Linear modeling I

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General notes on Homework

Do not use Rstudio-specific commands in R chunks

- View() is not an R command
- May prevent from knitting

Only set working directory once

• Add to top chunk with "root.dir"

Load all libraries in the top chunk

Chi-squared assumptions refer to expected counts

Don't give yourself more work than you have to

Don't make your hypotheses fancy

When to use median vs. mean for directional conclusions?

Announcing Final Project (25%)

Identify a dataset and ask **four scientific questions** about the data

- Use any of the statistical approaches we have learned to answer each question
- Make a descriptive figure for each scientific question

Scientific questions

Recall HW7:

- Do PHA levels tend to differ between the birds that received supplemental carotenoids and those that did not? Based on your results, can you infer anything about immune differences between birds that did and did not receive carotenoids?
 - Use a Mann Whitney U test to answer the scientific question
 - "Run a Mann Whitney U test on PHA levels between bird treatments" is not a scientific question
- What figure might be good to make here?
- What figures would not be good here?

Final project proposal

Homework due 11/28 will be a proposal

- Identify your dataset and give 1-2 paragraphs of background
 - IN YOUR OWN WORDS
- Pose four scientific questions
- Explain how you will solve each question
 - What statistical procedure and why
- Indicate how you will visualize your data

3-4 sentences total per question

Around 1 written page, single-spaced.

Updated schedule

Date	Торіс
10/24	Linear modeling I
10/31	Linear modeling II and logistic regression
11/7	Model selection and evaluation
11/14	Principal Components Analysis (PCA) and clustering
11/21	Thanksgiving break
11/28	Advanced R grabbag and/or overflow
12/5	Advanced R grabbag and/or in-class office hours for final project Email me for special topic requests.
12/12	Final project due (by 11:59 pm on 12/12)

Linear Modeling

- ANOVA and friends
- Correlation
- Regression
- Multiple regression

ANOVA: Analysis of Variance

Used to compare more than 2 means (among k groups)

Ho: All means are the same, i.e. $\mu_1 = \mu_2 = ... = \mu_k$

Ha: At least one mean is different, i.e. at least one $\mu_{<1-k>}$ differs

Why "can't" we use a *t*-test? • We can do all the comparisons and use a P-value correction • ANOVA is preferred

ANOVA Example

A clinical trial asks if there is a difference in mean daily calcium intake in adults with <u>normal bone density</u>, <u>adults with osteopenia</u>, and <u>adults with osteoporosis</u>. Each participant's daily calcium intake is measured based on reported food intake.

Normal Bone Density	Osteopenia	Osteoporosis
1200	1000	490
1000	1100	650
980	700	200
900	800	300
750	500	400
800	700	350

Is there a difference in mean calcium intake across groups?

ANOVA compares sources of *variance* using the **F statistic**

Variance *among* groups is the **group mean square** (MS_{group}) Variance *within* each group is the **error mean square** (MS_{error}) • Pooled sample variance



$$F = \frac{MS_{group}}{MS_{error}}$$

df group =
$$k - 1$$

df error = $N - k = \sum n - 1$



https://en.wikipedia.org/wiki/F-distribution#/media/File:F-distribution_pdf.svg

We use only the upper tail for P-value

If H_0 is true, $F \cong 1$ $MS_{group} \cong MS_{error}$



If H_0 is false, F > 1 $MS_{group} > MS_{error}$



Computing the F statistic

$$F = \frac{MS_{group}}{MS_{error}} MS = \frac{SS}{df} Su$$
Calculate an estimate of the variate structure in the variate of the variate of the variate structure in the struct

Sum of squares

 s_i^2 =Standard deviation of group i $df_i = n_i - 1$, where n_i = sample size of group i \overline{X}_i = mean of group i $SS_{error} = \sum df_i s_i^2$ \overline{X} = grand mean (mean of *all* numbers)

Error degrees of freedom =

$$df_{error} = \sum df_i = \sum (n_i - 1) = N - k$$

 S_{error} is like the pooled variance in a 2-sample *t*-test.

Computing the **F** statistic

$$F = \frac{MS_{group}}{MS_{error}} \quad MS = \frac{SS}{df}$$
$$MS_{error} = \frac{\sum s_i^2 (n_i - 1)}{N - k}$$

Normal Bone Density	Osteopenia	Osteoporosis
1200	1000	490
1000	1100	650
980	700	200
900	800	300
750	500	400
800	700	350

$$\frac{\sum n_i (\overline{X}_i - \overline{X})^2}{k-1}$$

Running the ANOVA

Ho: Groups have the same mean calcium intake.

Ha: At least one group has a different calcium intake.

```
group calcium
                                                                                         normal
                                                                                                  1200
                                                                                  1
2
3
4
5
6
data <- tibble("normal"
                                                                                                  1000
                                   = c(1200, 1000, 980, 900, 750, 800),
                                                                                         normal
                                                                                                   980
                                                                                         normal
                  "osteopenia" = c(1000, 1100, 700, 800, 500, 700),
                                                                                                   900
                                                                                         normal
                  "osteoporosis" = c(490, 650, 200, 300, 400, 350))
                                                                                                   750
                                                                                         normal
                                                                                                   800
                                                                                         normal
                                                                                  7
                                                                                                  1000
                                                                                     osteopenia
data %>% gather(group, calcium, normal:osteoporosis) -> tidy.data
                                                                                  8
                                                                                     osteopenia
                                                                                                  1100
                                                                                  9
                                                                                                   700
                                                                                     osteopenia
                                                                                 10
                                                                                     osteopenia
                                                                                                   800
                                                                                 11
                                                                                     osteopenia
                                                                                                   500
                                                                                 12
                                                                                     osteopenia
                                                                                                   700
                                                                                 . . .
```

Visualize the data

It is always the right idea to view your data before modeling it

ggplot(tidy.data, aes(x = group, y = calcium, fill= group)) + geom_violin()



Running the ANOVA

> aov(calcium ~ group, data = tidy.data) Call:

aov(formula = calcium ~ group, data = tidy.data)

Terms:

error

	group	Residuals
Sum of Squares	944144.4	493166.7
Deg. of Freedom	2	15

Residual standard error: 181.3223 Estimated effects may be unbalanced

Obtaining the ANOVA table

> summary(aov(calcium ~ group, data = tidy.data))



Reports and conclusions

Our P = 0.000328, which is less than alpha. We reject the null hypothesis and we have evidence that at least one mean (normal bone density, osteopenia, or osteoporosis calcium intake) differs from the other.

Unplanned comparisons with the Tukey-Kramer Method

AKA Tukey's test, Tukey's method, Tukey's HSD (honest significant difference) test

Tests all pairs of means0.8• Normal vs. osteopenia0.7• Normal vs. osteoporosis0.6• Osteopenia vs. osteoporosis0.6• Osteopenia vs. osteoporosis0.6• Osteopenia vs. osteoporosis0.4• Osteopenia vs. osteopenia vs. osteoporosis0.4• Osteopenia vs. osteopen

0

-2

0

2

Running Tukey's test on ANOVA

> TukeyHSD(aov(calcium ~ group, data = tidy.data))
Tukey multiple comparisons of means
95% family-wise confidence level

Fit: aov(formula = calcium ~ group, data = tidy.data)

\$group

diff lwr upr p adj osteopenia-normal -138.3333 -410.2534 133.5867 0.4054988 osteoporosis-normal -540.0000 -811.9200 -268.0800 0.0003238 osteoporosis-osteopenia -401.6667 -673.5867 -129.7466 0.0043335

Reports and conclusions, round 2

Our P = 0.000328, which is less than alpha. We reject the null hypothesis and we have evidence that at least one mean (normal bone density, osteopenia, or osteoporosis calcium intake) differs from the other.

After running the *post-hoc* Tukey's test, we find that osteoporosis groups have a significantly higher average calcium intake than normal groups (P=0.0003), and that osteoporosis groups have a significantly higher average calcium intake than osteopenia groups P=0.004). However, we do not find a significant difference in calcium intake between normal and osteopenia groups.

ANOVA assumptions

Samples are random

Samples are normally distributed

• Robust to violations when study is large

Samples have the same variance

• Robust to violations when study is **balanced**

Kruskal-Wallis is the non-parametric alternative

> kruskal.test(calcium ~ as.factor(group), data = tidy.data)

Kruskal-Wallis rank sum test

data: calcium by as.factor(group)
Kruskal-Wallis chi-squared = 11.439, df = 2, p-value = 0.003281

Unless something is **really really weird**, you "can" use ANOVA

Exercise break

Correlation

Measures the strength and direction of the linear association between two numeric variables

 $-1 \leq r \leq 1$ *r* = 0.5 *r* = −0.7 *r* = 0.9 r = 0.0> X X X X

Perfect correlations



Variability (error) has a substantial influence



Calculating the correlation coefficient

$$r = \frac{cov(X,Y)}{s_X s_Y}$$

$$=\frac{\sum(X_i-\overline{X})(Y_i-\overline{Y})}{\sqrt{\frac{\sum(X_i-\overline{X})^2}{n-1}}\sqrt{\frac{\sum(Y_i-\overline{Y})^2}{n-1}}}$$

$$= \frac{1}{n-1} \frac{\sum (X_i - \overline{X})(Y_i - \overline{Y})}{\sqrt{\sum (X_i - \overline{X})^2} \sqrt{\sum (Y_i - \overline{Y})^2}}$$

Example: correlation between irises

- > setosa <- iris %>% filter(Species == "setosa")
 > cor(setosa\$Sepal.Length, setosa\$Sepal.Width)
 [1] 0.7425467
- > cor(setosa\$Sepal.Width, setosa\$Sepal.Length)
 [1] 0.7425467



Example: correlation between irises

> cor(setosa\$Petal.Length, setosa\$Sepal.Width)
[1] 0.1777



Hypothesis testing with correlations

 H_0 : Petal length and sepal width are not correlated (r=0) H_A : Petal length and sepal width are correlated (r!=0)

```
> cor.test(setosa$Sepal.Width, setosa$Sepal.Length)
Pearson's product-moment correlation
```

Hypothesis testing

> cor.test(setosa\$Petal.Length, setosa\$Sepal.Width)

```
Pearson's product-moment correlation
```

We fail to reject the null hypothesis. There is no evidence that the correlation between petal lengths and sepal width in setosa irises differs from 0.

Nonlinear data



Nonlinear data



```
> cor.test(x, y)
```

Pearson's product-moment correlation

Spearman rank nonparametric correlation

Assumes data is *monotonic* (ordinal)

> cor.test(x, y, method = "spearman")

Spearman's rank correlation rho

```
data: x and y
S = 0, p-value < 2.2e-16
alternative hypothesis: true rho is not equal to 0
sample estimates:
rho
1</pre>
```

Assumptions: Check by plotting

Data are linearly related without any severe outliers

Both X and Y are normally distributed

• Robust to large N

Cloud of points is not "funnel-shaped" (fans out at end(s))



Exercise break

Regression

The simplest type of *linear model*

Predicts the value of one numeric variable from another via "line of best fit"

$$Y = a + bX$$

$$Y = \beta_o + \beta_1 X + \varepsilon$$

Residuals: \mathcal{E} is a random error component that measures how far above/below the line the **actual** value of Y for a given X lies. Mean is 0.

Least squares approach

Find the line which **minimizes** the sum of squared residuals





Least squares approach

Find the line which **minimizes** the sum of squared residuals



Calculating slope and intercept Y = a + bX

$$b = \frac{cov(X,Y)}{s_X^2} = \frac{\sum(X_i - \overline{X})(Y_i - \overline{Y})}{\frac{1}{n-1}\sum(X_i - \overline{X})^2}$$

 $a = \overline{Y} - b\overline{X}$

The point $(\overline{X}, \overline{Y})$ always goes through the regression line

Executing a linear model

```
### lm(Y ~ X, data = <data frame>) ###
```

> lm(Sepal.Length ~ Sepal.Width, data = setosa)

```
Call:
lm(formula = Sepal.Length ~ Sepal.Width, data = setosa)
```

```
Coefficients:
(Intercept) Petal.Width
2.6390 0.6905
```

$$Y = 2.64 + 0.69X$$

Testing a linear model

```
> summary( lm(Sepal.Length ~ Sepal.Width, data = setosa) )
Call:
lm(formula = Sepal.Length ~ Sepal.Width, data = setosa)
```

Residuals:

Min 1Q Median 3Q Max -0.52476 -0.16286 0.02166 0.13833 0.44428 Five number summary of the distribution of residuals

Coefficients:	
Estimate Std. Error t value Pr(> t)	
(Intercept) 2.6390 0.3100 8.513 3.74e-11 ***	
Sepal.Width 0.6905 0.0899 7.681 6.71e-10 ***	
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1	
Residual standard error: 0.2385 on 48 degrees of freedom SE of	8
Multiple R-squared: 0.5514, Adjusted R-squared: 0.542	
F-statistic: 58.99 on 1 and 48 DF, p-value: 6.71e-10 Test feedback	or model improvement over slope=0

Coefficients

Y X lm(Sepal.Length ~ Sepal.Width)



What can we conclude?

On average, setosa Sepal Length (Y) increases by 0.6905 cm (+/-0.0899 SE) for every 1 cm of Sepal Width (X).

[If P > alpha, don't conclude this..]

\mathbb{R}^2

R^2 is the percent of variation in Y than can be explained by X • $0 \le R^2 \le 1$

Multiple R-squared: 0.5514, Adjusted R-squared: 0.542

$$R^{2} = \left[\frac{cov(X,Y)}{s_{X}s_{Y}}\right]^{2} = 1 - \frac{SS_{res}}{SS_{total}} = \frac{Explained \ variation}{Total \ variation}$$

What can we conclude?

~55% of the variation in Setosa sepal lengths (Y) can be explained by Setosa sepals widths (X).

[If P < alpha, don't conclude this..]

Broom to the rescue

```
> summary( lm(Sepal.Length ~ Sepal.Width, data = setosa) )
Call:
lm(formula = Sepal.Length ~ Sepal.Width, data = setosa)
Residuals:
    Min
              10 Median 30
                                       Max
-0.52476 -0.16286 0.02166 0.13833 0.44428
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
                       0.3100 8.513 3.74e-11 ***
(Intercept) 2.6390
Sepal.Width 0.6905 0.0899 7.681 6.71e-10 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.2385 on 48 degrees of freedom
Multiple R-squared: 0.5514, Adjusted R-squared: 0.542
F-statistic: 58.99 on 1 and 48 DF, p-value: 6.71e-10
```

Broom to the rescue

```
> library(broom)
> model <- lm(Sepal.Length ~ Sepal.Width, data = setosa)</pre>
##### Coefficients and Pvalues #####
> tidy(model)
        term estimate std.error statistic p.value
1 (Intercept) 2.6390012 0.31001431 8.512514 3.742438e-11
2 Sepal.Width 0.6904897 0.08989888 7.680738 6.709843e-10
###### Concise *one row* summary #####
> glance(model)
 r.squared adj.r.squared sigma statistic p.value df logLik
1 0.5513756 0.5420292 0.2385422 58.99373 6.709843e-10 2 1.734067 2.531865
      BIC deviance df.residual
1 8.267934 2.731315
                           48
```

AIC

Broom to the rescue

Add columns from fit to the original data that was modeled
> augment(model)
Sepal.Length Sepal.Width .fitted .se.fit .resid .hat .sigma .cooksd .std.resid

1	5.1	3.5 5.055715 0.03435031 0.04428474 0.02073628 0.2409782 3.726311e-04 0.18760265
2	4.9	3.0 4.710470 0.05117134 0.18952960 0.04601750 0.2393991 1.596010e-02 0.81347001
3	4.7	3.2 4.848568 0.03947370 -0.14856834 0.02738325 0.2400630 5.614273e-03 -0.63152438
4	4.6	3.1 4.779519 0.04480537 -0.17951937 0.03528008 0.2395878 1.073468e-02 -0.76620575
5	5.0	3.6 5.124764 0.03710984 -0.12476423 0.02420180 0.2403616 3.476539e-03 -0.52947419
6	5.4	3.9 5.331911 0.05420835 0.06808885 0.05164186 0.2408507 2.339099e-03 0.29310589

ggplot(setosa, aes(x = Sepal.Width, y = Sepal.Length)) +
 geom_point() + geom_smooth(method = "lm")



ggplot(setosa, aes(x = Sepal.Width, y = Sepal.Length)) +
 geom_point() + geom_smooth(method = "lm") +
 geom_text(x = 2.75, y = 5.5, label = "y=2.639 + 0.69x", color="red")



ggplot(setosa, aes(x = Sepal.Width, y = Sepal.Length)) +
 geom_point() + geom_smooth(method = "lm") +
 geom_text(x=2.75, y=5.5, label="r^2 == 0.585", parse=TRUE, color="red")



Using the model: Predicting responses



- > new.data <-tibble(Sepal.Width = 2.6) ## Same column name as model's predictor</pre>
- > predict(model, new.data)
 1
 4.434275

Predicting with intervals

Confidence interval

• Range that is likely to contain the mean response

$$\hat{y}_h \pm t_{(\alpha/2, n-2)} \times \sqrt{MSE\left(\frac{1}{n} + \frac{(x_h - \bar{x})^2}{\sum (x_i - \bar{x})^2}\right)}$$

Prediction interval

- Range that is likely to contain the response value of a single new observation
- Wider than CI due to added uncertainty for predicting a single point

$$\hat{y}_h \pm t_{(\alpha/2, n-2)} \times \sqrt{MSE \times \left(1 + \frac{1}{n} + \frac{(x_h - \bar{x})^2}{\sum (x_i - \bar{x})^2}\right)}$$

Predicting with intervals

Assumptions of linear models

Residuals are normally distributed

The variance is the same for all predictors*

Predictors are independent of each other*

The relationship between response and any numeric predictors is linear*

Normality of residuals

- > augmented <- augment(model)</pre>
- > qqnorm(augmented\$.resid)
- > qqline(augmented\$.resid)



How to check regression assumptions

- **1.** Plot response and predictor against each other to ensure linearity
 - Critically important
- 2. Plot the residuals to ensure normality
 - Important, usually overlooked
 - Most times we are robust to departures

Exercise break

Linear Models

lm(Numeric response ~ <predictors>)

Linear Models

lm(Numeric response ~ <predictors>)

Single numeric predictor: Regression Single categorical predictor: ANOVA Multiple numeric predictors: multiple regression Multiple categorical predictors: *n*-way ANOVA Single categorical and *n* numeric predictors: ANCOVA Multiple categorical and *n* numeric predictors: linear model



Briefly, bootstrapping the regression

```
> library(slipper)
> setosa %>%
    slipper_lm(Sepal.Length ~ Sepal.Width, B=1e3)%>% head()
         term
                  value
                            type
1 (Intercept) 2.6390012 observed
2 Sepal.Width 0.6904897 observed
3 (Intercept) 2.0900929 bootstrap
4 Sepal.Width 0.8467474 bootstrap
5 (Intercept) 2.7629316 bootstrap
6 Sepal.Width 0.6527575 bootstrap
setosa %>%
    slipper_lm(Sepal.Length ~ Sepal.Width, B=1e3) %>%
    filter(type == "bootstrap", term == "Sepal.Width") %>%
    summarize(mean = mean(value),
              ci_low = quantile(value,0.025),
              ci_high = quantile(value,0.975))
               ci_low ci_high
       mean
1 0.6945918 0.5302098 0.8961058
```