Analysis of Covariance

Analysis of covariance

- ANOVA: explanatory variables categorical (divide data into groups)
- traditionally, analysis of covariance has categorical x's plus one numerical x ("covariate") to be adjusted for.
- 1m handles this too.
- Simple example: two treatments (drugs) (a and b), with before and after scores.
- Does knowing before score and/or treatment help to predict after score?
- Is after score different by treatment/before score?

Data

Treatment, before, after:

a 5 20

a 10 23

a 12 30

a 9 25

a 23 34

a 21 40

a 14 27

a 18 38

a 6 24

a 13 31

b 7 19

b 12 26

b 27 33

b 24 35

b 18 30

b 22 31

Packages

```
library(tidyverse)
library(broom)
library(marginaleffects)
```

the last of these for predictions.

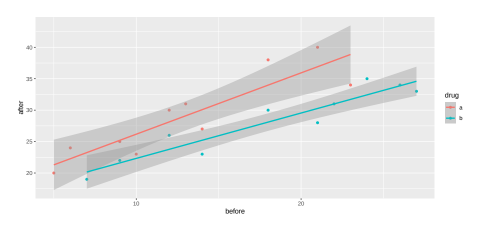
Read in data

```
url <- "http://ritsokiguess.site/datafiles/ancova.txt"
prepost <- read_delim(url, " ")
prepost</pre>
```

```
# A tibble: 20 x 3
   drug before after
   <chr> <dbl> <dbl>
              5
 1 a
                   20
2 a
             10 23
3 a
             12 30
4 a
                   25
 5 a
            23
                   34
 6 a
             21
                   40
7 a
             14
                   27
             18
                   38
8 a
 9 a
              6
                   24
10 a
             13
                   31
```

Making a plot

```
ggplot(prepost, aes(x = before, y = after, colour = drug)) +
geom_point() + geom_smooth(method = "lm")
```



Comments

- As before score goes up, after score goes up.
- Red points (drug A) generally above blue points (drug B), for comparable before score.
- Suggests before score effect and drug effect.

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The means

```
prepost %>%
  group_by(drug) %>%
  summarize(
   before_mean = mean(before),
   after_mean = mean(after)
)
```

- Mean "after" score slightly higher for treatment A.
- Mean "before" score much higher for treatment B.
- Greater improvement on treatment A.

Testing for interaction

```
prepost.1 <- lm(after ~ before * drug, data = prepost)
drop1(prepost.1, test = "F")</pre>
```

Single term deletions

```
anova(prepost.1)
```

Analysis of Variance Table

```
Response: after

Df Sum Sq Mean Sq F value Pr(>F)

hefers 1 420 02 420 02 62 6804 6 240 07 ***
```

Predictions

Set up values to predict for:

```
summary(prepost)
```

drug

```
Length:20
                  Min. : 5.00
                                  Min. :19.00
Class :character
                   1st Qu.: 9.75
                                  1st Qu.:23.75
                                  Median :29.00
Mode :character
                   Median :14.00
                   Mean :15.55
                                  Mean :28.65
                   3rd Qu.:21.25
                                  3rd Qu.:33.25
                   Max. :27.00
                                  Max. :40.00
new \leftarrow datagrid(before = c(9.75, 14, 21.25),
               drug = c("a", "b"), model = prepost.1)
```

before after

before drug rowid l 9.75 a 1

new

and then

2

3

4

5

6

b

b

```
cbind(predictions(prepost.1, newdata = new)) %>%
  select(drug, before, estimate, conf.low, conf.high)

drug before estimate conf.low conf.high
1   a  9.75 25.93250 24.05059 27.81442
```

9.75 22.14565 19.58681 24.70450

14.00 30.07784 28.43296 31.72271

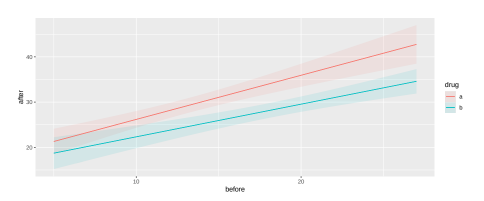
14.00 25.21304 23.32649 27.09959

21, 25, 30, 44565, 28, 64373, 32, 24758

a 21.25 37.14929 34.32557 39.97300

Predictions (with interaction included), plotted

plot_predictions(model = prepost.1, condition = c("before", "c



Lines almost parallel, but not quite.

Taking out interaction

```
prepost.2 <- update(prepost.1, . ~ . - before:drug)</pre>
drop1(prepost.2, test = "F")
Single term deletions
Model:
after ~ before + drug
      Df Sum of Sq RSS AIC F value Pr(>F)
                  122.32 42.218
<none>
before 1 540.18 662.50 74.006 75.074 1.211e-07 ***
drug 1 115.31 237.63 53.499 16.025 0.0009209 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
summary(prepost.2)
```

```
Call:
```

lm(formula = after ~ before + drug, data = prepost)

Predictions

```
cbind(predictions(prepost.2, newdata = new)) %>%
  select(drug, before, estimate)
```

```
drug before estimate

1 a 9.75 26.42794

2 b 9.75 21.27328

3 a 14.00 29.94473

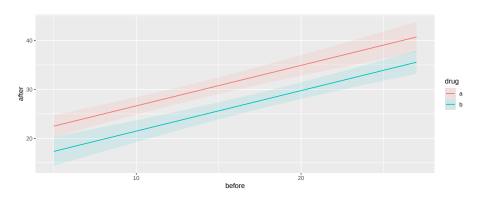
4 b 14.00 24.79007

5 a 21.25 35.94397

6 b 21.25 30.78931
```

Plot of predicted values

plot_predictions(prepost.2, condition = c("before", "drug"))



This time the lines are *exactly* parallel. No-interaction model forces them to have the same slope.

Different look at model output

• anova(prepost.2) tests for significant effect of before score and of drug, but doesn't help with interpretation.

Analysis of Covariance

• summary(prepost.2) views as regression with slopes:

```
summary(prepost.2)
```

Call:

```
lm(formula = after ~ before + drug, data = prepost)
Residuals:
   Min
            10 Median
                           30
                                  Max
-3.6348 -2.5099 -0.2038 1.8871 4.7453
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 18.3600 1.5115 12.147 8.35e-10 ***
before
         0.8275
                       0.0955 8.665 1.21e-07 ***
         -5.1547 1.2876 -4.003 0.000921 ***
drugb
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 2.682 on 17 degrees of freedom
Multiple R-squared: 0.817, Adjusted R-squared: 0.7955
```

Understanding those slopes

tidy(prepost.2)

- before ordinary numerical variable; drug categorical.
- 1m uses first category druga as baseline.
- Intercept is prediction of after score for before score 0 and *drug A*.
- before slope is predicted change in after score when before score increases by 1 (usual slope)
- Slope for drugb is *change* in predicted after score for being on drug B rather than drug A. Same for *any* before score (no interaction).

Summary

- ANCOVA model: fits different regression line for each group, predicting response from covariate.
- ANCOVA model with interaction between factor and covariate allows different slopes for each line.
- Sometimes those lines can cross over!
- If interaction not significant, take out. Lines then parallel.
- With parallel lines, groups have consistent effect regardless of value of covariate.

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