\hat{Y}_0) = \hat{\sigma} \sqrt{x_0(X^\top X)^{-1} x_0^\top} = 1.233132
Therefore, \hat{y}_0 = 467.58. The 95% confidence interval for the mean response given x_0 is [464.7929, 470.3719].
# Predict the value of the response for a unit with small nozzle and propellant ratio 5.5
# include a 95% PI
predict(
  object = m1,
  newdata = data.frame(nozzle = 0, propratio = 5.5),
  interval = "prediction",
  level = 0.95
)
##
            fit
                      lwr
                                 upr
## 1 467.5824 460.9612 474.2036
Therefore, y_0 = 467.5824. The 95% prediction interval for the response (y_0) given x_0 is [460.9612474.2036].
# Manual calculation for an individual
x0 \leftarrow matrix(c(1, 0, 5.5), nrow = 1)
y0_hat <- x0 %*% beta_hat
se_y0 <- sigma_hat * sqrt(1 + x0 %*% solve(t(X) %*% X) %*% t(x0))
se_y0
##
              [,1]
## [1,] 2.926941
crit_val <- qt(0.975, 9)
pi_lo <- y0_hat - crit_val * se_y0</pre>
pi_hi <- y0_hat + crit_val * se_y0</pre>
c(y0_hat, pi_lo, pi_hi)
## [1] 467.5824 460.9612 474.2036
```

•  $Se(Y_0 - \hat{Y}_0) = \hat{\sigma}\sqrt{1 + x_0(X^\top X)^{-1}x_0^\top} = 2.926941$ 

Handling categorical variables: when there are explanatory variables with values that fall into one of several categories.

- e.g., nozzle large/small, if just binary, code as 1 and 0
- ordered small, medium, large or not red, blue, green

Approach: can convert to indicator variables or treat as numerical if it makes sense to do so.

Example: Coffee Quality Institute (2018)

Extract a few variables:

	Acidity	Method
1	8.7	Washed-wet
2	8.3	Washed-wet
3	8.2	Natural-dry
4	8.4	Semi-washed/pulped

Flavour (response)

How to set up X? For example,

$$x_{i2} = \begin{cases} 0 & \text{dry} \\ 1 & \text{semi} \\ 2 & \text{wet} \end{cases}$$

Not generally appropriate unless we think a response is linear according to this scheme.

More flexible approach: indicator/dummy variables

$$x_{i2} = \begin{cases} 1 & \text{semi} \\ 0 & \text{otherwise} \end{cases}, \quad x_{i3} = \begin{cases} 1 & \text{wet} \\ 0 & \text{otherwise} \end{cases}$$

Therefore,

$$X = \begin{bmatrix} 1 & 8.7 & 0 & 1 \\ 1 & 8.3 & 0 & 1 \\ 1 & 8.2 & 0 & 0 \\ 1 & 8.4 & 1 & 0 \end{bmatrix}$$

Why not  $x_{i4} = \begin{cases} 1 & \text{dry} \\ 0 & \text{otherwise} \end{cases}$  ? If we did that, we would have

$$X = \begin{bmatrix} 1 & 8.7 & 0 & 1 & 0 \\ 1 & 8.3 & 0 & 1 & 0 \\ 1 & 8.2 & 0 & 0 & 1 \\ 1 & 8.4 & 1 & 0 & 0 \end{bmatrix}$$

This has linearly dependent columns since  $x_4 = 1 - x_2 - x_3$ . There is no new information and X would not have full rank.

Model: 
$$Y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$$
.

Interpretation:

- Mean flavour if acidity =  $x_{01}$  and method dry is  $\beta_0 + \beta_1 x_{01}$ .
- Mean flavour if acidity =  $x_{01}$  and method wet is  $\beta_0 + \beta_1 x_{01} + \beta_3$ .
- Mean flavour if acidity =  $x_{01}$  and method semi is  $\beta_0 + \beta_1 x_{01} + \beta_2$ .

- $\beta_2$  is the difference between semi and dry in expected response (holding acidity constant)
- $\beta_3$  is the difference between wet and dry in expected response (holding acidity constant)
- $\beta_2 \beta_3$  is the difference between semi and wet (holding other variables constant)

 $\hat{\boldsymbol{\beta}} \sim \text{MVN}(\boldsymbol{\beta}, \sigma^2 V) \text{ where } V = (X^{\top} X)^{-1}.$ 

- We know  $\hat{\beta}_j \sim \mathcal{N}(\beta_j, \sigma^2 V_{jj})$  with  $\mathrm{Se}(\hat{\beta}_j) = \hat{\sigma} \sqrt{V_{jj}}$  where  $j=0,\dots,p$ .
- What about  $\beta_2 \beta_3$ ?

$$\mathbb{V}(\hat{\beta}_2-\hat{\beta}_3)=\mathbb{V}(\hat{\beta}_2)-\mathbb{V}(\hat{\beta}_3)-2\mathrm{Cov}(\hat{\beta}_2,\hat{\beta}_3)=\sigma^2V_{22}+\sigma^2V_{33}-2\sigma^2V_{23}$$

Therefore,

$$Se(\hat{\beta}_2 - \hat{\beta}_3) = \hat{\sigma}\sqrt{V_{22} + V_{33} - 2V_{23}}$$

Now, we can construct a CI for  $\beta_2 - \beta_3$ .

In general, for an explanatory variable with k categories. We need k-1 indicator variables.

## LECTURE 9 | 2020-10-05

## 2.7 Analysis of Variance

Analysis of variance (ANOVA): how well does our regression model fit our response variable? Variability in response can be measured by "total sum of squares:"

$$\mathrm{SS}(\mathrm{Total}) = \sum_{i=1}^n (y_i - \bar{y})^2$$

as seen in A1, it's closely related to sample variance of  $y_1,\ldots,y_n$ , which is  $\mathrm{SS}(\mathrm{Total})/(n-1)$ .

ANOVA decomposes SS(Total) = SS(Reg) + SS(Res) where SS(Reg) is the regression sum of squares and SS(Res) is the residual sum of squares.

The regression sum of squares is variation explained by the model and the residual sum of squares is the variation not explained by the regression model.

Using the fact that

$$y_i - \bar{y} = y_i - \hat{\mu}_i + \hat{\mu}_i - \bar{y}$$

When regression fits data well, the observations  $y_i$  tend to be much closer to  $\hat{\mu}_i$ . Note that  $\bar{y}$  is line a regression line with  $\beta_1 = 0$ .

Mathematically,

since

$$\underbrace{\sum_{i=1}^n (y_i - \bar{y})^2}_{\text{SS(Total)}} = \underbrace{\sum_{i=1}^n (\hat{\mu}_i - \bar{y})^2}_{\text{SS(Reg)}} + \underbrace{\sum_{i=1}^n (y_i - \hat{\mu}_i)^2}_{\text{SS(Res)}}$$

since we showed that  $\sum\limits_{i=1}^n (\hat{\mu}_i - \bar{y}) \underbrace{(y_i - \hat{\mu}_i)}_{e_i} = 0$  in A1 for SLR. It's also true for multiple linear regression

$$\sum_{i=1}^n (\hat{\mu}_i - \bar{y}) e_i = \sum_{i=1}^n (e_i \hat{\mu}_i) - \bar{y} \sum_{i=1}^n e_i = \hat{\boldsymbol{\mu}}^\top \boldsymbol{e} - \bar{y} \mathbf{1}^\top \boldsymbol{e} = 0$$

Recall:  $\mathbf{1}^{\top}e=0$  is one of LS equations, and  $\hat{\boldsymbol{\mu}}=X\hat{\boldsymbol{\beta}}$  is in  $\mathrm{Span}(X)$ , so e is orthogonal to  $\mathrm{Span}(X)$ , so  $\hat{\boldsymbol{\mu}}^{\top}e=0$ .

*F* is used to test the overall significance of regression (later).

Table 2.1: ANOVA Table

Source	d.f.	SS	Mean Square	F
Regression Residual	$p \\ n-p-1$	$\begin{array}{c} {\rm SS(Reg)} \\ {\rm SS(Res)} \end{array}$	$\frac{\mathrm{SS}(\mathrm{Reg})/p}{\mathrm{SS}(\mathrm{Res})/(n\!-\!p\!-\!1)} = \hat{\sigma}^2$	MS(Reg)/MS(Res)
Total	n-1	SS(Total)		

## 2.8 Coefficient of Determination

#### **DEFINITION 2.8.1: Coefficient of Determination**

We define the coefficient of determination as

$$R^{2} = \frac{SS(Reg)}{SS(Total)} = 1 - \frac{SS(Res)}{SS(Total)}$$

Note that  $0 \le R^2 \le 1$ .

 $R^2$  is the proportion of variation (in our response variable) that is explained by the regression model. Larger  $R^2$  means the fitted values are closer to the observations  $y_i$ , which means the residuals are small; that is, smaller SS(Res). Note that (A1) in SLR,  $R^2$  is equivalent to the square of the sample correlation between x and y based on  $(x_1, y_1), \ldots, (x_n, y_n)$ .

Table 2.2: Rocket ANOVA Table

TABLE E.E. TO CACCULATE ABLE				
Source	d.f.	$\overline{SS}$	Mean Square	$\overline{F}$
Regression Residual	2 9	846.2 63.42	$423.1 \\ 7.05$	60
Total	11	909.62		

Response thrust  $R^2 = 846.2/909.62 \approx 0.93$ .  $R^2$  interpretation: regression model with nozzle size and propellant ratio explains 93% of variation in thrust (response).

LECTURE 10 | 2020-10-07

## 2.9 General Linear Hypothesis Tests Based on F Distribution

So far we've tested  $H_0$ :  $\beta_i = 0$  vs  $H_A$ :  $\beta_i \neq 0$  involving individual parameters, using t distribution.

Now consider hypothesis test of the form  $H_0$ :  $A\beta = \mathbf{0}$  where A is a matrix of constraints specifying linear combinations of parameters.

#### **EXAMPLE 2.9.1: Coffee Continued**

The full model is:

$$Y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$$

- $Y_i$  is the flavour
- $x_{i1}$  is acidity
- $x_{i2}$  is 1 if semi, and 0 otherwise.
- $x_{i3}$  is 1 if wet, and 0 otherwise.

Example 1.

- $H_0$ :  $\beta_1 = \beta_2 = \beta_3 = 0$  versus
- $H_A$ : at least one of  $\beta_1,\beta_2,\beta_3$  not 0. If  $H_0$  is true, the model reduces to  $Y_i=\beta_0+\varepsilon_i$ .
- This tests overall significance of regression (whether any of predictors impact response)
- $A = \begin{bmatrix} 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix}$ . Note that row i considers the constraint of  $\beta_i = 0$  for i = 1, 2, 3 in this

example.

#### Example 2.

- $H_0$ :  $\beta_2=\beta_3=0$  If  $H_0$  is true,  $Y_i=\beta_0+\beta_1x_{i1}+\varepsilon_i$  Q: Is reduced model with only acidity plausible?
- $A=\begin{bmatrix}0&0&1&0\\0&0&0&1\end{bmatrix}$ . Note that  $A\beta=\mathbf{0}_{1\times 2}$  Example 3.

- $H_0$ :  $\beta_2 \beta_3 = 0$
- $H_A$ :  $\beta_2 \neq \beta_3$
- Y<sub>i</sub> = β<sub>0</sub> + β<sub>1</sub>x<sub>i1</sub> + β<sub>2</sub>(x<sub>i2</sub> + x<sub>i3</sub>) + ε<sub>i</sub> where (x<sub>i2</sub> + x<sub>i3</sub>) is 1 if semi/wet and 0 if dry.
  Do the wet and semi methods have the same impact on the response (holding acidity constant)?
- $A = [0 \ 0 \ 1 \ -1]$

In general, with  $\ell$  constraints. A is an  $\ell \times (p+1)$  matrix with rank  $\ell$ . Recall that

$$\mathrm{Span}(X) = \{\beta_0 \mathbf{1} + \beta_1 \boldsymbol{x}_1 + \dots + \beta_p \boldsymbol{x}_p\}$$

Let

$$\mathrm{Span}(X)_A = \{\beta_0 \mathbf{1} + \beta_1 \boldsymbol{x}_1 + \dots + \beta_p \boldsymbol{x}_p : A\boldsymbol{\beta} = \mathbf{0}\}$$

which is a subspace of Span(X) since any vector in  $Span(X)_A$  is also in Span(X). We call  $Span(X)_A$  the Span(X) with constraint A on  $\beta$ .

Let  $\hat{\mu}_A$  denote the fitted values from fitting the reduced model. The residual if we fit the model with  $A\beta = 0$ is  $e_A = y - \hat{\mu}_A$ .

If  $H_0$ :  $A\beta = 0$  is true, then  $\hat{\mu}$  and  $\hat{\mu}_A$  should be close; that is, the model makes similar predictions whether we set  $A\beta = \mathbf{0}$  or not when fitting the model.

So to assess whether  $H_0$  is plausible, look at  $\|\hat{\mu} - \hat{\mu}_A\|$  where  $\|\cdot\|$  is Euclidean or  $L_2$  norm. That is,

$$\|\hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A\| = \sqrt{(\hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A)^\top (\hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A)}$$

If it's "large" or "small" (close to 0) where large gives evidence against  $H_0$  and small gives evidence for  $H_0$ .

By Pythagoras,

$$\| \boldsymbol{y} - \hat{\boldsymbol{\mu}}_A \|^2 = \| \boldsymbol{y} - \hat{\boldsymbol{\mu}} \|^2 + \| \hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A \|^2 \quad \text{ or } \quad \| \boldsymbol{e}_A \|^2 = \| \boldsymbol{e} \|^2 + \| \hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A \|^2$$

or equivalently  $e_A^\top e_A = e^\top e + \|\hat{\mu} - \hat{\mu}_A\|^2$  where  $e_A^\top e_A$  is the sum of squares residual in the reduced model and  $e^\top e$  is the sum of squares residual in the full model.

We define  $e_A^{\top} e_A = SS(Res)_A$  and  $e^{\top} e = SS(Res)$ .

Thus,  $\|\hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A\|^2 = \mathrm{SS}(\mathrm{Res})_A - \mathrm{SS}(\mathrm{Res}) \geqslant 0$  additional sum of squares explained by full model vs reduced one with constraints A.

Practical implications:

• SS(Res) cannot decrease when constraints applied.

ullet Equivalently, full model always has small (or equal) SS(Res) for a fixed SS(Tot) and thus higher  $R^2$  compared to a reduced model.

Define test statistic:

$$F = \frac{(\mathsf{SS}(\mathsf{Res})_A - \mathsf{SS}(\mathsf{Res}))/\ell}{\mathsf{SS}(\mathsf{Res})/(n-p-1)} = \frac{(\mathsf{SS}(\mathsf{Res})_A - \mathsf{SS}(\mathsf{Res}))/\ell}{\hat{\sigma}^2}$$

#### **DEFINITION 2.9.2:** F distribtion

If  $U \sim \chi^2(a)$  and  $V \sim \chi^2(b)$  are independent. We say F follows an F distribution if

$$F = \frac{U/a}{V/b}$$

and write  $F \sim F(a, b)$ .

Here, we have these facts when  $H_0$  is true

$$\begin{split} V &= \frac{\hat{\sigma}^2(n-p-1)}{\sigma^2} \sim \chi^2(n-p-1) \\ U &= \frac{\|\hat{\pmb{\mu}} - \hat{\pmb{\mu}}_A\|^2}{\sigma^2} \sim \chi^2(\ell) \end{split}$$

where U and V are independent. Therefore,

$$F = \frac{\frac{\|\hat{\boldsymbol{\mu}} - \hat{\boldsymbol{\mu}}_A\|^2}{\sigma^2} \bigg(\frac{1}{\ell}\bigg)}{\frac{\hat{\sigma}^2(n-p-1)}{\sigma^2} \bigg(\frac{1}{n-p-1}\bigg)} \sim F(\ell, n-p-1)$$

when  $H_0$  is true. Reject  $H_0$ :  $A\beta = \mathbf{0}$  at level  $\alpha$  if F is greater than  $(1 - \alpha)$  quantile of  $F(\ell, n - p - 1)$  and p-value is  $P(Y \geqslant F)$  where  $Y \sim F(\ell, n - p - 1)$ .

Relation to T distribution: Say  $Y \sim t(a)$ 

$$Y = \frac{Z}{\sqrt{U/a}}$$

where  $Z \sim \mathcal{N}(0,1)$  and  $U \sim \chi^2(a)$  are independent. Squaring everything

$$Y^2 = \frac{Z^2}{U/a}$$

and we know  $Z^2 \sim \chi^2(1)$ . Therefore,  $Y^2 \sim F(1,a)$  (we divide by 1 in the numerator).

Thus, if our hypothesis test has one constraint, then F test is equal to t test of same hypothesis; for example,  $H_0$ :  $\beta_1 = 0$  versus  $H_A$ :  $\beta_1 \neq 0$ .

## LECTURE 11 | 2020-10-19

## 2.10 ANOVA F Test

Recall the general linear hypothesis:  $H_0$ :  $A\beta = 0$  vs  $H_A$ :  $A\beta \neq 0$  where A gives  $\ell$  constraints.

$$F \text{ statistic} = \frac{(\text{SS}(\text{Res})_A - \text{SS}(\text{Res}))/\ell}{\text{SS}(\text{Res})/(n-p-1)} = \frac{(\text{SS}(\text{Res})_A - \text{SS}(\text{Res}))/\ell}{\hat{\sigma}^2}$$

compare to  $F(\ell, n-p-1)$ .

Special case: overall test of significance

"Are any predictors related to response?"

- $H_0$ :  $\beta_1 = \beta_2 = \cdots = \beta_p = 0$
- $H_A$ :  $\beta_j \neq 0$  for at least one j

$$A = \begin{bmatrix} 0 & 1 & 0 & 0 & \cdots & 0 & 0 \\ 0 & 0 & 1 & 0 & \cdots & 0 & 0 \\ \vdots & & & \ddots & & & \\ 0 & 0 & 0 & 0 & \cdots & 1 & 0 \\ 0 & 0 & 0 & 0 & \cdots & 0 & 1 \end{bmatrix} \begin{bmatrix} \beta_0 \\ \beta_1 \\ \vdots \\ \beta_{p-1} \\ \beta_p \end{bmatrix}$$

If  $H_0$  is true:  $Y_i = \beta_0 + \varepsilon_i$  where  $Y_i \sim \mathcal{N}(\beta_0, \sigma^2)$ .

Fit reduced model; that is, in this case estimate  $\beta_0$  using the least squares, minimize  $\sum_{i=1}^n (y_i - \beta_0)^2$ , which can be shown  $\hat{\beta}_0 = \bar{y}$ . So,

$$\mathrm{SS}(\mathrm{Res})_A = \sum_{i=1}^n e_i^2 = \sum_{i=1}^n (y_i - \hat{\mu}_i)^2 = \sum_{i=1}^n (y_i - \bar{y})^2 = \mathrm{SS}(\mathrm{Total})$$

Then,

$$F = \frac{(\mathsf{SS}(\mathsf{Total}) - \mathsf{SS}(\mathsf{Res}))/p}{\mathsf{SS}(\mathsf{Res})/(n-p-1)} = \frac{\mathsf{SS}(\mathsf{Reg})/p}{\mathsf{SS}(\mathsf{Res})/(n-p-1)} = \frac{\mathsf{MS}(\mathsf{Reg})}{\mathsf{MS}(\mathsf{Res})} \leftarrow F \text{ statistic on ANOVA table}$$

#### 2.10.1 R Demo

```
## R demo for Oct 19
## Plotting functions and histograms, F distribution,
## ANOVA tables, F tests, MLR with categorical variables
```

Evaluate the function at many x values, then plot it.

```
# Plotting functions (e.g., probability density functions)
# Create sequence from -4 to 4 increasing 0.01 each time.
x <- seq(-4, 4, 0.01)
head(x)</pre>
```

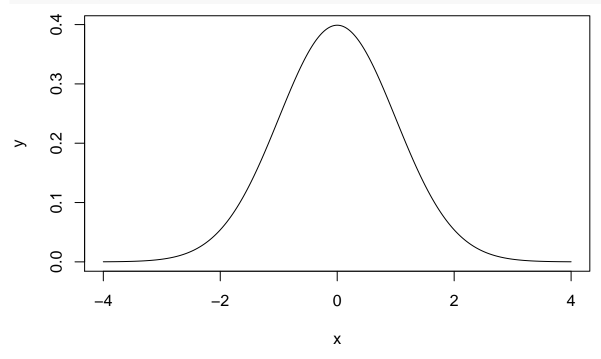
```
## [1] -4.00 -3.99 -3.98 -3.97 -3.96 -3.95

# Normal probability density function with mean 0, and standard deviation 1.

y <- dnorm(x, 0, 1)
```

dnorm is for density normal.

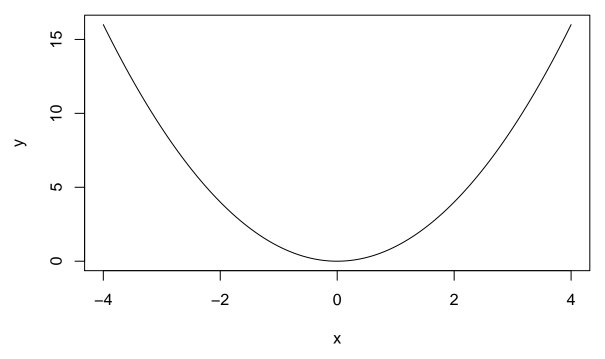
```
plot(x, y, type = "l")
```



type = "1" is for a smooth line (instead of dots).

We can also plot  $y = x^2$  for example.

```
y <- x ^ 2
plot(x, y, type = "1")
```

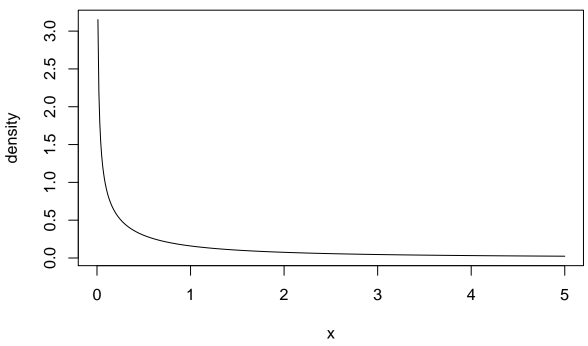


#### F-distribution Examples

```
x \leftarrow seq(0,5,0.01)
head(x)
```

```
## [1] 0.00 0.01 0.02 0.03 0.04 0.05
```

```
# df is degrees of freedom.
# type = "l" is for a smooth curve
plot(
    x,
    y = df(x, df1 = 1, df2 = 1),
    type = "l",
    xlab = "x",
    ylab = "density"
)
```



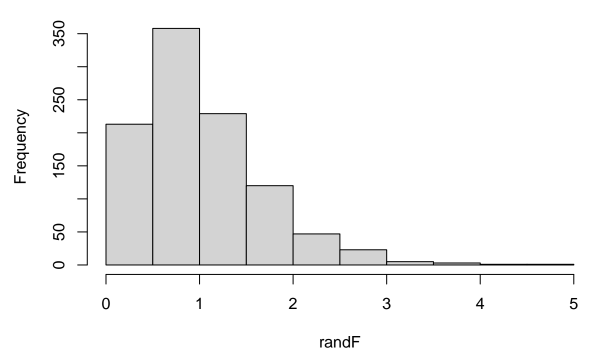
```
# ylim is for the y-axis limits
# lwd is for line width
plot(
  x,
 y = df(x, df1 = 1, df2 = 1),
 type = "1",
 col = "black",
 xlab = "x",
 ylab = "density",
 ylim = c(0, 2.5),
 lwd = 2
# Add lines to the existing plot.
lines(
 х,
 y = df(x, df1 = 1, df2 = 100),
 type = "1",
 col = "green",
 lwd = 2
)
lines(
 y = df(x, df1 = 5, df2 = 1),
 type = "1",
 col = "blue",
 lwd = 2
)
lines(
 y = df(x, df1 = 5, df2 = 100),
 type = "1",
col = "purple",
```

```
lwd = 2
)
lines(
  y = df(x, df1 = 10, df2 = 1),
  type = "1",
  col = "red",
  lwd = 2
)
lines(
  х,
  y = df(x, df1 = 10, df2 = 100),
  type = "1",
  col = "orange",
  lwd = 2
)
# Add a legend to the top-right.
# lty = 1 is for a straight solid line.
legend(
  "topright",
  legend = c(
    "df1=1, df2=1",
    "df1=1, df2=100",
    "df1=5, df2=1",
    "df1=5, df2=100",
    "df1=10, df2=1",
    "df1=10, df2=100"
  ),
  lty = 1,
  col = c("black", "green", "blue", "purple", "red", "orange")
     2.5
                                                                - df1=1, df2=1
                                                                   df1=1, df2=100
     2.0
                                                                   df1=5, df2=1
                                                                   df1=5, df2=100
     1.5
                                                                   df1=10, df2=1
density
                                                                   df1=10, df2=100
     1.0
     0.5
     0.0
             0
                           1
                                        2
                                                      3
                                                                                  5
                                                                    4
                                               Χ
```

#### Random numbers for the F-distribution

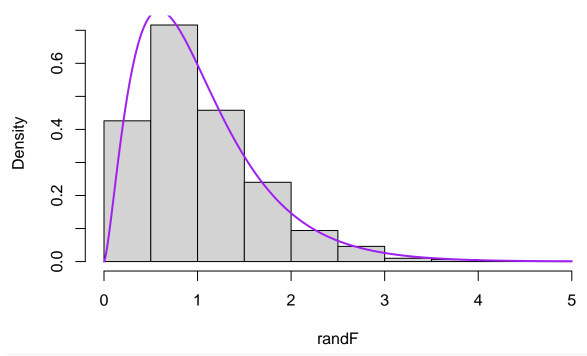
```
# set.seed allows for exact reproduction.
set.seed(12345678)
randF <- rf(1000, 5, 100)
# Generate histogram for the random numbers with exact.
hist(randF)</pre>
```

# Histogram of randF

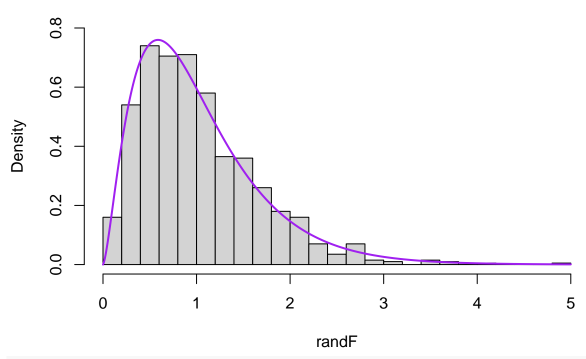


```
# Generate histogram for the random numbers with relative frequency.
# This is normalized, so we can superimpose an F-distribution to it.
hist(randF, freq = FALSE)
# Superimpose an F-distribution on the histogram.
lines(
    x,
    y = df(x, df1 = 5, df2 = 100),
    type = "l",
    col = "purple",
    lwd = 2
)
```

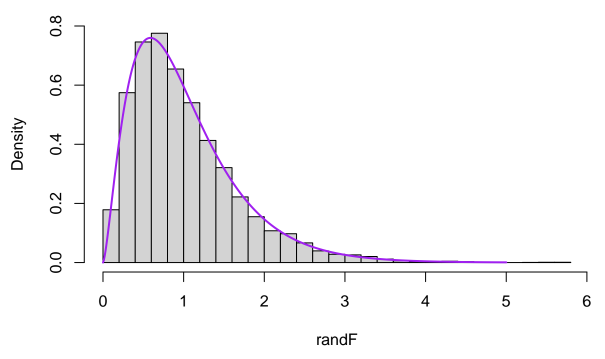
# Histogram of randF



# Histogram of randF



### Histogram of randF



#### Revisit Rocket Example

```
rocket <- read.csv("csv/rocket.csv")</pre>
m1 <- lm(thrust ~ nozzle + propratio, data = rocket)</pre>
summary(m1)
##
## lm(formula = thrust ~ nozzle + propratio, data = rocket)
##
## Residuals:
       Min
##
                1Q Median
                                3Q
                                       Max
## -3.8459 -1.7555 0.5934 1.2906 3.3008
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
                           4.7158 100.430 4.88e-15 ***
## (Intercept) 473.6039
## nozzle
                16.7383
                            1.5329 10.919 1.71e-06 ***
                -1.0948
## propratio
                            0.9414 - 1.163
                                              0.275
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 2.655 on 9 degrees of freedom
## Multiple R-squared: 0.9303, Adjusted R-squared: 0.9148
## F-statistic: 60.05 on 2 and 9 DF, p-value: 6.238e-06
# Compare summary with ANOVA table on board from Oct. 5.
anova(m1)
## Analysis of Variance Table
```

```
## Response: thrust
##
          Df Sum Sq Mean Sq F value Pr(>F)
## nozzle
           1 836.67 836.67 118.7377 1.743e-06 ***
## propratio 1 9.53
                       9.53
                                1.3524
                                         0.2748
## Residuals 9 63.42
                       7.05
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
anova(m1)$`Sum Sq`
## [1] 836.670000
                   9.529332 63.417335
sum(anova(m1)$`Sum Sq`[1:2])
## [1] 846.1993
SSRes <- anova(m1)$ Sum Sq [3]
# Test of overall significance.
m_red <- lm(thrust ~ 1, data = rocket)</pre>
summary(m_red)
##
## Call:
## lm(formula = thrust ~ 1, data = rocket)
## Residuals:
       Min
                 1Q Median
                                   3Q
## -13.4167 -7.1167 -0.2167 8.2333 11.3833
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 476.617 2.625 181.6 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 9.094 on 11 degrees of freedom
anova(m_red)
## Analysis of Variance Table
##
## Response: thrust
      Df Sum Sq Mean Sq F value Pr(>F)
## Residuals 11 909.62 82.692
SSRes_A <- anova(m_red) $ Sum Sq [1]
# Manually calculate F-statistic.
1 <- 2
n <- nrow(rocket)</pre>
p <- 2
Fstat <- ((SSRes_A - SSRes) / 1) / (SSRes / (n - p - 1))
Fstat
```

## [1] 60.04505

```
pval \leftarrow 1 - pf(Fstat, df1 = 1, df2 = n - p - 1)
pval
## [1] 6.238398e-06
# Automatically calculate F-statistic.
anova(m1, m red)$F[2]
## [1] 60.04505
Revist Coffee Example (Coffee Quality Institute, 2018)
coffee <- read.csv("csv/coffee arabica.csv")</pre>
mfull <-
 lm(
   Flavor ~ factor(Processing.Method) + Aroma + Aftertaste +
      Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
   dat = coffee
 )
summary(mfull)
##
## Call:
## lm(formula = Flavor ~ factor(Processing.Method) + Aroma + Aftertaste +
##
       Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
##
       data = coffee)
##
## Residuals:
       Min
                  1Q
                       Median
                                    30
                                             Max
## -0.68587 -0.08465 0.00079 0.08910 0.63633
##
## Coefficients:
##
                                                        Estimate Std. Error t value
                                                                   0.168516 -4.325
## (Intercept)
                                                       -0.728757
                                                                  0.022021 -0.063
## factor(Processing.Method)Semi-washed / Semi-pulped -0.001396
## factor(Processing.Method)Washed / Wet
                                                       -0.033061
                                                                   0.011024 - 2.999
## Aroma
                                                        0.220302
                                                                 0.020447 10.774
## Aftertaste
                                                                   0.023912 19.603
                                                        0.468759
## Body
                                                        0.096140 0.024334
                                                                              3.951
## Acidity
                                                        0.216751
                                                                   0.021194 10.227
## Balance
                                                        0.046806
                                                                   0.022558
                                                                              2.075
## Sweetness
                                                        0.025507
                                                                   0.010150
                                                                              2.513
                                                                   0.009803
## Uniformity
                                                        0.016297
                                                                              1.663
## Moisture
                                                        0.169012
                                                                   0.102480
                                                                              1.649
##
                                                       Pr(>|t|)
## (Intercept)
                                                       1.67e-05 ***
## factor(Processing.Method)Semi-washed / Semi-pulped 0.94947
## factor(Processing.Method)Washed / Wet
                                                        0.00277 **
## Aroma
                                                        < 2e-16 ***
## Aftertaste
                                                        < 2e-16 ***
## Body
                                                       8.28e-05 ***
                                                        < 2e-16 ***
## Acidity
## Balance
                                                        0.03823 *
## Sweetness
                                                        0.01211 *
                                                        0.09669 .
## Uniformity
```

```
0.09938 .
## Moisture
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.148 on 1108 degrees of freedom
## Multiple R-squared: 0.8091, Adjusted R-squared: 0.8073
## F-statistic: 469.5 on 10 and 1108 DF, p-value: < 2.2e-16
anova(mfull)
## Analysis of Variance Table
## Response: Flavor
                              Df Sum Sq Mean Sq F value
## factor(Processing.Method)
                               2 2.313
                                         1.156
                                                 52.8096 < 2.2e-16 ***
                               1 67.258 67.258 3071.2889 < 2.2e-16 ***
## Aroma
                               1 29.097 29.097 1328.6722 < 2.2e-16 ***
## Aftertaste
## Body
                              1 1.129
                                        1.129
                                                 51.5460 1.28e-12 ***
                               1 2.522
## Acidity
                                         2.522 115.1618 < 2.2e-16 ***
## Balance
                              1 0.116 0.116
                                                 5.2963 0.0215553 *
## Sweetness
                              1 0.251 0.251 11.4392 0.0007442 ***
                              1 0.064 0.064
## Uniformity
                                                2.9154 0.0880167 .
                               1 0.060 0.060
## Moisture
                                                2.7200 0.0993839 .
## Residuals
                          1108 24.264 0.022
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
SSRes <- anova(mfull) $`Sum Sq`[10]
# Reduced model without Uniformity and Moisture (beta9=beta10=0):
m red <-
 lm(
   Flavor ~ factor(Processing.Method) + Aroma + Aftertaste +
     Body + Acidity + Balance + Sweetness,
   dat = coffee
 )
summary(m red)
##
## lm(formula = Flavor ~ factor(Processing.Method) + Aroma + Aftertaste +
##
      Body + Acidity + Balance + Sweetness, data = coffee)
##
## Residuals:
                 1Q
                     Median
                                  3Q
## -0.67907 -0.08487 0.00054 0.08490 0.64763
## Coefficients:
                                                     Estimate Std. Error t value
## (Intercept)
                                                    -0.606791 0.159741 -3.799
## factor(Processing.Method)Semi-washed / Semi-pulped 0.002275 0.021969 0.104
## factor(Processing.Method)Washed / Wet
                                                    -0.031115 0.011009 -2.826
## Aroma
                                                     0.221362 0.020472 10.813
## Aftertaste
                                                     0.470849 0.023858 19.735
## Body
                                                     0.087671 0.024102 3.637
```

```
0.219257
                                                                 0.021182 10.351
## Acidity
## Balance
                                                      0.047526 0.022283
                                                                           2.133
## Sweetness
                                                      0.032406 0.009597
                                                                            3.377
##
                                                     Pr(>|t|)
## (Intercept)
                                                     0.000153 ***
## factor(Processing.Method)Semi-washed / Semi-pulped 0.917539
## factor(Processing.Method)Washed / Wet
                                                     0.004795 **
                                                      < 2e-16 ***
## Aroma
## Aftertaste
                                                      < 2e-16 ***
## Body
                                                     0.000288 ***
## Acidity
                                                      < 2e-16 ***
## Balance
                                                     0.033160 *
                                                     0.000759 ***
## Sweetness
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.1482 on 1110 degrees of freedom
## Multiple R-squared: 0.8081, Adjusted R-squared: 0.8067
## F-statistic: 584.2 on 8 and 1110 DF, p-value: < 2.2e-16
anova(m_red)
## Analysis of Variance Table
##
## Response: Flavor
                              Df Sum Sq Mean Sq F value
                                                            Pr(>F)
                                                 52.637 < 2.2e-16 ***
                                         1.156
## factor(Processing.Method)
                               2 2.313
                               1 67.258 67.258 3061.263 < 2.2e-16 ***
## Aroma
## Aftertaste
                               1 29.097 29.097 1324.335 < 2.2e-16 ***
                                                 51.378 1.387e-12 ***
## Body
                               1 1.129
                                         1.129
                                          2.522 114.786 < 2.2e-16 ***
## Acidity
                               1 2.522
## Balance
                               1 0.116 0.116 5.279 0.0217690 *
## Sweetness
                               1 0.251 0.251 11.402 0.0007591 ***
## Residuals
                            1110 24.387
                                          0.022
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
SSRes_A <- anova(m_red)$`Sum Sq`[8]
# Manually calculate F-statistic.
1 <- 2
n <- nrow(coffee)
p < -10
Fstat <- ((SSRes_A - SSRes) / 1) / (SSRes / (n - p - 1))
Fstat
## [1] 2.81769
pval \leftarrow 1 - pf(Fstat, df1 = 1, df2 = n - p - 1)
pval
## [1] 0.06017197
# Automatically calculate F-statistic.
anova(mfull, m_red)$F[2]
## [1] 2.81769
```

```
# Reduced model without Uniformity and Moisture and
# setting effect of Dry = Semi (beta1=beta9=beta10=0)
# 1 = wet, 0 otherwise
coffee$method2 <- ifelse(coffee$Processing.Method %in%</pre>
                          c('Natural / Dry', 'Semi-washed / Semi-pulped'),
                        0,
                        1)
# 1 = semi/dry, 0 o.w
coffee$wet <-
 ifelse(coffee$Processing.Method == 'Washed / Wet', 0, 1)
m_red2 <- lm(Flavor ~ method2 + Aroma + Aftertaste +
              Body + Acidity + Balance + Sweetness,
            dat = coffee)
summary(m_red2)
##
## Call:
## lm(formula = Flavor ~ method2 + Aroma + Aftertaste + Body + Acidity +
      Balance + Sweetness, data = coffee)
##
## Residuals:
##
       Min
                 1Q
                    Median
                                  30
                                          Max
## -0.67906 -0.08508 0.00052 0.08490 0.64722
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -0.606597   0.159659   -3.799   0.000153 ***
## method2
              -0.031543
                         0.010200 -3.092 0.002036 **
## Aroma
               ## Aftertaste 0.470861
                         0.023847 19.745 < 2e-16 ***
                                   3.638 0.000287 ***
## Body
              0.087561
                         0.024068
## Acidity
                        0.021173 10.356 < 2e-16 ***
               0.219266
## Balance
              0.047527
                         0.022273
                                   2.134 0.033077 *
## Sweetness 0.032462
                         0.009577 3.389 0.000725 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1482 on 1111 degrees of freedom
## Multiple R-squared: 0.8081, Adjusted R-squared: 0.8069
## F-statistic: 668.3 on 7 and 1111 DF, p-value: < 2.2e-16
anova(m red2)
## Analysis of Variance Table
## Response: Flavor
               Df Sum Sq Mean Sq F value
##
                                             Pr(>F)
## method2
               1 2.313
                         2.313 105.3648 < 2.2e-16 ***
                1 67.255 67.255 3063.8526 < 2.2e-16 ***
## Aroma
                1 29.100 29.100 1325.6571 < 2.2e-16 ***
## Aftertaste
## Body
                1 1.126
                         1.126 51.3088 1.434e-12 ***
## Acidity
               1 2.522
                         2.522 114.9115 < 2.2e-16 ***
## Balance
              1 0.116
                         0.116 5.2882 0.0216552 *
```

```
## Sweetness 1 0.252 0.252 11.4883 0.0007249 ***
## Residuals 1111 24.388 0.022
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
SSRes_A <- anova(m_red2) $ Sum Sq [8]
## Manually calculate F-statistic.
1 <- 3
n <- nrow(coffee)</pre>
p <- 10
Fstat \leftarrow ((SSRes_A - SSRes) / 1) / (SSRes / (n - p - 1))
Fstat
## [1] 1.882046
pval <-1 - pf(Fstat, df1 = 1, df2 = n - p - 1)
## [1] 0.1308207
{\it \# Automatically calculate F-statistic.}
anova(mfull, m_red2)$F[2]
## [1] 1.882046
```

#### LECTURE 12 | 2020-10-21

### 2.11 Multicollinearity

Multicollinearity: occurs when some explanatory variables have a **strong linear** relationship amongst themselves. For example, this might occur exactly

$$\boldsymbol{x}_3 = \alpha_0 \boldsymbol{1} + \alpha_1 \boldsymbol{x}_1 + \alpha_2 \boldsymbol{x}_2$$

in which case the columns of X would be **linearly dependent** and  $X^{T}X$  does not have an inverse. Practically, there is no new info including  $x_3$  when  $x_1, x_2$  are in the model. **Approximately**,

$$\boldsymbol{x}_3 \approx \alpha_0 \boldsymbol{1} + \alpha_1 \boldsymbol{x}_1 + \alpha_2 \boldsymbol{x}_2$$

in which case the columns of X are close to being linearly dependent which cases  $\mathbb{V}(\hat{\beta}_j)$  to be **inflated**, in turn leads to inaccurate confidence intervals and conclusions of hypothesis tests for the regression parameters, in practice.  $\mathrm{Se}(\hat{\beta}_j)$  when fitting models can change drastically when adding/removing variables from the model.

#### EXAMPLE 2.11.1: Hockey (NHL)

In the NHL we have Goals + Assists = Points. Suppose we want to predict a forward's salary. Define

- $x_1 = \text{Goals}$
- $x_2 = Assists$
- $x_3 = Points$

 $x_3 = x_1 + x_2$ , therefore we have exact multicollinearity.

#### **EXAMPLE 2.11.2: Burmese Pythons in Florida (2017)**

- y =fat content
- $x_1 = \text{mass}$
- $x_2$  = overall length
- $x_3 = \text{snout-to-vent length}$

It turns out that  $x_2$  and  $x_3$  are highly correlated. Including all variables in regression lead to inflated  $Se(\hat{\beta}_2)$  and  $Se(\hat{\beta}_3)$ .

## 2.12 Detection of Multicollinearity

If two predictors are related

- Scatter plot matrix [all possible pairs of scatter plots b/w  $y, x_1, x_2, \dots, x_n$ ]
- Correlation matrix (all pairwise correlations)

#### **DEFINITION 2.12.1: Variance inflation factor**

For multicollinearity between more than two predictors, we can define the **variance inflation factor** (VIF).

$$\text{VIF}_j = \frac{\mathbb{V}(\hat{\beta}_j)}{\mathbb{V}(\hat{\beta}_i^*)}$$

for  $j=1,\ldots,p$ , where  $\hat{\beta}_j$  is the estimate of  $\beta_j$  with all predictors in the model, and  $\hat{\beta}_j^*$  estimate of  $\beta_j$  based on regression with  $x_j$  only.

#### **THEOREM 2.12.2**

$$VIF_j \geqslant 1$$

Fit multiple linear regression model of  $\boldsymbol{x}_j$  in terms of other predictors; that is,

$$x_{ij} = \alpha_0 + \alpha_1 x_{i1} + \dots + \alpha_{j-1} x_{i(j-1)} + \alpha_{j+1} x_{i(j+1)} + \dots + \alpha_p x_{ip} + \varepsilon_{ij}$$

and compute  $R^2$  for this model, call it  $R_i^2$ .

Intuition: if  $R_j^2$  is close to 1,  $x_j$  is strongly related linearly to other predictors. It can be shown that

$$\text{VIF}_j = \frac{1}{1 - R_j^2}$$

If  ${
m VIF}_j \geq 10$ , then there is solid evidence of multicollinearity; that is,  $R_j^2 > 0.9$ .

Procedure:

• remove predictors with largest VIF, if it exceeds 10. Repeat until no more multicollinearity.

#### 2.12.1 R Demo

```
## Coffee example (Coffee Quality Institute, 2018) continued
coffee <- read.csv("csv/coffee_arabica.csv")</pre>
# cor(coffee) doesn't work as there's a categorical variable.
cor(coffee[, -1]) # e.g., remove first column
##
                     Aroma
                               Flavor Aftertaste
                                                          Body
                                                                                Balance
                                                                    Acidity
## Aroma
               1.00000000 0.7339782 0.6892744 0.56699932 0.60115765 0.6156508
## Flavor
               0.73397820 1.0000000 0.8582783 0.67694834
                                                                 0.73845546 0.7324530
## Aftertaste 0.68927440 0.8582783 1.0000000 0.67407704
                                                                 0.69408861 0.7657979
## Body
               0.56699932 0.6769483 0.6740770 1.00000000 0.60795391 0.6924568
## Acidity
               0.60115765 0.7384555 0.6940886 0.60795391 1.00000000 0.6417994
## Balance 0.61565084 0.7324530 0.7657979 0.69245676 0.64179938 1.0000000
## Sweetness 0.06955938 0.1345364 0.1185760 0.03977892
                                                                 0.06906093
                                                                             0.1016718
## Uniformity 0.14785498 0.2132347 0.2143116 0.07195778 0.14876428 0.2180726
## Moisture -0.11567549 -0.1327342 -0.1745366 -0.21009097 -0.10391684 -0.2161964
##
               Sweetness Uniformity
                                         Moisture
              0.06955938 0.14785498 -0.11567549
## Aroma
## Flavor 0.13453644 0.21323472 -0.13273418
## Aftertaste 0.11857600 0.21431157 -0.17453658
## Body
              0.03977892 0.07195778 -0.21009097
## Acidity
              0.06906093 0.14876428 -0.10391684
              0.10167183 0.21807265 -0.21619640
## Balance
## Sweetness 1.00000000 0.34756414 0.08049300
## Uniformity 0.34756414 1.00000000 0.02105693
## Moisture 0.08049300 0.02105693 1.00000000
Plot the pairs (disabled due to loading time on PDF).
  ~ Flavor + Aroma + Aftertaste + Body +
    Acidity + Balance + Sweetness + Uniformity + Moisture,
  data = coffee
# Code our own indicators, so that we can more easily interpret VIFs.
# 1 = wet, 0 otherwise
coffee$wet <-</pre>
  ifelse(coffee$Processing.Method == 'Washed / Wet', 1, 0)
# 1 = semi/dry, 0 otherwise
coffee$semi <-
  ifelse(coffee$Processing.Method == 'Semi-washed / Semi-pulped',
         1, 0)
Model:
     y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \beta_4 x_{i4} + \beta_5 x_{i5} + \beta_6 x_{i6} + \beta_7 x_{i7} + \beta_8 x_{i8} + \beta_9 x_{i9} + \beta_{10} x_{i(10)} + \varepsilon_i
where
  • y = \text{flavour}
  • x_1 = 1 if wet, 0 otherwise
  • x_2 = 1 if semi, 0 otherwise
  • x_3 = \text{Aroma}
  • x_4 = \text{Aftertaste}
```

```
• x_5 = \text{Body}
  • x_6 = Acidity
  • x_7 = Balance
  • x_8 = \text{Sweetness}
  • x_9 = \text{Uniformity}
  • x_{10} = Moisture
# Full MLR with our manually coded indicators.
mfull <- lm(
 Flavor ~ wet + semi + Aroma + Aftertaste +
   Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
 dat = coffee
summary(mfull)
##
## Call:
## lm(formula = Flavor ~ wet + semi + Aroma + Aftertaste + Body +
      Acidity + Balance + Sweetness + Uniformity + Moisture, data = coffee)
##
## Residuals:
##
       Min
                1Q
                    Median
                                 3Q
## -0.68587 -0.08465 0.00079 0.08910 0.63633
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.728757
                        0.168516 -4.325 1.67e-05 ***
                        0.011024 -2.999 0.00277 **
## wet
              -0.033061
             ## semi
## Aroma
              ## Aftertaste 0.468759
                        0.023912 19.603 < 2e-16 ***
## Body
              0.096140
                        0.024334
                                  3.951 8.28e-05 ***
                        0.021194 10.227 < 2e-16 ***
## Acidity
              0.216751
## Balance
              0.010150
## Sweetness
              0.025507
                                 2.513 0.01211 *
## Uniformity 0.016297
                         0.009803
                                  1.663 0.09669 .
## Moisture
              0.169012
                         0.102480
                                 1.649 0.09938 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.148 on 1108 degrees of freedom
## Multiple R-squared: 0.8091, Adjusted R-squared: 0.8073
## F-statistic: 469.5 on 10 and 1108 DF, p-value: < 2.2e-16
# Full MLR alternative, using the factor command.
mfull_alternative <-
 lm(
   Flavor ~ factor(Processing.Method) + Aroma + Aftertaste +
     Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
   dat = coffee
```

Suppose we want to check the VIF for j = 1; that is,  $x_1$ . Now, we fit:

```
x_{i1} = \alpha_0 + \alpha_2 x_{i2} + \alpha_3 x_{i3} + \alpha_4 x_{i4} + \alpha_5 x_{i5} + \alpha_6 x_{i6} + \alpha_7 x_{i7} + \alpha_8 x_{i8} + \alpha_9 x_{i9} + \alpha_{10} x_{i(10)} + \varepsilon_i
```

```
wet_reg <-
  lm(
   wet ~ semi + Aroma + Aftertaste + Body + Acidity + Balance +
     Sweetness + Uniformity + Moisture,
   dat = coffee
summary(wet_reg)
##
## Call:
## lm(formula = wet ~ semi + Aroma + Aftertaste + Body + Acidity +
       Balance + Sweetness + Uniformity + Moisture, data = coffee)
##
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
  -1.0015 -0.0283 0.1770 0.2522 0.7704
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.81748 0.45838
                                     1.783 0.074794 .
## semi
               -0.75675
                           0.05551 -13.632 < 2e-16 ***
               0.09690
                           0.05562
                                    1.742 0.081774 .
## Aroma
## Aftertaste -0.13169
                           0.06502 -2.026 0.043054 *
                           0.06596 -3.318 0.000936 ***
## Body
              -0.21885
## Acidity
               0.18696
                           0.05746
                                    3.254 0.001173 **
## Balance
              -0.10804
                           0.06136 -1.761 0.078563 .
## Sweetness
               0.08373
                           0.02753
                                    3.041 0.002413 **
## Uniformity
               0.03547
                           0.02668
                                     1.329 0.184053
## Moisture
                0.59486
                           0.27858
                                     2.135 0.032956 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.4031 on 1109 degrees of freedom
## Multiple R-squared: 0.1911, Adjusted R-squared: 0.1845
## F-statistic: 29.11 on 9 and 1109 DF, p-value: < 2.2e-16
r2_wet <- summary(wet_reg)$r.squared
r2_wet
## [1] 0.191077
R_i: In our case, R_1 = 0.191077.
VIF_wet <- 1 / (1 - r2_wet)</pre>
VIF_wet
```

#### ## [1] 1.236212

 $VIF_j$ :  $VIF_1 = 1.236212$ . Interpretation: in a regression with all the variables compared to a regression with just this one, the estimated variance has increased by a factor of 1.24, which is not a very large inflation. The variable wet is not very linearly correlated or dependent on the other predictors that we have in the model.

```
Aroma_reg <- lm(
  Aroma ~ wet + semi + Aftertaste +
   Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
)
r2_Aroma <- summary(Aroma_reg)$r.squared
r2_Aroma
## [1] 0.5204716
VIF_Aroma <- 1 / (1 - r2_Aroma)</pre>
VIF_Aroma
## [1] 2.085382
R_3 = 0.5204716, VIF<sub>3</sub> = 2.085382.
Aftertaste_reg <- lm(
  Aftertaste ~ wet + semi + Aroma +
   Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
  dat = coffee
r2_Aftertaste <- summary(Aftertaste_reg)$r.squared
r2_Aftertaste
## [1] 0.7101012
VIF_Aftertaste <- 1 / (1 - r2_Aftertaste)</pre>
VIF_Aftertaste
## [1] 3.449479
# Load car library for automatic VIF calculation using vif()
library(car)
vif(mfull)
##
                              Aroma Aftertaste
          wet
                    semi
                                                      Body
                                                              Acidity
                                                                         Balance
##
    1.236212
              1.178004
                           2.085382 3.449479
                                                 2.317728 2.232210
                                                                        3.002813
##
   Sweetness Uniformity
                           Moisture
   1.159602
               1.209901
                           1.086101
No serious signs of inflation, all VIFs are less than 10.
## Python in FL everglades example (2017)
## Sex, length, total mass, fat mass, and specimen condition data for
## 248 Burmese pythons (Python bivittatus) collected in the Florida Everglades
python <- read.csv("csv/FLpython.csv")</pre>
head(python)
##
    sex svl mass length
                             fat
## 1 F 70.0 186
                    77.5 6.000
## 2 M 76.0 310
                    83.8 11.000
## 3 M 77.0 260
                    86.1 6.000
## 4 M 78.0 262 87.1 8.000
## 5 M 81.0 306 91.1 4.000
## 6 M 93.5 605 104.6 18.959
```

```
python\frac{s}{male} \leftarrow ifelse(python sex == 'M', 1, 0) # 1 = M, 0 = F
cor(python[, -1])
##
                 svl
                            mass
                                     length
                                                    fat
                                                              male
## svl
           1.0000000 0.8843022 0.9994935 0.8098652 -0.1602418
## mass
           0.8843022 \quad 1.0000000 \quad 0.8858256 \quad 0.9419114 \quad -0.2190993
## length 0.9994935 0.8858256 1.0000000 0.8114658 -0.1593512
           0.8098652  0.9419114  0.8114658  1.0000000  -0.2933111
## fat
          -0.1602418 -0.2190993 -0.1593512 -0.2933111 1.0000000
## male
pairs(python[, -1])
mpf <- lm(fat ~ male + svl + mass + length, data = python)</pre>
summary(mpf)
##
## Call:
## lm(formula = fat ~ male + svl + mass + length, data = python)
##
## Residuals:
##
        Min
                  1Q
                        Median
                                     3Q
                                              Max
## -2445.77 -137.41
                        -5.29
                                 110.00 1527.27
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.021e+02 1.331e+02
                                        1.518
                                                  0.130
               -1.971e+02 4.732e+01 -4.165 4.32e-05 ***
## male
## svl
               -3.370e+00 1.125e+01 -0.300
                                                  0.765
                1.178e-01
                           5.302e-03 22.210
                                                < 2e-16 ***
## mass
                1.594e+00 1.010e+01
                                        0.158
                                                  0.875
## length
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 360.9 on 243 degrees of freedom
## Multiple R-squared: 0.897, Adjusted R-squared: 0.8953
## F-statistic: 529 on 4 and 243 DF, p-value: < 2.2e-16
vif(mpf)
##
                        svl
                                              length
          male
                                   mass
##
      1.058699 994.546545
                               4.813078 1007.484200
mpf_l <- lm(length ~ male + svl + mass, data = python)</pre>
1 / (1 - summary(mpf_l)$r.squared)
## [1] 1007.484
Misleading conclusion: svl and length are both irrelevant (this is not the case). Also, the standard errors
are very large.
# remove "length" based on VIF
mpf2 <- lm(fat ~ male + mass + svl, data = python)</pre>
summary(mpf2)$adj
```

## [1] 0.8957164

```
vif(mpf2)
     male
            mass
## 1.056139 4.720065 4.611903
anova(mpf2)
## Analysis of Variance Table
## Response: fat
##
      Df
             Sum Sq Mean Sq F value Pr(>F)
        1 26435988 26435988 203.7689 < 2e-16 ***
## male
## Residuals 244 31655372
                       129735
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
AIC(mpf2)
## [1] 3629.527
```

svl now has a significant t-statistic.

### LECTURE 13 | 2020-10-26

### 2.13 Model Selection Criteria

Given p explanatory variables, find the subset  $k \le p$  of explanatory variables ("reduced model") that gives us the "best" model:

- · Goodness of fit.
- Interpretability.
- Predictive performance.

Some related concepts:

1. *F* tests compare between 2 specific models where test adequacy of a "reduced" model (subset, "nested") relative to full model.

Quiz 4: 
$$\beta_1 = \beta_2$$
 in part d-f

- 2. Multicollinearity: can affect interpretability of  $\hat{\beta}_j$  usual interpretation "holding other variables constant" doesn't really work when  $x_j$  is strongly correlated with other predictors.
- 3.  $R^2$  is the proportion of variability in the response explained by the regression model. It always increases when adding variables.
- 4.  $\hat{\sigma}^2$  is estimated residual variable, used for prediction, want  $\hat{\sigma}^2$  small to give good predictive performance Two key ingredients:
  - Metric (or criterion) for comparing different models with potentially different number of predictors
  - Selection/search strategy (which models should we fit and test?)

Examples of metrics for model selection:

#### **DEFINITION 2.13.1:** Adjusted $R^2$

$$R_{\mathrm{adj}}^2 = 1 - \frac{\mathrm{SS}(\mathrm{Res})/(n-k-1)}{\mathrm{SS}(\mathrm{Total})/(n-1)} = 1 - \frac{\hat{\sigma}^2}{s^2}$$

for model with k predictors.

Compared to

$$R^2 = 1 - \frac{\mathrm{SS}(\mathrm{Res})}{\mathrm{SS}(\mathrm{Total})} = 1 - \frac{\hat{\sigma}^2(n-k-1)}{s^2(n-1)}$$

- SS(Res)/(n-k-1) estimated  $\hat{\sigma}^2$  for model with k predictors
- SS(Total)/(n-1) is the sample variance of responses  $y_i$ .

$$\begin{split} R_{\rm adj}^2 &= 1 - \frac{n-1}{n-k-1}(1-R^2) = 1 - \left(1 + \frac{k}{n-k-1}\right)(1-R^2) \\ &= 1 - \left[1(1-R^2) + \left(\frac{k}{n-k-1}\right)(1-R^2)\right] \\ &= 1 - \left[1 - R^2 + \left(\frac{k}{n-k-1}\right)(1-R^2)\right] \\ &= 1 - 1 + R^2 - \frac{k}{n-k-1}(1-R^2) \\ &= R^2 - (1-R^2)\frac{k}{n-k-1} \end{split}$$

Intuition:  $R_{\rm adj}^2$  accounts for number variables in model, penalizes inclusion of unimportant predictors; that is, SS(Res) has little decrease when adding such variables. Meanwhile,  $R^2$  always increases with more predictors, but  $R_{\text{adi}}^2$  can decrease if SS(Res) change is small.

While  $R_{\rm adi}^2$  loses its usual interpretation of  $R^2$ , but can be used as a measure of "goodness of fit" and model selection criterion (e.g., pick subset of predictors that gives the highest  $R_{adi}^2$ ).

#### **EXAMPLE 2.13.2**

#### Given

- n = 25
- SS(Total) = 20

Suppose we're considering on a subset of k = 4 predictors, and find:

	Reduced	Full
	k = 4	p=6
SS(Total)	20	20
SS(Res)	10	9.8
$R^2$	10/20 = 0.5	9.8/20 = 0.49
$R^2_{ m adj} \ \widehat{\sigma}^2$	$1 - \frac{10/(25 - 4 - 1)}{20/(25 - 1)} = 0.4$	$1 - \frac{9.8/(25-6-1)}{20/(25-1)} \approx 0.347$
$\hat{\sigma}^2$	10/(25-4-1)=0.5	$9.8/(25-6-1) \approx 0.544$

- n-k-1 d.f. Res in reduced
- n-p-1 d.f. Res in full

- $R_{\mathrm{adj}}^2 < R^2$ , but as  $n \to \infty$ ,  $R_{\mathrm{adj}}^2 \to R^2$ . Model with higher  $R_{\mathrm{adj}}^2$  has lower  $\hat{\sigma}^2$ , thus is a reasonable metric for model selection.

#### **DEFINITION 2.13.3: Akaike Information Criterion (AIC)**

Let n be sample size and q be the number of estimated parameters.

• MLR: q = p + 2 since we have p predictors + 1 intercept  $(\beta_0) + 1$   $(\sigma^2)$ .

The Akaike information criterion (AIC) is defined as

$$\mathrm{AIC} = 2q - 2\ln[L(\hat{\theta})]$$

where  $L(\hat{\theta})$  is the likelihood function evaluated at  $\hat{\theta}$  (parameter estimates).

#### **REMARK 2.13.4**

- The least square estimates of  $\beta$  are equivalent to maximum likelihood estimates under the usual normal assumptions on  $\varepsilon$ .
- 2q is the penalty for including more predictors. With more parameters,  $L(\hat{\theta})$  increases, offset by penalty 2q.
- The model with lower AIC is preferred; that is, differences in AIC matter not the value itself.

#### **REMARK 2.13.5:** †

If we want to measure just the difference of AIC, we can do

$$AIC = n \ln \left[ \frac{SS(Res)}{n} \right] + 2q$$

#### **DEFINITION 2.13.6: Bayesian Information Criterion (BIC)**

Let n be sample size and q be the number of estimated parameters.

• MLR: q = p + 2 since we have p predictors + 1 intercept  $(\beta_0) + 1$   $(\sigma^2)$ .

The Bayesian information criterion (BIC) is defined as

$$\mathrm{BIC} = q \ln(n) - 2 \ln[L(\hat{\theta})]$$

where  $L(\hat{\theta})$  is the likelihood function evaluated at  $\hat{\theta}$  (parameter estimates).

#### **REMARK 2.13.7**

AIC is similar to BIC, but BIC strongly penalizes inclusion of more variables. Note that in BIC,  $q \ln(n)$  depends on sample size.

#### **REMARK 2.13.8:** †

If we want to measure just the difference of BIC, we can do

$$\mathrm{BIC} = n \ln \left[ \frac{\mathrm{SS}(\mathrm{Res})}{n} \right] + q \ln(n)$$

#### Recap:

- $R^2$ , AIC, BIC are all based on comparing the fitted models. In other words, they look at the explanatory power of the model.
- They all have penalties to try to prevent "overfitting." That is, having too many variables might end up modelling spurious relationships that are actually noise.

#### **Mean Square Prediction Error (MSPE)**

Consider predictive performance of model on *new* data; that is, data *not* used in fitting of models. "Is model generalisable to new data?" Overfitted models tend to have high prediction error.

For example, via cross-validation schemes. We've given 4 examples of metrics/criteria for comparing models. Imagine we have p predictors:

$$\begin{array}{ccc} \binom{p}{1} & 1 \text{ predictor} \\ \binom{p}{2} & 2 \text{ predictors} \\ \vdots & & \\ \binom{p}{n} & p \text{ predictors} \end{array} \Longrightarrow \sum_{j=0}^{p} \binom{p}{j} = 2^{p}$$

Occam's Razor: "The simplest explanation is usually the best one."—William Ockham

### LECTURE 14 | 2020-10-28

### 2.14 Model Selection Basic Strategies

- Criteria:  $R_{\rm adi}^2$ , AIC, BIC, MSPE, etc. explicitly penalizes unnecessarily complex models.
- Search strategies (use with chosen criterion)
- (i) Brute force: fit all possible regressions. With p predictors, we have  $\sum_{i=0}^{p} {p \choose i} = 2^p$  possible models to fit.
  - Finds optimal model that may be computationally intensive (or infeasible) if *p* is large.

<u>Idea</u>: Find a "good" (useful) model in reasonable computational time (not necessarily optimal). Many strategies focus on adding/removing variables one at a time.

- (ii) Forward selection: add one variable at a time to model.
  - Start with a model that only has an intercept  $(\beta_0)$ .
  - Fit p simple linear regression models

$$\mathbf{y} = \beta_0 \mathbf{1} + \beta_1 \mathbf{x}_j + \boldsymbol{\varepsilon} \quad j = 1, \dots, p$$

- Pick the best of p models (with 1 predictor) according to chosen criterion, and add that variable  $x_i$  to model.
- Fit (p-1) models containing  $x_i$  and one other variable.
  - If none of (p-1) models improves criterion, stop.
  - Pick the best of (p-1) models according to criterion, so now we have 2 variables in the model.

Continue adding 1 variable at a time in this way until we can no more variables improve the criterion. The final model is one with the best criterion after we *stop*; that is, no further improvement is possible.

Note: Much faster than brute force as the maximum number of models to fit is:

$$p + (p-1) + \dots + 2 + 1 = \sum_{i=1}^{p} i = \frac{p(p+1)}{2}$$

which is  $\mathcal{O}(p^2)$  compared to  $\mathcal{O}(2^p)$  for all possible regressions.

- (iii) Backward direction: remove one variable at a time to model.
  - Start with model that has *p* predictors.
  - Fit p models that result from removing one variable from the regression; that is, each one has (p-1) variables.
  - Pick the best of p models according to criterion.
    - Eliminate that variable  $x_i$  from model.
    - Fit (p-1) models that remove  $x_i$  and one other variable from model.
    - Pick best of (p-1) models (2 variables removed).

Continue removing 1 variable at a time in this way until we can no more variables improve the criterion. Same computational complexity as forward selection.

(iv) Forward-backwards (allows individual variables to be both added/removed)

- Start as in forward selection
- If we have *k* variables in model:
  - Backwards: fit k models with (k-1) variables. If any of these improve criterion, remove the variable.
  - Forwards: fit (p-k) models with (k+1) variables. If any of these improve criterion, add that variable.
- These are the basic "stepwise" selection models to get a "good" (useful) model.
- Many other have sophisticated procedures available. For example, stochastic search, lasso.
- We've assumed that n>p because otherwise  $(X^{\top}X)$  is not invertible. More specialized methods needed if number of predictors is larger than sample size.

#### 2.14.1 R Demo

```
## Coffee example (Coffee Quality Institute, 2018) continued.
coffee <- read.csv("csv/coffee_arabica.csv")</pre>
mfull <-
  lm(
    Flavor ~ factor(Processing.Method) + Aroma + Aftertaste +
      Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
    dat = coffee
  )
summary(mfull)$adj.r.squared
AIC(mfull)
BIC(mfull)
library(leaps)
# Exhaustive, brute-force search.
all_regs <-
  regsubsets(
    Flavor ~ .,
    data = coffee,
    nvmax = 10,
    nbest = 2 ^10,
    really.big = TRUE
  )
all_regs_summ <- summary(all_regs)</pre>
all_regs_summ$which
all_regs_summ$adjr2
all_regs_summ$bic
# Organize results according to number of variables in model.
p <- 10
k < -c(
  rep(1, choose(p, 1)),
  rep(2, choose(p, 2)),
  rep(3, choose(p, 3)),
  rep(4, choose(p, 4)),
  rep(5, choose(p, 5)),
  rep(6, choose(p, 6)),
  rep(7, choose(p, 7)),
  rep(8, choose(p, 8)),
  rep(9, choose(p, 9)),
  rep(10, choose(p, 10))
boxplot(
  all_regs_summ$adjr2 ~ k,
 xlab = "Number of predictors",
 ylab =
    expression(R[adj] ^ 2),
  ylim = c(0, 1)
abline(h = c(0, 1), lty = 2, col = "red")
boxplot(all_regs_summ$bic ~ k, xlab = "Number of predictors", ylab = "BIC")
```

```
max(all_regs_summ$adjr2)
bestR2adj <- which.max(all_regs_summ$adjr2)</pre>
min(all_regs_summ$bic)
bestBIC <- which.min(all_regs_summ$bic)</pre>
# Find out which predictors in those models.
all_regs_summ$which[bestR2adj, ]
all regs summ$which[bestBIC, ]
coffee$wet <- ifelse(coffee$Processing.Method == 'Washed / Wet', 1,</pre>
                      0) # 1 = wet, 0 otherwise
coffee$semi <-
  ifelse(coffee$Processing.Method == 'Semi-washed / Semi-pulped',
         1, 0) # 1 = semi/dry, 0 otherwise
coffee$Processing.Method <- NULL</pre>
m_bestr2adj <- lm(</pre>
  Flavor ~ wet + Aroma + Aftertaste +
    Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
  dat = coffee
summary(m_bestr2adj)
AIC(m_bestr2adj)
BIC(m_bestr2adj)
m bestBIC <- lm(Flavor ~ wet + Aroma + Aftertaste +
                   Body + Acidity + Sweetness , dat = coffee)
summary(m_bestBIC)
AIC(m_bestBIC)
BIC(m_bestBIC)
# Let's also try stepwise methods.
library(MASS)
# Full model and empty model with just intercept.
full <- lm(Flavor ~ ., data = coffee)</pre>
empty <- lm(Flavor ~ 1, data = coffee)</pre>
# Default stepAIC uses AIC criterion.
m_f_AIC <- stepAIC(</pre>
 object = empty,
 scope = list(upper = full, lower = empty),
 direction = "forward"
# Let's get stepAIC to use BIC by specifying the penalty k = log(n).
# Add 'trace = 0' to hide the output.
# Forward.
m_f <-
 stepAIC(
   object = empty,
    scope = list(upper = full, lower = empty),
```

```
direction = "forward",
   trace = 0,
   k = log(nrow(coffee))
  )
summary(m_f)
# Backward.
m_b <-
  stepAIC(
    object = full,
    scope = list(upper = full, lower = empty),
    direction = "backward",
   trace = 0,
    k = log(nrow(coffee))
summary(m_b)
# Forward-backward.
m_h <-
 stepAIC(
    object = empty,
   scope = list(upper = full, lower = empty),
   direction = "both",
   trace = 0,
    k = log(nrow(coffee))
  )
summary(m_h)
```

10 variables is still a fairly small problem: in this example all 3 approaches identify the same BIC-based model as the exhaustive search.

#### LECTURE 15 | 2020-11-02

### 2.15 Residual Plots for Linear Regression Assumptions

Recall:  $\mathbf{Y} = X\boldsymbol{\beta} + \boldsymbol{\varepsilon}$  where  $\boldsymbol{\varepsilon} \sim \text{MVN}(0, \sigma^2 I_n)$ . Practically, this means

$$\varepsilon \overset{\mathrm{iid}}{\sim} \mathcal{N}(0,\sigma^2)$$

- · Independence among all error terms
- · Normally distributed
- Since  $\mathbb{E}[Y_i] = \beta_0 + \beta_1 x_{i1} + \dots + \beta_p x_{ip}$  is implied by  $\mathbb{E}[\varepsilon_i] = 0$  for any  $x_{i1}, \dots, x_{ip}$ , the linear model is appropriate; that is, it correctly explains response on average.
- Constant error variance  $\sigma^2$

We could assess these assumptions via  $\varepsilon_i$ 's, but we can't observe  $\varepsilon_i$  directly. Rather, we do have an approximation via residuals  $e_i$  from the fitted model.

Recall,  $e \sim \text{MVN}(0, (I - H)\sigma^2)$ . So e and  $\varepsilon$  are related:

$$e = Y - X\hat{\beta} = (I - H)Y = (I - H)(X\beta + \varepsilon) = (X\beta - HX\beta) + (I - H)\varepsilon = (I - H)\varepsilon$$

Note: we can't "solve"  $\varepsilon$  since (I-H) is not invertible since it is not full rank: recall H and (I-H) are idempotent, so

$$\operatorname{tr}(H) = p + 1 = \operatorname{rank} X$$

$$\operatorname{tr}(I-H) = n - (p+1) < n$$

Similarly, this does not imply  $Y = \varepsilon$ . So,  $e_i = \varepsilon_i - \sum_{j=1}^n h_{ij}\varepsilon_j$  which means  $e_i$  is a good approximation to  $\varepsilon$  when entries of  $h_{ij}$  of H are small (which is "usually" the case, especially when n is large).

$$e_i \sim \mathcal{N}(0, \sigma^2(1 - h_{ii})) \iff \frac{e_i - 0}{\sigma \sqrt{1 - h_{ii}}} \sim \mathcal{N}(0, 1)$$

If we plug in  $\hat{\sigma}$ , that defines the studentized residuals.

$$d_i \equiv \frac{e_i}{\hat{\sigma}\sqrt{1 - h_{ii}}}$$

Common practice is to use e for the following residual plots/diagnostics, (using  $d_i$  is also possible) to check model assumptions:

Plot e versus  $\hat{\mu}$  which was shown in A2 were mutually independent since they are multivariate normal with covariance 0.

Typical "good" scatter plot will have a random scatter around y = 0 (no visible patterns).

Problematic scatter plot is when variance of  $e_i$  is not constant (cone) increases with fitted values.

In general, plot of e and  $\hat{\mu}$  can show deviations from independence, constant variance if those assumptions are violated.

Plot e versus  $x_j$  for each  $j=1,\ldots,p$  in model when not many predictors. This can help detect non-linearity between  $x_j$  and y, not as practical when p is large.

Typical "good" scatter plot will have a random scatter around y = 0 (no visible patterns).

If the observation numbers were collected in some order (time, space, etc.), also plot  $e_i$  versus indices i to check for any patterns (again, look for random scatter)

Histogram of e: is it bell shaped and symmetric to assess Normal assumption, but the histogram can't easily detect overly fat/thing tails.

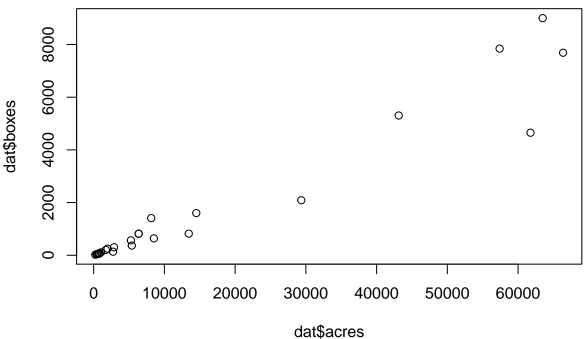
Q-Q plot of e more formally assess Normality: scatter plot of ordered quantiles from 2 distributions. In our case: empirical quantiles from residuals (data) versus theoretical quantiles from assumed Normal distribution. If quantiles roughly match, the points will roughly fall on the  $45^{\circ}$  line through origin.

If these sets of residuals plots show violations, then that could affect the validity of confidence intervals, hypothesis tests, etc.

#### 2.15.1 R Demo

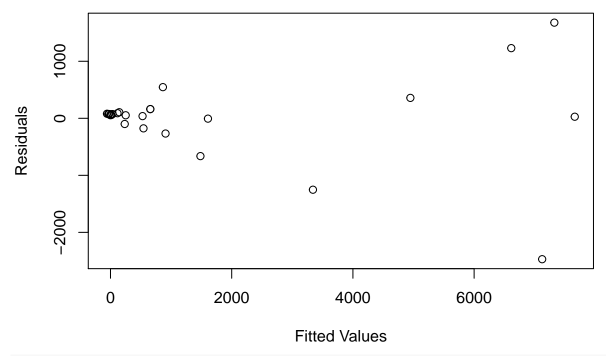
```
### Residual plots/diagnostics demo.

## Florida oranges revisited.
dat <- read.csv("csv/florange.csv")
plot(dat$acres, dat$boxes)</pre>
```

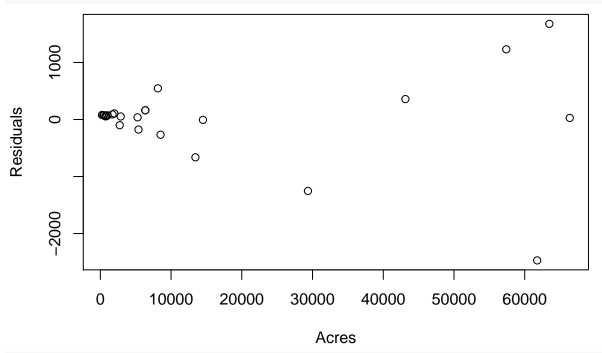


```
lm.1 <- lm(dat$boxes ~ dat$acres)
summary(lm.1)</pre>
```

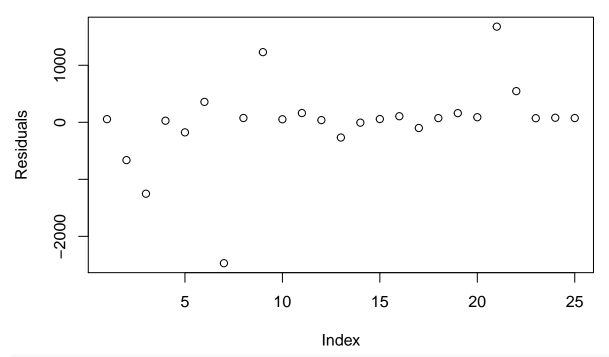
```
##
## Call:
## lm(formula = dat$boxes ~ dat$acres)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                    30
                        71.72
## -2470.81
               -6.17
                                106.46 1677.32
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) -85.391989 186.178031 -0.459
                            0.006761 17.263 1.16e-14 ***
## dat$acres
                 0.116717
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 754.4 on 23 degrees of freedom
## Multiple R-squared: 0.9284, Adjusted R-squared: 0.9252
## F-statistic:
                 298 on 1 and 23 DF, p-value: 1.164e-14
# Residual plot: vs fitted values.
plot(lm.1$fitted.values,
     lm.1$residuals,
     xlab = "Fitted Values",
    ylab = "Residuals")
```



# Residual plot: vs predictor (just one in this case).
plot(dat\$acres, lm.1\$residuals, xlab = "Acres", ylab = "Residuals")

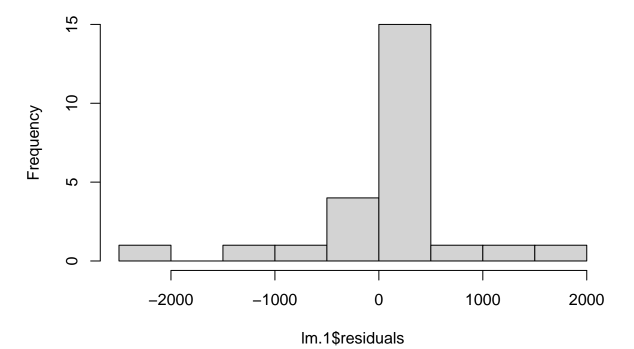


# Residual plot: vs i (just to demo plot; no time/space ordering here).
plot(1:nrow(dat), lm.1\$residuals, xlab = "Index", ylab = "Residuals")

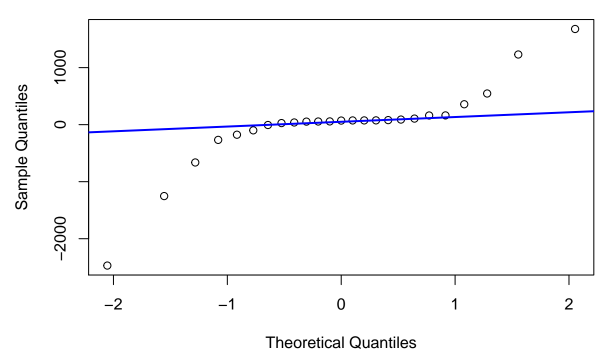


# Histogram of residuals.
hist(lm.1\$residuals)

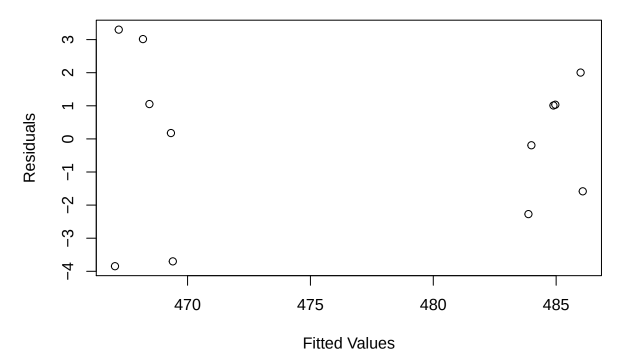
# Histogram of Im.1\$residuals



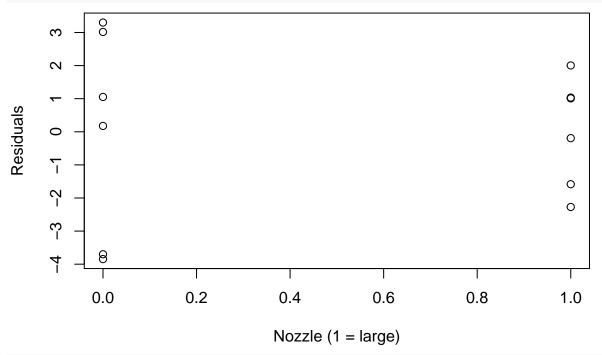
### Normal Q-Q Plot



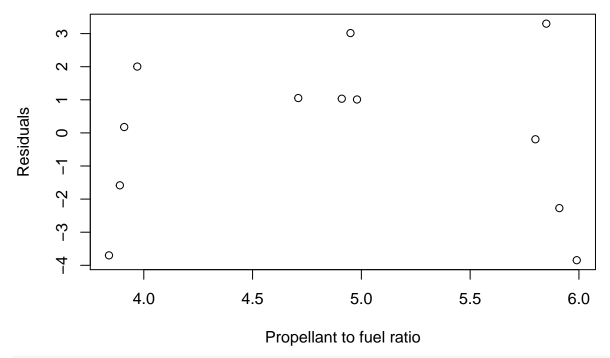
```
## Rocket data revisited.
rocket <- read.csv("csv/rocket.csv")</pre>
mr <- lm(thrust ~ nozzle + propratio, data = rocket)</pre>
summary(mr)
##
## Call:
## lm(formula = thrust ~ nozzle + propratio, data = rocket)
##
## Residuals:
       Min
                1Q Median
                                3Q
                                       Max
## -3.8459 -1.7555 0.5934 1.2906 3.3008
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 473.6039
                            4.7158 100.430 4.88e-15 ***
## nozzle
                16.7383
                            1.5329 10.919 1.71e-06 ***
## propratio
                -1.0948
                            0.9414 -1.163
                                              0.275
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2.655 on 9 degrees of freedom
## Multiple R-squared: 0.9303, Adjusted R-squared: 0.9148
## F-statistic: 60.05 on 2 and 9 DF, p-value: 6.238e-06
# Residual plot: vs fitted values.
plot(mr$fitted.values,
     mr$residuals,
     xlab = "Fitted Values",
    ylab = "Residuals")
```



```
# Residual plot: vs predictors.
plot(rocket$nozzle, mr$residuals, xlab = "Nozzle (1 = large)",
    ylab = "Residuals")
```

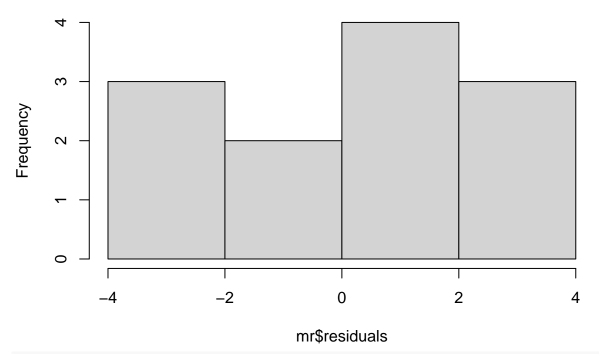


```
plot(rocket$propratio,
    mr$residuals,
    xlab = "Propellant to fuel ratio",
    ylab = "Residuals")
```

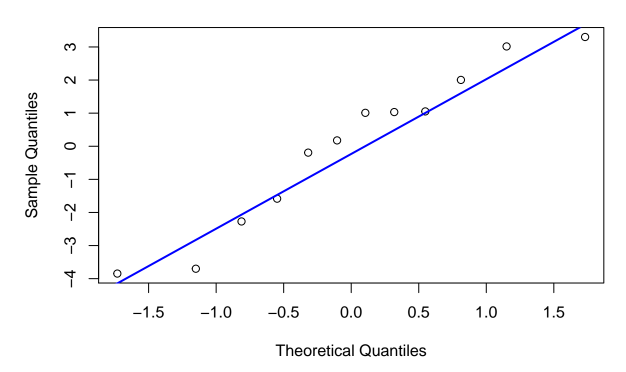


# Histogram of residuals,
hist(mr\$residuals)

# Histogram of mr\$residuals



```
# QQ plot of residuals,
qqnorm(mr$residuals)
qqline(mr$residuals, col = "blue", lwd = 2)
```



## LECTURE 16 | 2020-11-04

## 2.16 Addressing Problems With Regression Model Assumptions

If residual plots reveal problems with assumptions (although plots don't fully check linearity, independence), we might be able to address via  $\underline{\text{transformations}}$ , adding variables to model, using different error distribution on  $\varepsilon$ 

(1) Variance-stabilizing transformations on responses  $y_i$ , can help constant variance identified on e vs  $\hat{\mu}$  plot.

<u>Idea</u>: apply function  $g(\cdot)$  and fit regression on transformed  $g(y_i)$ ; that is,

$$g(Y_i) = \beta_0 + \beta_1 x_{i1} + \dots + \beta_p x_{ip} + \varepsilon_i$$

Note this can drastically change SS(Res) and  $\hat{\sigma}$ , so cannot directly use those among different choices  $g(\cdot)$ .

<u>Rationale</u>: variance of response might be a function of mean  $\mu_i = \mathbb{E}[Y_i]$ ; could be expressed

$$\mathbb{V}(Y_i) = \mathbb{V}(\varepsilon_i) = h(\mu_i)\sigma^2$$

for some  $h(\cdot) > 0$ . In which case, we want

$$\mathbb{V}(g(Y_i)) \approx \sigma^2$$

1st order Taylor expansion

$$g(Y_i) \approx g(\mu_i) + (Y_i - \mu_i)g'(\mu_i)$$
$$\mathbb{V}(g(Y_i)) \approx [g'(\mu_i)]^2 \mathbb{V}(Y_i)$$

Thus, for  $\mathbb{V}(g(Y_i))$  to be constant, we need

$$[g'(\mu_i)]^2 \propto \frac{1}{h(\mu_i)}$$

Examples:

(i)  $h(\mu_i)=\mu_i$ ; that is,  $\mathbb{V}(Y_i)=\sigma^2\mu_i\propto\mu_i$ . Variance in responses proportional to mean response. We need

$$g'(\mu_i) \propto \frac{1}{\sqrt{h(\mu_i)}} = \frac{1}{\sqrt{\mu_i}}$$

and so  $g(\mu_i)=\sqrt{\mu_i}$  works, and we apply  $g(y_i)=\sqrt{y_i}$  to obtain approximately constant variance

(ii)  $h(\mu_i) = \mu_i^2$ ; that is,  $\mathbb{V}(Y_i) = \sigma^2 \mu_i^2 \propto \mu_i^2$  or  $Sd(Y_i) \propto \mu_i$ . We need

$$g'(\mu_i) \propto \frac{1}{\mu_i}$$

and so  $g(\mu_i) = \ln(\mu_i)$  works, and we apply  $g(y_i) = \ln(y_i)$  to stabilize variance.

(iii) Class of power transformations (Box-Cox)

$$g(y_i) = \begin{cases} \frac{y_i^{\lambda} - 1}{\lambda} & \lambda \neq 0 \\ \ln(y_i) & \lambda = 0 \end{cases}$$

$$g'(\mu_i) = \begin{cases} \mu_i^{\lambda-1} & \lambda \neq 0 \\ \frac{1}{\mu_i} & \lambda = 0 \end{cases} \iff h(\mu_i) \propto \frac{1}{[g'(\mu_i)]^2} = \mu_i^C \quad C \in \mathbf{R}$$

Box-Cox transformation can help address non-constant variance of the form

$$\mu_i^C \sigma^2 = \mathbb{V}(Y_i)$$

Special cases include:

- $\lambda = \frac{1}{2}$  is  $\sqrt{\cdot}$
- $\lambda = 0$  is  $\ln(\cdot)$
- $\lambda = 1$  is identity
- $\lambda = -1$  is reciprocal

can automatically try a sequence of  $\lambda$  and find the choice that gives the best value of likelihood

Note that interpreting  $\hat{\beta}_j$  can be less intuitive as a result of transformation, since now increasing  $x_j$  by 1 unit corresponds to an estimated change of  $\hat{\beta}_j$  in  $g(y_i)$ . For  $g(y_i) = \ln(y_i)$ ,  $\hat{\beta}_j$  represents estimate of expected change in  $\ln(y_i)$  which corresponds to  $e^{\hat{\beta}_j}$  being the expected multiplicative change applied to the (original) response. But for an arbitrary  $\lambda$ , the transformation might be less interpretable.

- (2) Transforming and/or adding explanatory variables.
  - If y (or g(y)) has a clear non-linear relationship with some  $x_j$ , we can consider transforming  $x_j$  (e.g.,  $\ln(\cdot)$ ,  $\sqrt{\cdot}$ , etc.)
  - Could add polynomial terms (e.g.,  $x^2, x^3, ...$ ). For example,

$$Y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \varepsilon_i$$

suppose we think adding  $x_1^2$  is appropriate, then we define  $x_{i3} = x_{i1}^2$  and fit

$$Y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$$

which is still <u>linear</u> in  $\beta$  and note that  $x_1$  and  $x_1^2$  are linearly independent.

• Add interaction terms: if we think the effect of  $x_i$  on response depends on the value of  $x_j$ , e.g., suppose we think  $x_1$  and  $x_2$  interact, then we might fit

$$Y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \varepsilon_i$$

where  $x_{i3} = x_{i1} + x_{i2}$ , so that

$$Y_{i} = \beta_{0} + (\beta_{1} + \beta_{3}x_{i2})x_{i1} + \beta_{2}x_{i2} + \varepsilon_{i}$$

Note: in general, there's  $\binom{p}{2}$  possible interactions, consider whether interactions are conceptually plausible.

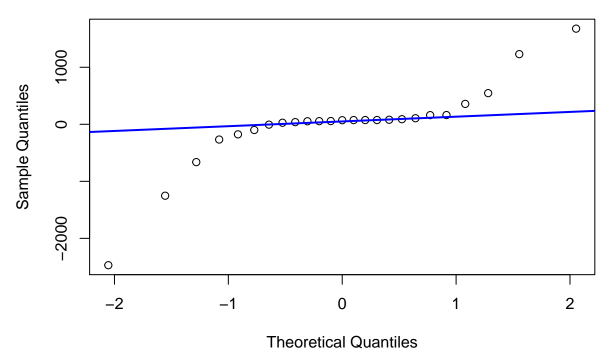
- (3) Q-Q plot of residuals not normal (even after any appropriate transformations)
  - Consider using different distribution for error term  $\varepsilon$  (e.g., t, Cauchy, Laplace, etc.)

#### 2.16.1 R Demo

```
## Demo for transformations and interactions
## Florida oranges revisited
dat <- read.csv("csv/florange.csv")</pre>
lm.1 <- lm(dat$boxes ~ dat$acres)</pre>
summary(lm.1)
##
## Call:
## lm(formula = dat$boxes ~ dat$acres)
## Residuals:
        Min
                  1Q
                       Median
                                     ЗQ
## -2470.81
                        71.72
               -6.17
                                106.46 1677.32
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) -85.391989 186.178031 -0.459
                                                 0.651
## dat$acres
                 0.116717
                            0.006761 17.263 1.16e-14 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 754.4 on 23 degrees of freedom
## Multiple R-squared: 0.9284, Adjusted R-squared: 0.9252
## F-statistic: 298 on 1 and 23 DF, p-value: 1.164e-14
# Recall: residuals had non-constant variance
# (variance increases with fitted values)
plot(lm.1$fitted.values,
     lm.1$residuals,
     xlab = "Fitted Values",
     ylab = "Residuals")
                                                                              0
     1000
                                                                        0
                     0
                                                         0
Residuals
                                                                                 0
     0
                           0
                          0
                                           0
                                                                            0
             0
                             2000
                                               4000
                                                                 6000
```

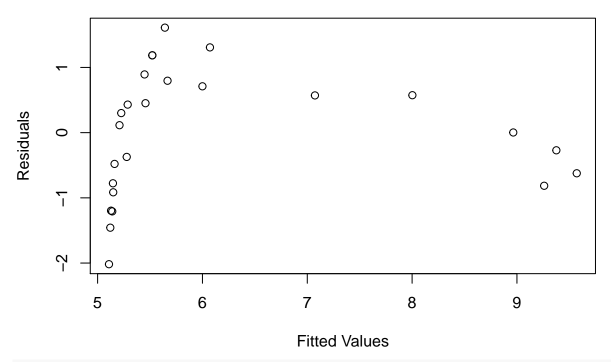
Fitted Values

```
qqnorm(lm.1$residuals)
qqline(lm.1$residuals, col = "blue", lwd = 2)
```

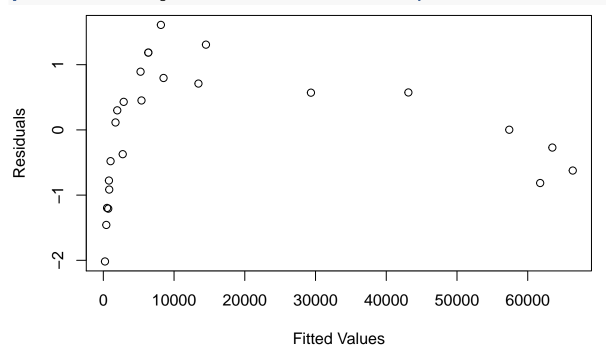


# Try log-transforming y
lm.log <- lm(log(dat\$boxes) ~ dat\$acres)
summary(lm.log)</pre>

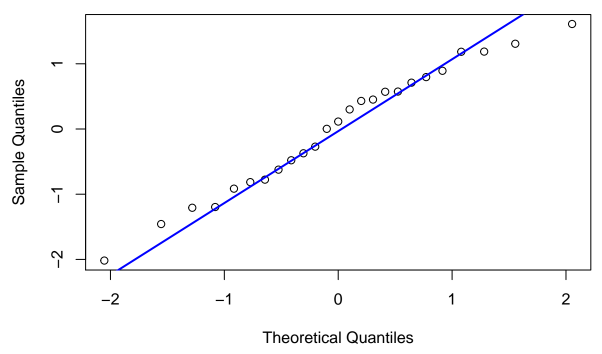
```
##
## Call:
## lm(formula = log(dat$boxes) ~ dat$acres)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -2.0175 -0.7767 0.1142 0.7106 1.6102
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 5.093e+00 2.425e-01 20.997 < 2e-16 ***
## dat$acres
                                   7.661 8.95e-08 ***
              6.748e-05 8.808e-06
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.9828 on 23 degrees of freedom
## Multiple R-squared: 0.7184, Adjusted R-squared: 0.7062
## F-statistic: 58.69 on 1 and 23 DF, p-value: 8.948e-08
plot(lm.log$fitted.values,
     lm.log$residuals,
     xlab = "Fitted Values",
    ylab = "Residuals")
```



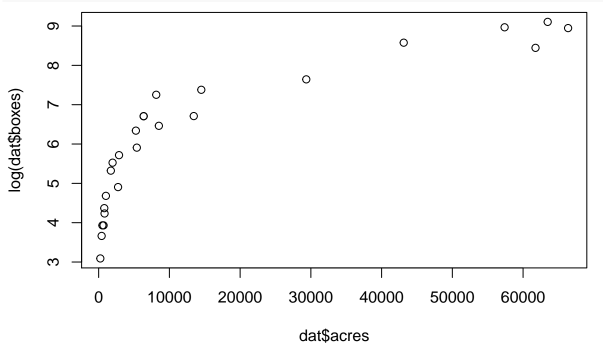
plot(dat\$acres, lm.log\$residuals, xlab = "Fitted Values", ylab = "Residuals")



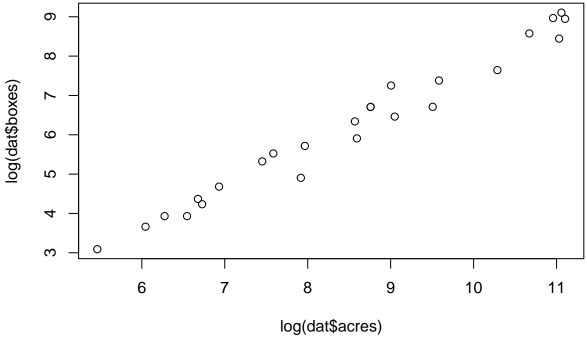
```
qqnorm(lm.log$residuals)
qqline(lm.log$residuals, col = "blue", lwd = 2)
```



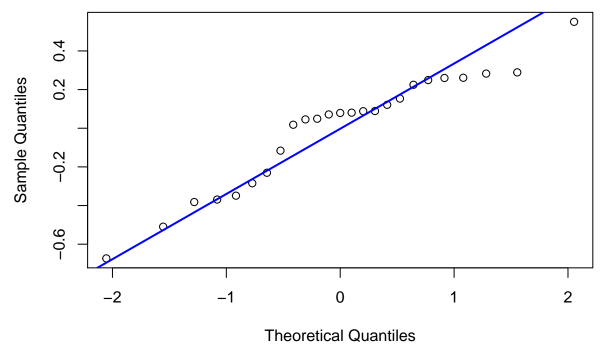
# Does the plot of residuals vs x suggest a problem
# Let's take a closer look
plot(dat\$acres, log(dat\$boxes)) # evidently not linear!



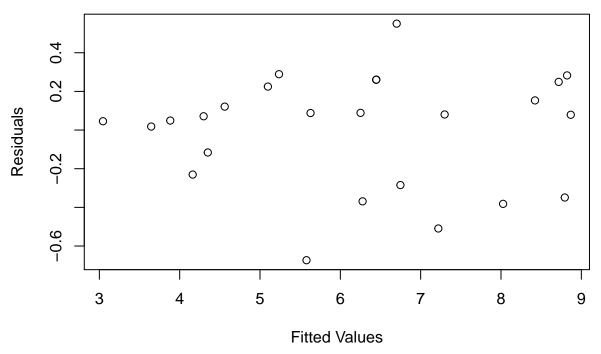
# Log-transform x as well
plot(log(dat\$acres), log(dat\$boxes)) # looks much more linear!



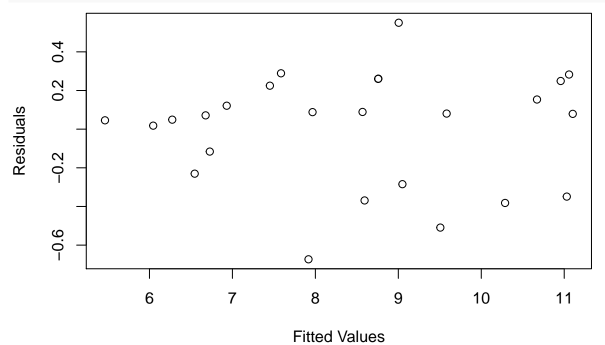
```
lm.loglog <- lm(log(dat$boxes) ~ log(dat$acres))
qqnorm(lm.loglog$residuals)
qqline(lm.loglog$residuals, col = "blue", lwd = 2)</pre>
```



```
plot(lm.loglog$fitted.values,
    lm.loglog$residuals,
    xlab = "Fitted Values",
    ylab = "Residuals")
```



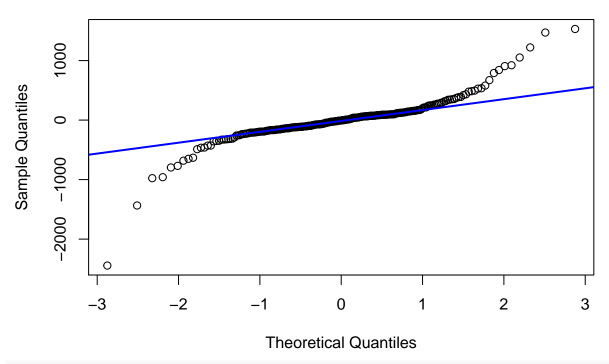
```
plot(log(dat$acres),
    lm.loglog$residuals,
    xlab = "Fitted Values",
    ylab = "Residuals")
```



```
## Python data revisited
python <- read.csv("csv/FLpython.csv")
python$male <- ifelse(python$sex == 'M', 1, 0) # 1 = M, 0 =F
mpf2 <- lm(fat ~ male + mass + svl, data = python)
summary(mpf2)</pre>
```

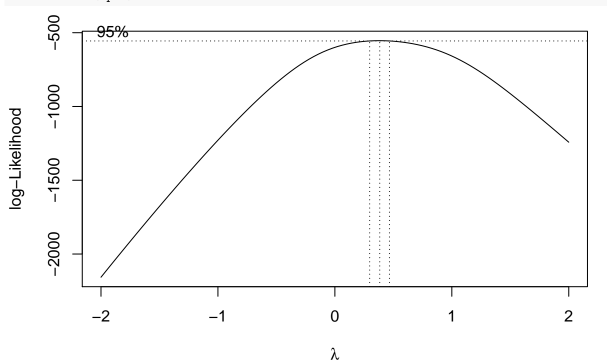
```
##
## Call:
## lm(formula = fat ~ male + mass + svl, data = python)
## Residuals:
##
        Min
                       Median
                                     3Q
                  1Q
                                             Max
   -2444.44 -137.38
                        -6.66
                                109.22 1530.81
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 204.09840
                          132.30121
                                        1.543
                                                0.1242
               -196.71705
                            47.16396
                                      -4.171 4.22e-05 ***
## male
                             0.00524 22.495 < 2e-16 ***
## mass
                  0.11788
                                                0.0375 *
## svl
                 -1.59841
                             0.76433
                                      -2.091
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 360.2 on 244 degrees of freedom
## Multiple R-squared: 0.897, Adjusted R-squared: 0.8957
## F-statistic: 708.2 on 3 and 244 DF, p-value: < 2.2e-16
# Residual plot: vs fitted values
plot(mpf2$fitted.values,
     mpf2$residuals,
     xlab = "Fitted Values",
     ylab = "Residuals")
                                     0
                                                         0
     1000
                                                  0
                                         0
                                         0
                                                                                  0
Residuals
     0
                                     0
                                           0
     -2000 -1000
                                                    0
                                               0
                                                               0
                                                          0
             0
                            2000
                                                             6000
                                                                             8000
                                            4000
                                         Fitted Values
## QQ plot of residuals
qqnorm(mpf2$residuals)
```

qqline(mpf2\$residuals, col = "blue", lwd = 2)



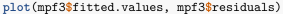
## library(MASS)

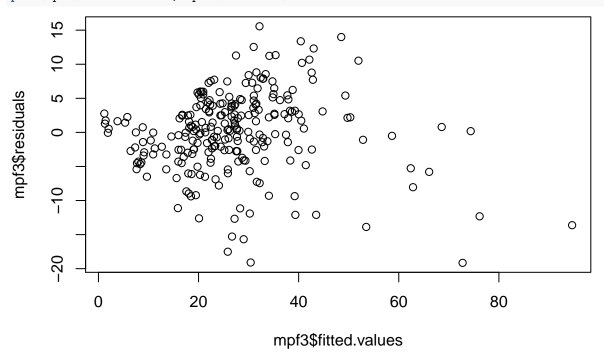
# Try a Box-Cox transformation
bc <- boxcox(mpf2)</pre>



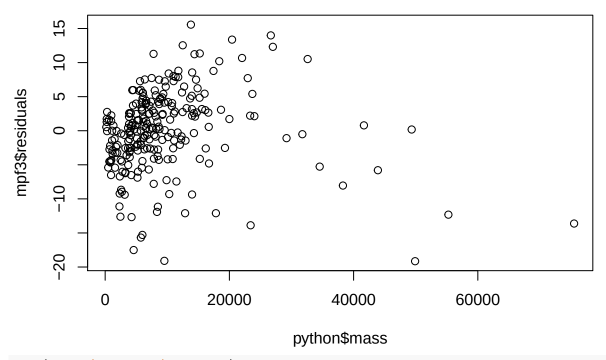
```
lambda <- bc$x[which.max(bc$y)]
mpf3 <- lm((fat ^ lambda - 1) / lambda ~ male + mass + svl, data = python)
summary(mpf3)</pre>
```

```
##
## lm(formula = (fat^lambda - 1)/lambda ~ male + mass + svl, data = python)
##
## Residuals:
##
       Min
                1Q
                    Median
                                3Q
                                       Max
  -19.146 -2.910
                     0.297
                             3.688
                                    15.568
##
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                                      -3.693 0.000273 ***
## (Intercept) -8.0558134
                          2.1813183
##
               -1.7849310
                          0.7776166
                                      -2.295 0.022560 *
## mass
                0.0004461
                           0.0000864
                                       5.164 5.03e-07 ***
                0.1431492
                           0.0126019
                                      11.359 < 2e-16 ***
## svl
##
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## Residual standard error: 5.939 on 244 degrees of freedom
## Multiple R-squared: 0.8356, Adjusted R-squared: 0.8336
## F-statistic: 413.5 on 3 and 244 DF, p-value: < 2.2e-16
```

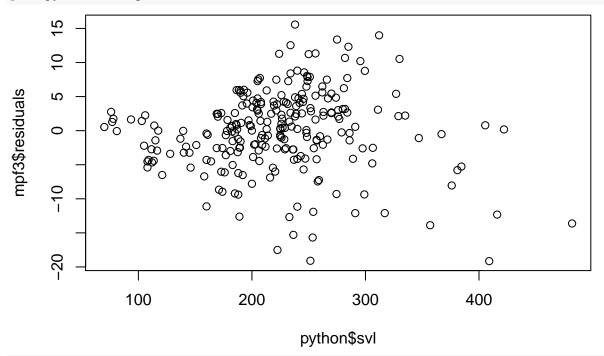




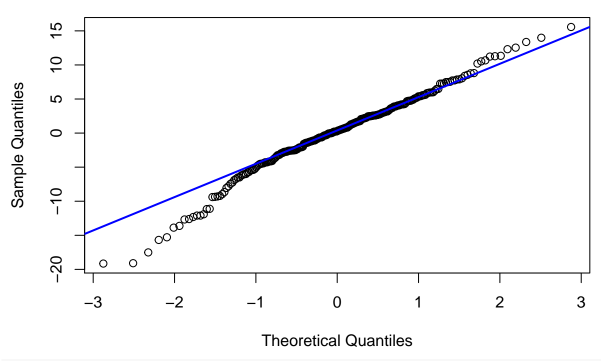
plot(python\$mass, mpf3\$residuals)







```
qqnorm(mpf3$residuals)
qqline(mpf3$residuals, col = "blue", lwd = 2)
```



# still some skew, but better!

## LECTURE 17 | 2020-11-09

## 2.17 Effects of Individual Observations

- (1) We say an observation  $(y_i, x_{i1}, \dots, x_{ip})$  is an outlier if it is substantially different from other observations. This can occur if its response and/or some of its explanatory variables have values that are unusual or extreme compared to the others. Outliers can occur for different reasons. For example, an extraordinary subject, data entry errors. Generally, we don't recommend removing outliers unless we have a strong reason to believe that observation is an error. But it can be useful to investigate what effect it has on our fitted model and quality of fit to the rest of data.
- (2) How to detect and characterize outliers: Studentized residual

$$d_i = \frac{e_i}{\hat{\sigma}\sqrt{1 - h_{ii}}}$$

are standardized to have approximately variance 1 where

$$e_i \sim \mathcal{N}(0, \sigma^2(1-h_{ii}))$$

So, if  $|d_i|$  is large, observation i could be considered an outlier in the sense of having an extreme value of response  $y_i$  (e.g.,  $|d_i| > 3$ )

(3) Recall that  $h_{ii}$  is the  $i^{th}$  diagonal element of hat matrix H. We call  $h_{ii}$  the **leverage** of observation i.

$$\hat{\boldsymbol{\mu}} = X\hat{\boldsymbol{\beta}} = H\boldsymbol{Y}$$

so

$$\hat{\mu}_i = \begin{bmatrix} h_{i1} & h_{i2} & \cdots & h_{in} \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix} = \sum_{j=1}^n h_{ij} y_j = h_{ii} y_i + \sum_{j \neq i} h_{ij} y_j$$

so  $h_{ii}$  captures contribution of  $Y_i$  in determining its corresponding fitted value. If the leverage is large relative to  $h_{ij}$ 's then  $\hat{\mu}_i$  is mostly determined by  $Y_i$ . In A3, you will show the leverage is between  $\frac{1}{n}$  and 1; that is,

$$\frac{1}{n} \le h_{ii} \le 1$$

If  $h_{ii}\approx 1$ , then  $\mathbb{V}(e_i)=\sigma^2(1-h_{ii})\approx 0$ , which in turn implies  $y_i\approx \hat{\mu}_i$ ; that is, residuals with observations with high leverage tend to be small.

Rule of thumb: an observation with leverage higher than twice the average leverage is considered high.

$$h_{ii} > 2\bar{h}$$

where 
$$\bar{h} = \frac{1}{n} \sum_{i=1}^{n} h_{ii}$$
.

$$tr(H) = rank X = p + 1$$

Recall that  $H = X(X^{T}X)^{-1}X^{T}$  only involves predictors. Thus, leverage is useful to help identify outliers n the sense of having explanatory variables with extreme or unusual values. In simple linear regression,

$$h_{ii} = \frac{1}{n} + \frac{(x_i - \bar{x})^2}{S_{xx}}$$

Which means if  $x_i$  is far from  $\bar{x}$ , that point will have high leverage. This generalizes to multiple linear regression: an observation with high leverage is an outlier with extreme values in one or more explanatory variables. However, leverage does not tell us directly whether that observation is also an outlier in response, in fact,  $y_i \approx \hat{\mu}_i$  for such observations with high leverage.

(4) An observation i is quite influential if its presence in fitting regression considerably changes estimates compared to when observation i is not used to fit a model.

Start with fitting  $Y = X\beta + \varepsilon$  using all observations and call estimates  $\hat{\beta}$  as usual in the least squares.

Let  $\hat{\beta}^{(i)}$  denote the least squares estimates based on fitting a model with the  $i^{th}$  observation removed.

<u>Idea</u>: If  $\hat{\beta}^{(i)}$  is quite different from  $\hat{\beta}$ , then observation i is highly influential. Measure this via **Cook's distance** between  $\hat{\beta}$  and  $\hat{\beta}^{(i)}$ :

$$D_i = \frac{(\hat{\beta}^{(i)} - \hat{\beta})^\top X^\top X (\hat{\beta}^{(i)} - \hat{\beta})}{\hat{\sigma}^2(p+1)}$$

To see this intuition, let  $\hat{\mu}^{(i)} = X \hat{\beta}^{(i)}$  fitted values based on removing  $i^{\text{th}}$  observation and estimating  $\beta$ . Then,

$$D_i = \frac{(X\hat{\pmb{\beta}}^{(i)} - X\hat{\pmb{\beta}})^\top (X\hat{\pmb{\beta}}^{(i)} - X\hat{\pmb{\beta}})}{\hat{\sigma}^2(p+1)} = \frac{(\hat{\pmb{\mu}}^{(i)} - \hat{\pmb{\mu}})^\top (\hat{\pmb{\mu}}^{(i)} - \hat{\pmb{\mu}})}{\hat{\sigma}^2(p+1)} = \frac{\|\hat{\pmb{\mu}}^{(i)} - \hat{\pmb{\mu}}\|^2}{\hat{\sigma}^2(p+1)}$$

 $D_i$  measures the Euclidean distance between fitted values of two regressions that give  $\hat{\mu}^{(i)}$  and  $\hat{\mu}$ , up to a scaling factor. Further, it can be shown

$$D_i = d_i^2 \bigg(\frac{h_{ii}}{1-h_{ii}}\bigg) \bigg(\frac{1}{p+1}\bigg)$$

where we can see that both  $d_i$  and  $h_{ii}$  are key quantities to calculate  $D_i$  and observation i that are most influential will have larges of  $|d_i|$  and  $h_{ii}$ .

- High  $|d_i| \to \text{outlier}$  in response
- High  $h_{ii} \rightarrow$  outlier in explanatory variable

So high influential observations tend to be outliers in the sense of having extreme values in *both* response and one/more predictors. Check if some  $D_i$ 's are much larger than others, as they would be the most influential observations.

## LECTURE 18 | 2020-11-11

For the  $i^{th}$  observation, the values of its predictors are

$$\begin{aligned} \boldsymbol{v}_i &= \begin{bmatrix} 1 & x_{i1} & x_{i2} & \cdots & x_{ip} \end{bmatrix}^\top \\ & X &= \begin{bmatrix} \boldsymbol{v}_1^\top \\ \boldsymbol{v}_2^\top \\ \vdots \\ \boldsymbol{v}_n^\top \end{bmatrix} \\ & X^\top X &= \begin{bmatrix} \boldsymbol{v}_1 & \cdots & \boldsymbol{v}_n \end{bmatrix} \begin{bmatrix} \boldsymbol{v}_1^\top \\ \vdots \\ \boldsymbol{v}_n^\top \end{bmatrix} = \sum_{i=1}^n \boldsymbol{v}_j \boldsymbol{v}_j^\top \end{aligned}$$

Recall  $H = X(X^{\top}X)^{-1}X^{\top}$  so

$$h_{ii} = \boldsymbol{v}_i^\top (X^\top X)^{-1} \boldsymbol{v}_i^\top$$

Let  $X^{(i)}$  be X with  $i^{\text{th}}$  observation deleted,  $\boldsymbol{y}^{(i)}$  be  $\boldsymbol{y}$  with  $i^{\text{th}}$  response deleted. Then,

$$(X^{(i)})^\top X^{(i)} = \sum_{j \neq i} \boldsymbol{v}_j \boldsymbol{v}_j^\top \implies X^\top X = (X^{(i)})^\top X^{(i)} + \boldsymbol{v}_i \boldsymbol{v}_i^\top$$

Similarly,

$$X^{\top}\boldsymbol{y} = \sum_{j=1}^{n} \boldsymbol{v}_{j} y_{j} \implies (X^{(i)})^{\top} \boldsymbol{y}^{(i)} = \sum_{j \neq i} \boldsymbol{v}_{j} y_{j} \implies X^{\top} \boldsymbol{y} = (X^{(i)})^{\top} \boldsymbol{y}^{(i)} + \boldsymbol{v}_{i} y_{i}$$

Let A be an  $n \times n$  invertible matrix

$$(A - aa^{\top})^{-1} = A^{-1} + \frac{A^{-1}aa^{\top}A^{-1}}{1 - a^{\top}A^{-1}a}$$

Fit the least squares using  $X^{(i)}$  and  $\boldsymbol{y}^{(i)}$  to obtain

$$\begin{split} \hat{\boldsymbol{\beta}}^{(i)} &= [(\boldsymbol{X}^{(i)})^{\intercal}\boldsymbol{X}^{(i)}]^{-1}(\boldsymbol{X}^{(i)})^{\intercal}\boldsymbol{y}^{(i)} \\ &= (\boldsymbol{X}^{\intercal}\boldsymbol{X} - \boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal})^{-1}(\boldsymbol{X}^{\intercal}\boldsymbol{y} - \boldsymbol{v}_{i}\boldsymbol{y}_{i}) \\ &= \left[ (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1} + \frac{(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}}{1 - \boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}} \right] (\boldsymbol{X}^{\intercal}\boldsymbol{y} - \boldsymbol{v}_{i}\boldsymbol{y}_{i}) \\ &= (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{X}^{\intercal}\boldsymbol{y} - (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i} + \frac{(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{X}^{\intercal}\boldsymbol{y} - (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}(\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\boldsymbol{v}_{i}^{\intercal}\boldsymbol{\beta} - \boldsymbol{h}_{ii}\boldsymbol{y}_{i}} \\ &= \hat{\boldsymbol{\beta}} - (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\left[ \frac{\boldsymbol{y}_{i} - \boldsymbol{h}_{ii}\boldsymbol{y}_{i} - \boldsymbol{v}_{i}^{\intercal}\hat{\boldsymbol{\beta}} + \boldsymbol{h}_{ii}\boldsymbol{y}_{i}}{1 - \boldsymbol{h}_{ii}} \right] \\ &= \hat{\boldsymbol{\beta}} - (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\left[ \frac{\boldsymbol{y}_{i} - \boldsymbol{v}_{i}^{\intercal}\hat{\boldsymbol{\beta}}}{1 - \boldsymbol{h}_{ii}} \right] \\ &= \hat{\boldsymbol{\beta}} - (\boldsymbol{X}^{\intercal}\boldsymbol{X})^{-1}\boldsymbol{v}_{i}\left[ \frac{\boldsymbol{e}_{i}}{1 - \boldsymbol{h}_{ii}} \right] \end{split}{1 - \boldsymbol{h}_{ii}}$$

Therefore,

$$\hat{\boldsymbol{\beta}}^{(i)} - \hat{\boldsymbol{\beta}} = \frac{-e_i}{1 - h_{ii}} (\boldsymbol{X}^\top \boldsymbol{X})^{-1} \boldsymbol{v}_i$$

In fact, we can get  $\hat{\beta}^{(i)}$  from the regression with all n observations; that is, we don't need to fit a separate model.

$$\begin{split} D_i &= \frac{(\hat{\beta}^{(i)} - \hat{\beta})^\top X^\top X (\hat{\beta}^{(i)} - \hat{\beta})}{\hat{\sigma}^2(p+1)} \\ &= \frac{\left(\frac{-e_i}{1-h_{ii}}\right) \boldsymbol{v}_i^\top (X^\top X)^{-1} X^\top X \left(\frac{-e_i}{1-h_{ii}}\right) (X^\top X)^{-1} \boldsymbol{v}_i}{\hat{\sigma}^2(p+1)} \\ &= \frac{e_i^2}{\hat{\sigma}^2(1-h_{ii})(1-h_{ii})} \left(\frac{h_{ii}}{p+1}\right) \\ &= d_i^2 \left(\frac{h_{ii}}{1-h_{ii}}\right) \left(\frac{1}{p+1}\right) \end{split}$$

as claimed, so we can calculate Cook's distance in terms of  $|d_i|$  and  $h_{ii}$ . Therefore, the most influential observation on estimates of  $\beta$  are those with high  $|d_i|$  and  $h_{ii}$ .

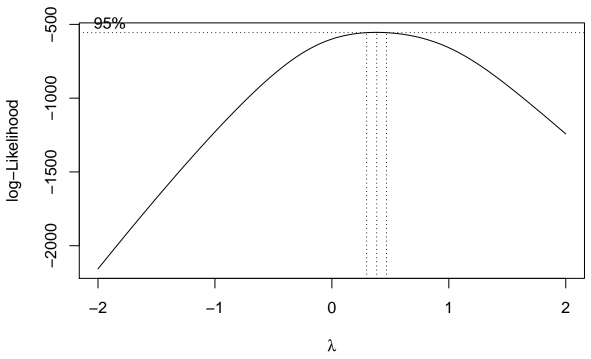
#### 2.17.1 R Demo

```
## Effect of individual observations

## Python data revisited
python <- read.csv("csv/FLpython.csv")
python$male <- ifelse(python$sex == 'M', 1, 0) # 1 = M, 0 =F

mpf2 <- lm(fat ~ male + mass + svl, data = python)

# Last time we used a Box-Cox transformation
library(MASS)
bc <- boxcox(mpf2)</pre>
```

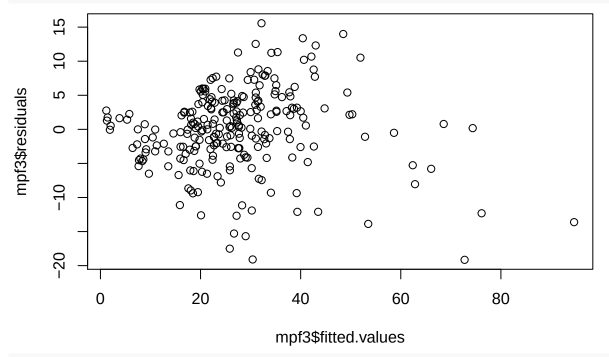


```
lambda <- bc$x[which.max(bc$y)]
mpf3 <- lm((fat ^ lambda - 1) / lambda ~ male + mass + svl, data = python)
summary(mpf3)</pre>
```

```
##
## Call:
## lm(formula = (fat^lambda - 1)/lambda ~ male + mass + svl, data = python)
## Residuals:
      Min
               1Q Median
                               3Q
## -19.146 -2.910
                    0.297
                            3.688 15.568
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -8.0558134 2.1813183 -3.693 0.000273 ***
              -1.7849310 0.7776166 -2.295 0.022560 *
## male
               0.0004461 0.0000864
                                     5.164 5.03e-07 ***
## mass
               0.1431492  0.0126019  11.359  < 2e-16 ***
## svl
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

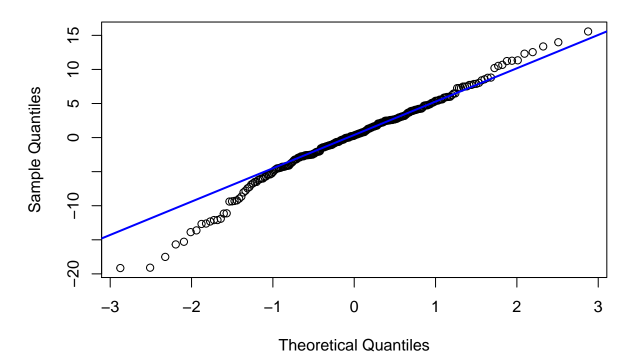
```
## Residual standard error: 5.939 on 244 degrees of freedom
## Multiple R-squared: 0.8356, Adjusted R-squared: 0.8336
## F-statistic: 413.5 on 3 and 244 DF, p-value: < 2.2e-16</pre>
```

plot(mpf3\$fitted.values, mpf3\$residuals)



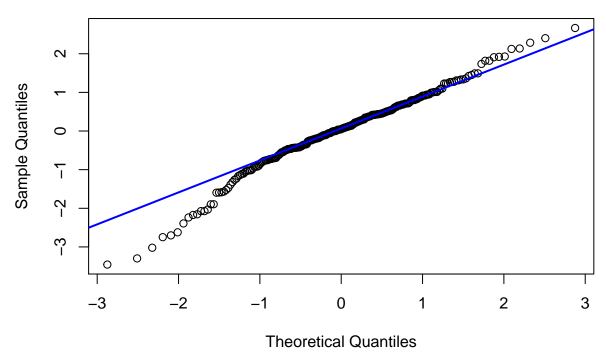
```
qqnorm(mpf3$residuals)
qqline(mpf3$residuals, col = "blue", lwd = 2)
```

# Normal Q-Q Plot

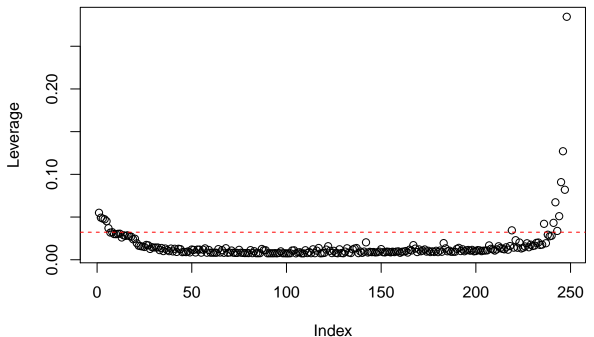


```
# Quantities for individual observations
studres(mpf3) # studentized residuals
hatvalues(mpf3) # leverage
cooks.distance(mpf3) # Cook's distance
# Residual plots with studentized residuals
plot(mpf3$fitted.values,
     studres(mpf3),
     xlab = "Fitted values",
     ylab = "Studentized residuals")
abline(h = c(3, -3), col = "red", lty = 2)
                                     0
                                                  0
      \sim
                                                    0
Studentized residuals
                                                  0
                                                   0
      0
                                                     0
                                                             0
                                           0
      7
                                           0 0
                                                                       0
                                                      0
                                 00
                                                                                     0
      က
                                    0
                                                                    0
             0
                           20
                                          40
                                                                         80
                                                          60
                                           Fitted values
which(abs(studres(mpf3)) > 3)
```

```
## 122 181 245
## 122 181 245
qqnorm(studres(mpf3))
qqline(studres(mpf3), col = "blue", lwd = 2)
```







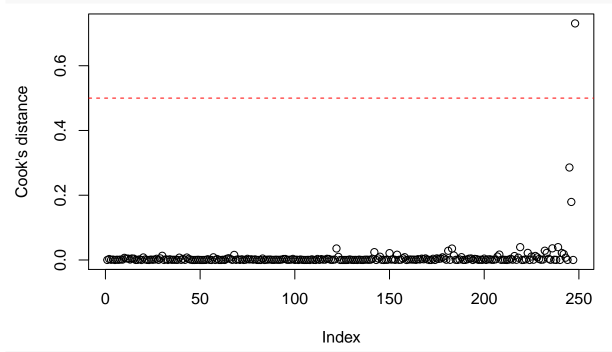
which(hatvalues(mpf3) > 2 \* mean(hatvalues(mpf3)))

## 1 2 3 4 5 6 8 219 236 241 242 243 244 245 246 247 248

```
python[which(hatvalues(mpf3) > 2 * mean(hatvalues(mpf3))), ]
```

```
mass length
##
       sex
             svl
                                     fat male
## 1
         F
            70.0
                          77.5
                                   6.000
                    186
## 2
         М
            76.0
                    310
                          83.8
                                  11.000
                                            1
## 3
         М
            77.0
                    260
                          86.1
                                   6.000
                                            1
## 4
         М
            78.0
                    262
                          87.1
                                   8.000
                                            1
                                   4.000
## 5
         M 81.0
                    306
                          91.1
                                            1
## 6
         M 93.5
                         104.6
                    605
                                  18.959
                                            1
         F 105.0
## 8
                    800
                         117.5
                                  17.000
                                            0
## 219
         M 285.0 27000
                         316.2 3230.000
## 236
         M 330.0 32600
                         370.9 4374.000
                                            1
## 241
         F 376.0 38280
                         424.2 3156.000
                                            0
         F 381.0 43910
## 242
                         424.9 4002.000
                                            0
## 243
         F 384.5 34540
                         432.4 3500.000
                                            0
## 244
         F 405.5 41660
                         455.3 5688.000
                                            0
## 245
         F 409.0 49900
                         460.2 2988.000
                                            0
## 246
         F 416.0 55260
                         469.1 4618.000
                                            0
## 247
         F 422.0 49350
                         473.4 6818.000
                                            0
         F 482.0 75500
                         545.0 8406.000
## 248
                                            0
```

# # Cook's distance plot(cooks.distance(mpf3), ylab = "Cook's distance") abline(h = 0.5, col = "red", lty = 2)



```
which(cooks.distance(mpf3) > 0.5)
```

## 248 ## 248

# Let's look at actual changes in beta estimates mpf3\$coefficients # with all the data

```
## (Intercept) male mass svl
## -8.0558134354 -1.7849310101 0.0004461197 0.1431491887
# e.g., fit without obs 248
mpf4 <-
 lm((fat ^ lambda - 1) / lambda ~ male + mass + svl, data = python[-248, ])
mpf4$coefficients
## (Intercept)
                        male
                                                    svl
                                     {\tt mass}
## -6.6475616056 -1.6605858218 0.0005743312 0.1313793189
# e.g., fit without obs 50
mpf5 <-
 lm((fat ^ lambda - 1) / lambda ~ male + mass + svl, data = python[-50, ])
mpf5$coefficients
## (Intercept)
                       \mathtt{male}
                                     mass
## -8.0628754675 -1.7805093651 0.0004462354 0.1431573753
```

## LECTURE 19 | 2020-11-16

## 2.18 Prediction Error

Sometimes a key application of a fitted model is to do <u>prediction</u> on new data (and is more important than interpretability).

Previously, we saw model selection criteria that are computed on fitted models (AIC, BIC, Adjusted  $R^2$ ) which assess the explanatory power of a model on the *data used to fit the model* (or *train*).

While these criteria incorporate penalty terms to try to prevent overfitting, they don't directly assess how well a model would perform in predicting the response on new data given predictors.

We mentioned metrics such as MSPE as measures of predictive accuracy.

To assess accuracy in prediction, we need metrics for measuring prediction error, e.g., evaluated over m observations.

#### **DEFINITION 2.18.1: Mean-squared Error (MSE)**

If we have m observations,

$$\text{MSE} = \frac{1}{m} \sum_{i=1}^{m} (y_i - \hat{y}_i)^2$$

where  $y_i$  is the actual value,  $\hat{y}_i$  is the predicted value (or equivalently, if we apply MSE on the fitted data, this would be the fitted value,  $\hat{\mu}_i$ ; if calculated on training data). Measured on the scale of  $\sigma^2$ .

#### **REMARK 2.18.2**

MSE is equivalently called MSPE if applied on new data.

#### **DEFINITION 2.18.3: Root-mean-squared Error (RMSE)**

$$RSME = \sqrt{MSE}$$

#### **REMARK 2.18.4**

RMSE is measured on the scale of  $\sigma$ .

#### **DEFINITION 2.18.5: Mean Absolute Error (MAE)**

$$\text{MAE} = \frac{1}{m} \sum_{i=1}^{m} |y_i - \hat{y}_i|$$

Ideally, we have lots of data, conceptualize having three parts.

	Validation $(y_{n+1}, \dots, y_{n+v})$	$\text{Test } (y_{n+v+1}, \dots, y_{n+v+t})$
<ul><li> n observations</li><li> Fit models, as many as we want.</li></ul>	<ul> <li>v observations</li> <li>Estimate prediction error for each fitted model.</li> </ul>	<ul> <li>t observations</li> <li>Used at the very end for final assessment of our selected model.</li> </ul>

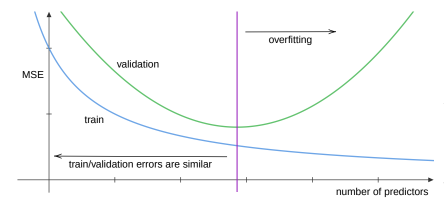
We have access to train and validation. Test data is from the future, we won't have access to this until the data is released (say the actual stock prices were released); that is, assume we don't get to see this.

For example, using MSE as a metric, based on a fitted model we can compute and compare:

MSE Train	MSE Validation	MSE Test
$\frac{1}{n}\sum_{i=1}^n(y_i-\hat{y}_i)^2$	$\frac{1}{m}\sum_{i=n+1}^{n+v}(y_i-\hat{y_i})^2$	$\frac{1}{t}\sum_{i=n+v+1}^{n+v+t}(y_i-\hat{y}_i)^2$

- Observe "MSE Train" is equal to  $\frac{SS(Res)}{n}$  and our usual estimate of  $\hat{\sigma}^2$  is a scaled version of this quantity to compensate for the number of predictors. Specifically,  $\hat{\sigma}^2 = \frac{SS(Res)}{n-p-1}$  so it's unbiased.
- Consider "MSE Validation" as an estimate of MSPE on new data.
- "MSE Test" is the actual test of prediction, we call this the actual MSPE.

<u>Idea</u>: We hope that "MSE Validation"  $\approx$  "MSE Test" since neither sets were used to fit the model.



So, if we're using MSE/RMSE as a metric (as related to  $\hat{\sigma}^2/\hat{\sigma}$ ) and the MSE/RMSE is significantly larger on validation compared to train, then we probably overfitted the model; that is, we can't expect the model to generalize well to new data.

## 2.19 Cross-Validation

How to use framework in practice:

- Simplest: randomly divide available data between train/validation, say 80%/20% split.

Weakness:

- 1. Don't use all data for training.
- 2. Only get one estimate of prediction error.
- Better: Use cross-validation scheme (CV). How to do CV with K folds:
  - Divide available data for train and validation into K roughly equally sized sets (folds), usually randomly.
  - For CV k, use data in fold k as validation, and train on the rest of data.
  - Thus, to estimate the prediction error for a given model, we fit it K times, each time treating the data in folds, 1, 2, ..., K as validation. Therefore, we get K estimates of prediction error for that particular model.

- For example, using RMSE, we get

$$\mathsf{RMSE}_1, \mathsf{RMSE}_2, \dots, \mathsf{RSME}_K$$

and can take the average

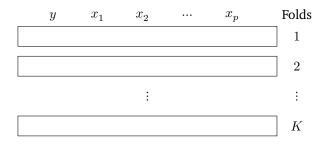
$$\overline{\text{RSME}} = \frac{1}{K} \sum_{k=1}^{K} \text{RSME}_k$$

as an estimate for RMSPE on new data (test set).

## 2.19.1 R Demo

```
## Cross-validation
## Coffee example (Coffee Quality Institute, 2018) continued
coffee <- read.csv("csv/coffee_arabica.csv")</pre>
# 1 = wet. O otherwise
coffee$wet <-</pre>
  ifelse(coffee$Processing.Method == 'Washed / Wet', 1, 0)
# 1 = semi/dry, 0 otherwise
coffee$semi <-
  ifelse(coffee$Processing.Method == 'Semi-washed / Semi-pulped', 1, 0)
coffee$Processing.Method <- NULL</pre>
N <- nrow(coffee)
## Train and validation set split
set.seed(12345678)
trainInd <- sample(1:N, round(N * 0.8), replace = F)</pre>
trainSet <- coffee[trainInd,]</pre>
validSet <- coffee[-trainInd,]</pre>
# Calculate RMSE on three models each with different variables included
m1 <-
 lm(Flavor ~ wet + semi + Aroma + Aftertaste + Body, dat = trainSet)
pred1 <- predict(m1, newdata = validSet)</pre>
sqrt(mean((validSet$Flavor - pred1) ^ 2)) # RMSE
## [1] 0.1577479
mean(abs(validSet$Flavor - pred1)) # MAE
## [1] 0.113643
m2 < -1m(
  Flavor ~ wet + Aroma + Aftertaste +
    Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
  dat = trainSet
pred2 <- predict(m2, newdata = validSet)</pre>
sqrt(mean((validSet$Flavor - pred2) ^ 2))
## [1] 0.1426565
m3 <- lm(Flavor ~ Aroma + Aftertaste, dat = trainSet)
pred3 <- predict(m3, newdata = validSet)</pre>
sqrt(mean((validSet$Flavor - pred3) ^ 2))
## [1] 0.1615385
# K fold cross validation
K <- 5
validSetSplits <- sample((1:N) %% K + 1)</pre>
RMSE1 <-c()
RMSE2 \leftarrow c()
RMSE3 <-c()
for (k in 1:K) {
validSet <- coffee[validSetSplits == k,]</pre>
```

```
trainSet <- coffee[validSetSplits != k,]</pre>
 m1 <-
   lm(Flavor ~ wet + semi + Aroma + Aftertaste + Body, dat = trainSet)
  pred1 <- predict(m1, newdata = validSet)</pre>
  RMSE1[k] <- sqrt(mean((validSet$Flavor - pred1) ^ 2))</pre>
 m2 < -1m(
   Flavor ~ wet + Aroma + Aftertaste +
      Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
    dat = trainSet
  pred2 <- predict(m2, newdata = validSet)</pre>
  RMSE2[k] <- sqrt(mean((validSet$Flavor - pred2) ^ 2))</pre>
 m3 <- lm(Flavor ~ Aroma + Aftertaste, dat = trainSet)
  pred3 <- predict(m3, newdata = validSet)</pre>
 RMSE3[k] <- sqrt(mean((validSet$Flavor - pred3) ^ 2))</pre>
}
RMSE1
## [1] 0.1479415 0.1653329 0.1556385 0.1656876 0.1482716
RMSE2
## [1] 0.1427025 0.1525461 0.1478815 0.1620440 0.1384244
## [1] 0.1513836 0.1667202 0.1616626 0.1675113 0.1532496
mean(RMSE1)
## [1] 0.1565744
mean(RMSE2)
## [1] 0.1487197
mean(RMSE3)
## [1] 0.1601055
```



LECTURE 20 | 2020-11-18

## 2.20 Combining Cross Validation With Model Selection

- Given some candidate models (e.g., with different variables included) we can do K-fold cross-validation using each of them, and choose the one with the lowest average prediction error across the folds, e.g.,  $\frac{1}{K} \sum_{k=1}^{K} \text{RSME}_k.$
- How to choose *K*? Some common choices are:
  - -K = 5
  - -K = 10
  - K=N (number of observations for train/valid). This is commonly known as "leave one out" (LOO).

The more folds, the higher the computational time, but it can give a better estimate of prediction error.

• What if we are not given a list of models to compare? (and if there are too many predictors to do all possible regressions). Then, we can combine cross-validation with *model selection procedures* (criteria based on training data and search strategy).

To estimate the prediction error for a given *model selection procedure* we apply it K times, each time treating observations in fold  $1, 2, \ldots, K$  as validation sample, e.g., use the *average* RMSE and choose the model selection procedure with the lowest estimated prediction error.

Note: actual variables selected in each fold might be different.

<u>Recommendation</u>: apply chosen model selection *procedure* (lowest estimated prediction error) to the entire set of observations available for training and validation to get a final model (for applications to test data).

Recall we have:

- AIC =  $2q 2\ln[L(\hat{\theta})]$
- BIC =  $q \ln(n) 2 \ln[L(\hat{\theta})]$

We can go further (and beyond PLUS ULTRA!) and consider the following.

# DEFINITION 2.20.1: $L_0$ Penalized Likelihood For any $\lambda>0$ , $q\lambda-2\ln[L(\hat{\theta})]$

### Beyond Stepwise Regression

• Stepwise is a deterministic algorithm which is fast, but might not give optimal model selected.

• Could try stochastic algorithm, e.g., iterative conditional minimization (ICM)

## 2.21 Iterative Conditional Minimization

Start with a model with just intercept. For a given random seed, take predictors  $x_1,\ldots,x_p$  and randomly re-order them into  $x_{(1)},\ldots,x_{(p)}$ . For  $j=1,\ldots,p$ : (let  $\mathcal S$  be the predictors currently in model, so start at  $\mathcal S=\emptyset$ ).

- If  $x_{(i)}$  is not in  $\mathcal{S}$ :
  - Fit model with  $\mathcal{S}$ . If addition of  $x_{(j)}$  improves criterion, then add  $x_{(j)}$  to  $\mathcal{S}$ .
- If  $x_{(i)}$  is in  $\mathcal{S}$ :
  - Fit model removing  $x_{(j)}$ . If excluding  $x_{(j)}$  improves criterion, then remove  $x_{(j)}$  from  $\mathcal{S}$ .

Repeat for loop until no predictors are added/removed from an entire pass through the for loop.

#### **REMARK 2.21.1**

Different random orderings could give different sets S at end of procedure, could pick one that has the best criterion overall.

#### 2.21.1 R Demo

```
## Cross-validation
## Coffee example (Coffee Quality Institute, 2018) continued
coffee <- read.csv("csv/coffee_arabica.csv")</pre>
# 1 = wet. O otherwise
coffee$wet <-</pre>
  ifelse(coffee$Processing.Method == 'Washed / Wet', 1, 0)
# 1 = semi/dry, 0 otherwise
coffee$semi <-
  ifelse(coffee$Processing.Method == 'Semi-washed / Semi-pulped', 1, 0)
coffee$Processing.Method <- NULL</pre>
N <- nrow(coffee)
## Train and validation set split
set.seed(12345678)
trainInd <- sample(1:N, round(N * 0.8), replace = F)</pre>
trainSet <- coffee[trainInd,]</pre>
validSet <- coffee[-trainInd,]</pre>
# Calculate RMSE on three models each with different variables included
m1 <-
 lm(Flavor ~ wet + semi + Aroma + Aftertaste + Body, dat = trainSet)
pred1 <- predict(m1, newdata = validSet)</pre>
sqrt(mean((validSet$Flavor - pred1) ^ 2)) # RMSE
## [1] 0.1577479
mean(abs(validSet$Flavor - pred1)) # MAE
## [1] 0.113643
m2 < -1m(
  Flavor ~ wet + Aroma + Aftertaste +
    Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
  dat = trainSet
pred2 <- predict(m2, newdata = validSet)</pre>
sqrt(mean((validSet$Flavor - pred2) ^ 2))
## [1] 0.1426565
m3 <- lm(Flavor ~ Aroma + Aftertaste, dat = trainSet)
pred3 <- predict(m3, newdata = validSet)</pre>
sqrt(mean((validSet$Flavor - pred3) ^ 2))
## [1] 0.1615385
# K fold cross validation
K <- 5
validSetSplits <- sample((1:N) %% K + 1)</pre>
RMSE1 <-c()
RMSE2 \leftarrow c()
RMSE3 <-c()
for (k in 1:K) {
validSet <- coffee[validSetSplits == k,]</pre>
```

```
trainSet <- coffee[validSetSplits != k,]</pre>
 m1 <-
   lm(Flavor ~ wet + semi + Aroma + Aftertaste + Body, dat = trainSet)
  pred1 <- predict(m1, newdata = validSet)</pre>
  RMSE1[k] <- sqrt(mean((validSet$Flavor - pred1) ^ 2))</pre>
 m2 < -1m(
   Flavor ~ wet + Aroma + Aftertaste +
      Body + Acidity + Balance + Sweetness + Uniformity + Moisture,
    dat = trainSet
  pred2 <- predict(m2, newdata = validSet)</pre>
  RMSE2[k] <- sqrt(mean((validSet$Flavor - pred2) ^ 2))</pre>
 m3 <- lm(Flavor ~ Aroma + Aftertaste, dat = trainSet)
  pred3 <- predict(m3, newdata = validSet)</pre>
 RMSE3[k] <- sqrt(mean((validSet$Flavor - pred3) ^ 2))</pre>
}
RMSE1
## [1] 0.1479415 0.1653329 0.1556385 0.1656876 0.1482716
RMSE2
## [1] 0.1427025 0.1525461 0.1478815 0.1620440 0.1384244
## [1] 0.1513836 0.1667202 0.1616626 0.1675113 0.1532496
mean(RMSE1)
## [1] 0.1565744
mean(RMSE2)
## [1] 0.1487197
mean(RMSE3)
## [1] 0.1601055
```

# Chapter 3

# **Generalized Linear Models**

Lecture 21 2020-11-23

#### Beyond STAT 331: Generalized Linear Models and Logistic Regres-3.1 sion

We have a vector of responses Y which are independent, and predictors  $x_1, \dots, x_n$ .

#### **DEFINITION 3.1.1: Generalized Linear Model (GLM)**

Three ingredients of a GLM:

- 1. Random component. Response  $Y_i$  is a random variable with a distribution that is a member of the exponential family, e.g., Normal, Binomial, Poisson.
- 2. Systematic component. Typically, a linear predictor based on  $x_{i1}, \dots, x_{ip}$  which we denote as

$$\eta_i = \beta_0 + \beta_1 x_{i1} + \dots + \beta_p x_{ip}$$

3. Link function: A function  $g(\cdot)$  that defines a relationship between  $\mathbb{E}[Y_i]$  and  $\eta_i$ ; that is,  $\eta_i$  $g(\mathbb{E}[Y_i])$ .

### **EXAMPLE 3.1.2: Multiple Linear Regression**

$$\begin{split} \text{MLR: } Y_i &= \beta_0 + \beta_1 x_{i1} + \dots + \beta_p x_{ip} + \varepsilon \text{ where } \varepsilon_i \overset{\text{iid}}{\sim} \mathcal{N}(0, \sigma^2) \\ & \bullet \text{ Random component: } Y_i \text{ is Normal and } \mathbb{E}[Y_i] = \eta_i. \end{split}$$

- Link function:  $g(\mathbb{E}[Y_i]) = \mathbb{E}[Y_i]$  (identity).

#### **EXAMPLE 3.1.3: Logistic Regression**

Logistic regression: can be used when response  $Y_i$  is binary; that is,

$$Y_i = \begin{cases} 1 & \text{success/on/yes/goose} \\ 0 & \text{failure/off/no/non-goose} \end{cases}$$

- Random component:  $\mathbb{P}(Y_i=1)=\pi_i$  and  $\mathbb{P}(Y_i=0)=1-\pi_i$ ; that is,  $Y_i\sim \mathrm{Binomial}(1,\pi_i)$ .
- · Link function for logistic regression sets

$$\ln\!\left(\frac{\pi_i}{1-\pi_i}\right) = \beta_0 + \beta_1 x_{i1} + \dots + \beta_p x_{ip} = \eta_i$$

Note:  $\mathbb{E}[Y_i] = 1(\pi_i) + 0(1-\pi_i) = \pi_i$  so logistic regression takes  $g(\pi_i) = \ln\left(\frac{\pi_i}{1-\pi_i}\right)$ . Say A is an event, then the *odds* of event A is defined as

$$\frac{\mathbb{P}(A)}{1 - \mathbb{P}(A)}$$

Therefore,  $\frac{\pi_i}{1-\pi_i}$  is the odds that  $Y_i=1$  and  $\ln\left(\frac{\pi_i}{1-\pi_i}\right)$  is the log-odds that  $Y_i=1$ . Since  $\eta_i=\ln\left(\frac{\pi_i}{1-\pi_i}\right)$ , inverting this function gives

$$e^{\eta_i} = \frac{\pi_i}{1 - \pi_i} \iff \pi_i = \frac{e^{\eta_i}}{1 + e^{\eta_i}}$$

Plot 
$$y = \frac{e^x}{1 + e^x}$$
.

Note that  $\pi_i$  is bounded by 0 to 1, while  $\ln\left(\frac{\pi_i}{1-\pi_i}\right)$  can take any real number.

• Since  $\ln\left(\frac{\pi_i}{1-\pi_i}\right)=\beta_0+\beta_1x_{i1}+\cdots+\beta_px_{ip}$  and  $\mathbb{P}(Y_i=1)=\pi_i$ , then  $\beta_j$  is the expected additive change in the log-odds of the event that  $Y_i=1$  for a unit increase in  $x_j$  (holding other variables constant). Also,

$$\frac{\pi_i}{1-\pi_i} = e^{\beta_0+\beta_1 x_{i1}+\cdots+\beta_p x_{ip}}$$

would say that  $e^{\beta_j}$  is the expected *multiplicative change* in odds of  $Y_i=1$  or a unit increase in  $x_j$  (holding other variables constant).

• Model is fit using MLE (glm() function in R).

#### EXAMPLE 3.1.4: Logistic Regression (Music Analysis)—String Quartet Classification

Define

$$Y_i = \begin{cases} 1 & \text{Haydn} \\ 0 & \text{Mozart} \end{cases}$$

The model (for illustration purposes, the actual model had 7 predictors) is given by:

$$\ln\left(\frac{\pi_i}{1-\pi_i}\right) = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2}$$

From training data (observation with  $Y_i$ 's known and predictors given) we have

$$\hat{\boldsymbol{\beta}} = (-1.12, -15.47, 16.71)^{\top}$$

Also,

- $x_1 = SD$  of note duration in 1st violin where  $x_1 \in (0.05, 0.35)$ .
- $x_2$  = proportion descending pairwise intervals in 1st violin where  $x_2 \in (0.2, 0.5)$ .
- Higher values of  $x_1$  are more likely to be Mozart
- Higher values of  $x_2$  are more likely to be Haydn

Suppose we are given  $\bar{\boldsymbol{x}}_1 = (0.05, 0.2, 0.35)^{\top}$  and  $\boldsymbol{x}_2 = (0.4, 0.4, 0.4)^{\top}$ , then the higher estimated probability is Mozart since  $\hat{\beta}_1$  is negative.

Estimate log odds of  $Y_i = 1$  is

$$-1.12 - 15.47(0.05) + 16.71(0.4) = \hat{\eta}_i$$

Estimate  $\pi_i$  is

$$\frac{e^{\hat{\eta}_i}}{1 + e^{\hat{\eta}_i}} = 0.992$$

which is the estimate probability an observation is Haydn for  $x_1 = 0.05$  and  $x_2 = 0.4$ .

- Second probability: 0.992
- Third probability: 0.537

Interpretation for  $\beta_1$ :

• For unit increase in SD of note duration in 1st violin, estimated expected change in log-odds of the piece being Haydn is -15.47.

## LECTURE 22 | 2020-11-25

#### 3.2 Last Lecture!

**Project Tips:** 

- · Explore the data
- No one "right" model
- · Understand what you're fitting and why
- Present it well (no raw R output in report)
- Data set has many variables (be careful about overfitting)—don't wait until last minute.

Residual Surrealism:

- Want  $(\hat{\mu}_i, e_i)$ , i = 1, ..., n to represent black pixel locations in the image.
- Recall in fitting MLR that e and  $\hat{\mu}$  satisfy

$$\hat{m{\mu}} = Hm{Y}$$
  
 $m{e} = (I - H)m{Y}$ 

which is a linear system of 2n equations.

- Reverse engineer what  $Y, x_1, ..., x_p$  could be. There are n(p+1) free variables.
- Fix  $\beta$  (choose our regression coefficient).
- Recall  $Y = X\beta + \varepsilon$ , so the equations can be expressed as

$$\hat{\mu} = H(X\beta + \varepsilon) = X\beta + H\varepsilon$$
  
 $e = (I - H)\varepsilon = \varepsilon - H\varepsilon$ 

· Residuals also satisfy condition

$$\hat{\boldsymbol{\mu}}^{\mathsf{T}}\boldsymbol{e} = 0$$

since  $\boldsymbol{e}$  are orthogonal to  $\mathrm{Span}(X)$  which might not be satisfied by an arbitrary image.

Solution. Add a few pixels to corners (outliers, influential points) to satisfy orthogonality.

• Least squares solution implies  $\mathbf{1}^{\mathsf{T}}e=0$  for the intercept, and  $\tilde{X}^{\mathsf{T}}e=\mathbf{0}$  for the predictors.

$$X = \begin{bmatrix} \mathbf{1} & \tilde{X} \end{bmatrix}$$

So if we let

$$\tilde{X} = \bigg(I_n - \frac{ee^\top}{e^\top e}\bigg) M_{n \times p}$$

then  $\tilde{X}e = 0$  is satisfied, for any matrix M.

• Simulate  ${\bf Z}$  i.i.d. from  $\mathcal{N}(0,\tau^2)$  controls  $R^2$  in MLR and let

$$\boldsymbol{arepsilon} = \boldsymbol{e} + H \boldsymbol{Z}$$

and substitute in 1st equation to get

$$\hat{m{\mu}} = X m{eta} + H m{e} + H m{Z} = \underbrace{X m{eta}}_{ ext{function of } M} + \underbrace{H m{Z}}_{ ext{function of } M}$$

• Iterative solution to update M until it satisfies the equation.

The program provided by the author that has the easiest version to use runs on Windows, so I had to boot up a Virtual Box for Windows.—Samuel Wong

## **REMARK 3.2.1: Author's Note**

I use Arch btw.