STAD29: Statistics for the Life and Social Sciences

Lecture notes

Section 1

Course Outline

Course and instructor

- Lecture: Wednesday 14:00-16:00 in HW 215. Optional computer lab Monday 16:00-17:00 in BV 498.
- Instructor: Ken Butler
- Office: IC 471.
- E-mail: butler@utsc.utoronto.ca
- Office hours: Monday 11:00-13:00. I am often around otherwise. See if I'm in. Or make an appointment. E-mail always good.
- Course website: link.
- Using Quercus for assignments/grades only; using website for everything else.

Texts

- There is no official text for this course.
- You may find "R for Data Science", link helpful for R background.
- I will refer frequently to my book of Problems and Solutions in Applied Statistics (PASIAS), link.
- Both of these resources are and will remain free.

Programs, prerequisites and exclusions

- Prerequisites:
- For undergrads: STAC32. Not negotiable.
- For grad students, a first course in statistics, and some training in regression and ANOVA. The less you know, the more you'll have to catch up!
- This course is a required part of Applied Statistics minor.
- Exclusions: this course is not for Math/Statistics/CS
 majors/minors. It is for students in other fields who wish to learn
 some more advanced statistical methods. The exclusions in the
 Calendar reflect this.
- If you are in one of those programs, you won't get program credit for this course, or for any future STA courses you take.

Computing

- Computing: big part of the course, not optional. You will need to demonstrate that you can use R to analyze data, and can critically interpret the output.
- For grad students who have not come through STAC32, I am happy to offer extra help to get you up to speed.

Assessment 1/2

 Grading: (2 hour) midterm, (3 hour) final exam. Assignments most weeks, due Tuesday at 11:59pm. Graduate students (STA 1007) also required to complete a project using one or more of the techniques learned in class, on a dataset from their field of study. Projects due on the last day of classes.

Assessment:

| | STAD29 | STA 1007 |
|--------------|--------|----------|
| Assignments | 20% | 20% |
| Midterm exam | 30% | 20% |
| Project | - | 20% |
| Final exam | 50% | 40% |

Assessment 2/2

- Assessments missed with documentation will cause a re-weighting of other assessments of same type. No make-ups.
- You must pass the final exam to guarantee passing the course. If you fail the final exam but would otherwise have passed the course, you receive a grade of 45.

Plagiarism

- This link defines academic offences at this university. Read it. You are bound by it.
- Plagiarism defined (at the end) as
 The wrongful appropriation and purloining, and publication as
 one's own, of the ideas, or the expression of the ideas ... of another.
- The code and explanations that you write and hand in must be yours and yours alone.
- When you hand in work, it is implied that it is your work. Handing in work, with your name on it, that was actually done by someone else is an academic offence.
- If I am suspicious that anyone's work is plagiarized, I will take action.

Getting help

- The English Language Development Centre supports all students in developing better Academic English and critical thinking skills needed in academic communication. Make use of the personalized support in academic writing skills development. Details and sign-up information: link.
- Students with diverse learning styles and needs are welcome in this course. In particular, if you have a disability/health consideration that may require accommodations, please feel free to approach the AccessAbility Services Office as soon as possible. I will work with you and AccessAbility Services to ensure you can achieve your learning goals in this course. Enquiries are confidential. The UTSC AccessAbility Services staff are available by appointment to assess specific needs, provide referrals and arrange appropriate accommodations: (416) 287-7560 or by e-mail: ability@utsc.utoronto.ca.

Course material

- Regression-like things
 - review of (multiple) regression
 - logistic regression (including multi-category responses)
 - survival analysis
- ANOVA-like things
 - more ANOVA
 - multivariate ANOVA
 - repeated measures
- Multivariate methods
 - discriminant analysis
 - cluster analysis
 - multidimensional scaling
 - principal components
 - factor analysis
- Miscellanea
 - time series
 - multiway frequency tables

Section 2

Review of (multiple) regression

Regression

- Use regression when one variable is an outcome (response, y).
- See if/how response depends on other variable(s), explanatory, x_1, x_2, \ldots
- Can have one or more than one explanatory variable, but always one response.
- Assumes a straight-line relationship between response and explanatory.
- Ask:
 - is there a relationship between y and x's, and if so, which ones?
 - what does the relationship look like?

Packages

```
library(MASS) # for Box-Cox, later
library(tidyverse)
library(broom)
```

A regression with one x

13 children, measure average total sleep time (ATST, mins) and age (years) for each. See if ATST depends on age. Data in sleep.txt, ATST then age. Read in data:

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/sleep.txt"
sleep <- read_delim(my_url, " ")

## Parsed with column specification:
## cols(
## atst = col_double(),
## age = col double()</pre>
```

)

Check data

summary(sleep)

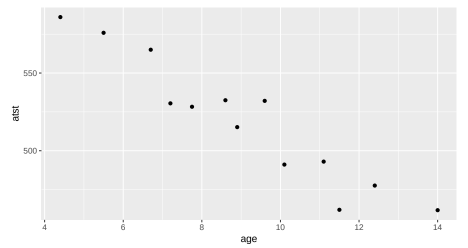
atst

##

```
age
##
   Min. :461.8
                  Min. : 4.400
##
   1st Qu.:491.1
                  1st Qu.: 7.200
   Median :528.3
##
                  Median: 8.900
   Mean :519.3
##
                  Mean : 9.058
##
   3rd Qu.:532.5
                  3rd Qu.:11.100
##
   Max. :586.0
                  Max. :14.000
```

Make scatter plot of ATST (response) vs. age (explanatory) using code overleaf:

The scatterplot



Correlation

• Measures how well a straight line fits the data:

```
with(sleep, cor(atst, age))
```

```
## [1] -0.9515469
```

- ullet 1 is perfect upward trend, -1 is perfect downward trend, 0 is no trend.
- This one close to perfect downward trend.
- Can do correlations of all pairs of variables:

cor(sleep)

```
## atst age
## atst 1.0000000 -0.9515469
## age -0.9515469 1.0000000
```

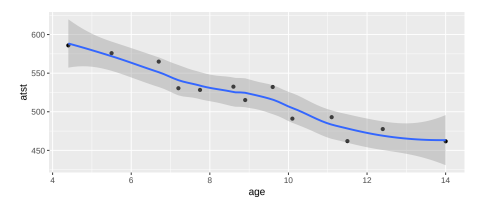
Lowess curve

- Sometimes nice to guide the eye: is the trend straight, or not?
- Idea: lowess curve. "Locally weighted least squares", not affected by outliers, not constrained to be linear.
- Lowess is a *guide*: even if straight line appropriate, may wiggle/bend a little. Looking for *serious* problems with linearity.
- Add lowess curve to plot using geom_smooth:

Plot with lowess curve

```
ggplot(sleep, aes(x = age, y = atst)) + geom_point() +
  geom_smooth()
```

```
## geom_smooth() using method = 'loess' and formula 'y ~ x'
```



The regression

Scatterplot shows no obvious curve, and a pretty clear downward trend. So we can run the regression:

```
sleep.1 <- lm(atst ~ age, data = sleep)</pre>
```

The output

summary(sleep.1)

```
##
## Call:
## lm(formula = atst ~ age, data = sleep)
##
## Residuals:
      Min 10 Median 30
##
                                    Max
## -23.011 -9.365 2.372 6.770 20.411
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 646.483 12.918 50.05 2.49e-14 ***
## age
              -14.041 1.368 -10.26 5.70e-07 ***
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 13.15 on 11 degrees of freedom
## Multiple R-squared: 0.9054, Adjusted R-squared: 0.8968
## F-statistic: 105.3 on 1 and 11 DF, p-value: 5.7e-07
```

Conclusions

- The relationship appears to be a straight line, with a downward trend.
- F-tests for model as a whole and t-test for slope (same) both confirm this (P-value $5.7 \times 10^{-7} = 0.00000057$).
- \bullet Slope is -14, so a 1-year increase in age goes with a 14-minute decrease in ATST on average.
- R-squared is correlation squared (when one x anyway), between 0 and 1 (1 good, 0 bad).
- Here R-squared is 0.9054, pleasantly high.

Doing things with the regression output

- Output from regression (and eg. *t*-test) is all right to look at, but hard to extract and re-use information from.
- Package broom extracts info from model output in way that can be used in pipe (later):

```
tidy(sleep.1)
```

```
## # A tibble: 2 x 5
##
    term
                estimate std.error statistic p.value
    <chr>
                   <dbl>
                             <dbl>
                                      <dbl>
                                               <dbl>
##
## 1 (Intercept)
                   646. 12.9
                                       50.0 2.49e-14
                             1.37
                                      -10.3 5.70e- 7
  2 age
                   -14.0
```

also one-line summary of model:

glance(sleep.1)

Broom part 2

A tibble: 8 x 9

```
sleep.1 %>% augment(sleep) %>% slice(1:8)
```

```
##
     atst
          age .fitted .se.fit .resid .hat .sigma .cooksd
##
    <dbl> <dbl>
                <dbl>
                      <dbl> <dbl> <dbl>
                                        <dbl>
                                               <dbl>
        4.4
                 585. 7.34 1.30 0.312
## 1
     586
                                         13.8 0.00320
    462. 14 450. 7.68 11.8
                                         13.0 0.319
## 2
                                  0.341
   491. 10.1 505.
                    3.92 -13.6 0.0887
                                         13.0 0.0568
## 3
## 4
     565
        6.7
                552.
                     4.87 12.6 0.137
                                          13.1 0.0844
    462 11.5
                485. 4.95 -23.0 0.141
                                          11.3 0.294
## 5
## 6
    532. 9.6
                512.
                       3.72 20.4 0.0801
                                          12.0 0.114
```

472. 5.85 5.23 0.198

3.65 -6.32 0.0772

Useful for plotting residuals against an *x*-variable.

522.

... with 1 more variable: .std.resid <dbl>

515.

478. 12.4

8.9

7

8

13.7 0.0243

13.6 0.0105

CI for mean response and prediction intervals

Once useful regression exists, use it for prediction:

- To get a single number for prediction at a given x, substitute into regression equation, eg. age 10: predicted ATST is 646.48-14.04(10)=506 minutes.
- To express uncertainty of this prediction:
- CI for mean response expresses uncertainty about mean ATST for all children aged 10, based on data.
- Prediction interval expresses uncertainty about predicted ATST for a new child aged 10 whose ATST not known. More uncertain.
- Also do above for a child aged 5.

Intervals

Make new data frame with these values for age

```
my.age <- c(10, 5)
ages.new <- tibble(age = my.age)
ages.new

## # A tibble: 2 x 1</pre>
```

```
## # A tibble: 2 x 1

## age

## <dbl>

## 1 10

## 2 5
```

• Feed into predict:

```
pc <- predict(sleep.1, ages.new, interval = "c")
pp <- predict(sleep.1, ages.new, interval = "p")</pre>
```

The intervals

Confidence intervals for mean response:

```
cbind(ages.new, pc)
```

```
## age fit lwr upr
## 1 10 506.0729 497.5574 514.5883
## 2 5 576.2781 561.6578 590.8984
```

Prediction intervals for new response:

```
cbind(ages.new, pp)
```

```
## age fit lwr upr
## 1 10 506.0729 475.8982 536.2475
## 2 5 576.2781 543.8474 608.7088
```

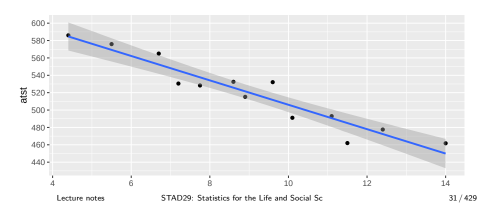
Comments

- Age 10 closer to centre of data, so intervals are both narrower than those for age 5.
- Prediction intervals bigger than CI for mean (additional uncertainty).
- Technical note: output from predict is R matrix, not data frame, so Tidyverse bind_cols does not work. Use base R cbind.

That grey envelope

Marks confidence interval for mean for all x:

```
ggplot(sleep, aes(x = age, y = atst)) + geom_point() +
geom_smooth(method = "lm") +
scale_y_continuous(breaks = seq(420, 600, 20))
```



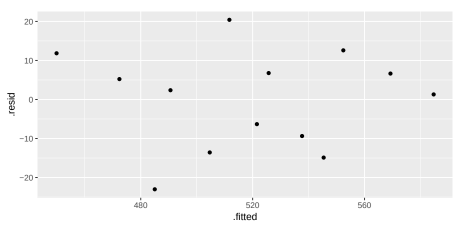
Diagnostics

How to tell whether a straight-line regression is appropriate?

- Before: check scatterplot for straight trend.
- After: plot residuals (observed minus predicted response) against predicted values. Aim: a plot with no pattern.

Residual plot

Not much pattern here — regression appropriate.



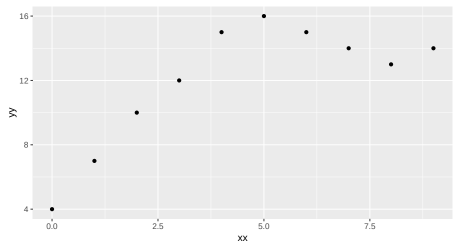
An inappropriate regression

Different data:

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/curvy.txt"
curvy <- read_delim(my_url, " ")
## Parsed with column specification:
## and and are all are all and are all and are all and are all all are all and are all and are all are all and are all are all all are all are all and are all are all and are all are all are all and are all all are al
```

```
## cols(
## xx = col_double(),
## yy = col_double()
## )
```

Scatterplot

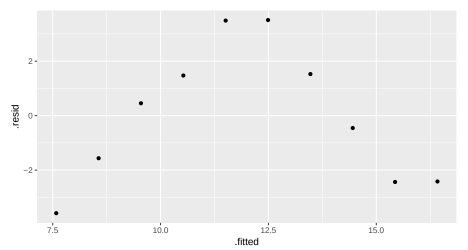


Regression line, anyway

```
curvy.1 \leftarrow lm(yy \sim xx, data = curvy)
summary(curvy.1)
##
## Call:
## lm(formula = yy ~ xx, data = curvy)
##
## Residuals:
##
     Min 1Q Median 3Q
                               Max
## -3.582 -2.204 0.000 1.514 3.509
##
## Coefficients:
##
             Estimate Std. Error t value Pr(>|t|)
## (Intercept) 7.5818 1.5616 4.855 0.00126 **
         ## xx
## ---
## Signif. codes:
## 0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' ' 1
##
## Residual standard error: 2.657 on 8 degrees of freedom
## Multiple R-squared: 0.5848, Adjusted R-squared: 0.5329
## F-statistic: 11.27 on 1 and 8 DF, p-value: 0.009984
```

Residual plot

ggplot(curvy.1, aes(x = .fitted, y = .resid)) + geom_point()



No good: fixing it up

- Residual plot has *curve*: middle residuals positive, high and low ones negative. Bad.
- Fitting a curve would be better. Try this:

```
curvy.2 <- lm(yy ~ xx + I(xx^2), data = curvy)</pre>
```

- Adding xx-squared term, to allow for curve.
- Another way to do same thing: specify how model changes:

```
curvy.2a <- update(curvy.1, . ~ . + I(xx^2))</pre>
```

Regression 2

tidy(curvy.2)

r.squared adj.r.squared sigma statistic p.value

<dbl> <dbl>

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0.936 0.983 66.8 2.75e-5

... with 5 more variables: logLik <dbl>, AIC <dbl>,
BIC <dbl>, deviance <dbl>, df.residual <int>

df

<dbl> <dbl> <int>

##

##

1

A tibble: 1 x 11

<dbl>

0.950

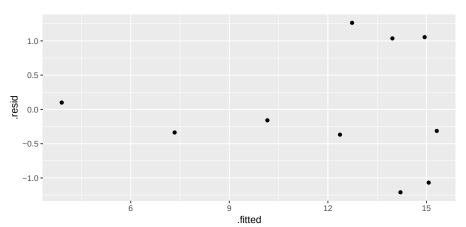
Comments

- xx-squared term definitely significant (P-value 0.000182), so need this curve to describe relationship.
- Adding squared term has made R-squared go up from 0.5848 to 0.9502: great improvement.
- This is a definite curve!

The residual plot now

No problems any more:

ggplot(curvy.2, aes(x = .fitted, y = .resid)) + geom_point()



Another way to handle curves

- Above, saw that changing x (adding x^2) was a way of handling curved relationships.
- Another way: change y (transformation).
- Can guess how to change y, or might be theory:
- example: relationship $y = ae^{bx}$ (exponential growth):
- take logs to get $\ln y = \ln a + bx$.
- ullet Taking logs has made relationship linear ($\ln y$ as response).
- Or, estimate transformation, using Box-Cox method.

Box-Cox

- Install package MASS via install.packages("MASS") (only need to do once)
- Every R session you want to use something in MASS, type library(MASS)

Some made-up data

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/madeup.csv"
madeup <- read_csv(my_url)</pre>
madeup
```

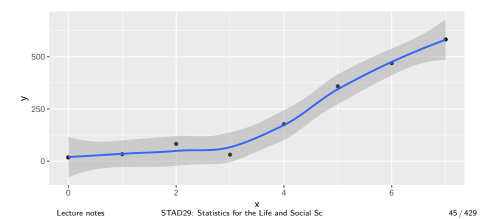
```
## # A tibble: 8 x 3
##
       row
                X
     <dbl> <dbl> <dbl>
##
## 1
                0 17.9
## 2
                1 33.6
         3
                2 82.7
## 3
                3 31.2
## 4
                4 177.
## 5
         5
         6
                5 359.
## 6
## 7
                6 469.
         8
                7 583.
## 8
```

Seems to be faster-than-linear growth, maybe exponential growth. STAD29: Statistics for the Life and Social Sc

Scatterplot: faster than linear growth

```
ggplot(madeup, aes(x = x, y = y)) + geom_point() +
  geom_smooth()
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'

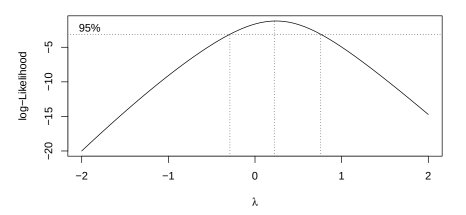


Running Box-Cox

- library(MASS) first.
- Feed boxcox a model formula with a squiggle in it, such as you would use for lm.
- Output: a graph (next page):

```
boxcox(y ~ x, data = madeup)
```

The Box-Cox output



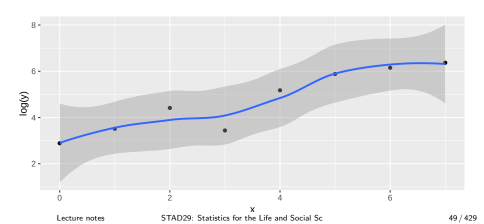
Comments

- λ (lambda) is the power by which you should transform y to get the relationship straight (straighter). Power 0 is "take logs"
- Middle dotted line marks best single value of λ (here about 0.1).
- Outer dotted lines mark 95% CI for λ , here -0.3 to 0.7, approx. (Rather uncertain about best transformation.)
- Any power transformation within the CI supported by data. In this case, $\log (\lambda = 0)$ and square root $(\lambda = 0.5)$ good, but no transformation $(\lambda = 1)$ not.
- Pick a "round-number" value of λ like 2, 1, 0.5, 0, -0.5, -1. Here 0 and 0.5 good values to pick.

Did transformation straighten things?

ullet Plot transformed y against x. Here, log:

```
ggplot(madeup, aes(x = x, y = log(y))) + geom_point() +
  geom_smooth()
```



Regression with transformed y

```
madeup.1 <- lm(log(y) ~ x, data = madeup)
glance(madeup.1)

## # A tibble: 1 x 11

## r.squared adj.r.squared sigma statistic p.value df

## <dbl> <dbl> <dbl> <dbl> <dbl> <int>
## 1 0.883 0.864 0.501 45.3 5.24e-4 2

## # ... with 5 more variables: logLik <dbl>, AIC <dbl>,
## BIC <dbl>, deviance <dbl>, df.residual <int>
tidy(madeup.1)
```

R-squared now decently high.

Multiple regression

- What if more than one x? Extra issues:
 - Now one intercept and a slope for each x: how to interpret?
 - Which x-variables actually help to predict y?
 - Different interpretations of "global" F-test and individual t-tests.
 - R-squared no longer correlation squared, but still interpreted as "higher better".
 - In 1m line, add extra xs after ~.
 - Interpretation not so easy (and other problems that can occur).

Multiple regression example

Study of women and visits to health professionals, and how the number of visits might be related to other variables:

timedrs: number of visits to health professionals (over course of study)

phyheal: number of physical health problems

menheal: number of mental health problems

stress: result of questionnaire about number and type of life changes timedrs response, others explanatory.

The data

##

)

```
my_url <-
  "http://www.utsc.utoronto.ca/~butler/d29/regressx.txt"
visits <- read delim(my url, " ")</pre>
## Parsed with column specification:
## cols(
##
     subjno = col_double(),
##
     timedrs = col_double(),
     phyheal = col_double(),
##
     menheal = col_double(),
##
```

stress = col_double()

Check data

visits

```
## # A tibble: 465 x 5
##
      subjno timedrs phyheal menheal stress
       <dbl>
                <dbl> <dbl>
                                   dbl>
##
                                           <dbl>
                               5
                                              265
##
                                        8
                     3
                                              415
##
                                        6
##
    3
                                        4
                                               92
                    13
                                        2
                                              241
##
                               3
##
    5
            5
                    15
                                        6
                                               86
##
    6
            6
                               5
                                        5
                                              247
                               5
##
                                        6
                                               13
##
            8
                                        5
                                               12
##
    9
            9
                               5
                                        4
                                              269
##
  10
           10
                                        9
                                              391
## # ... with 455 more rows
```

Fit multiple regression

```
visits.1 <- lm(timedrs ~ phyheal + menheal + stress,
 data = visits)
glance(visits.1)
## # A tibble: 1 x 11
##
    r.squared adj.r.squared sigma statistic p.value
                                                        df
        <dhl>
                      <dbl> <dbl>
                                      <dbl> <dbl> <int>
##
        0 219
                      0.214 9.71 43.0 1.56e-24
## 1
## # ... with 5 more variables: logLik <dbl>, AIC <dbl>,
      BIC <dbl>, deviance <dbl>, df.residual <int>
## #
```

The slopes

Model as a whole strongly significant even though R-sq not very big (lots of data). At least one of the x's predicts timedrs.

```
tidy(visits.1)
```

```
## # A tibble: 4 x 5
##
              estimate std.error statistic
    term
                                        p.value
    <chr>>
                 <dbl>
                          <dbl>
                                  <dbl>
                                          <dbl>
##
## 1 (Intercept) -3.70
                      1.12
                                -3.30 1.06e- 3
  2 phyheal 1.79 0.221
                                 8.08 5.60e-15
## 3 menheal
              -0.00967 0.129 -0.0749 9.40e- 1
                                        1.85e- 4
## 4 stress
               0.0136
                        0.00361
                                 3.77
```

The physical health and stress variables initely help to predict the number of visits, but with those in the model we don't need menheal. However, look at prediction of timedrs from menheal by itself:

visits.2 <- lm(timedrs ~ menheal, data = visits)</pre>

Just menheal

```
## # A tibble: 1 x 11
## r.squared adj.r.squared sigma statistic p.value df
## <dbl> <dbl> <dbl> <dbl> <dbl> <int>
## 1 0.0653 0.0633 10.6 32.4 2.28e-8 2
## # ... with 5 more variables: logLik <dbl>, AIC <dbl>,
## BIC <dbl>, deviance <dbl>, df.residual <int>
tidy(visits.2)
```

menheal by itself

- menheal by itself does significantly help to predict timedrs.
- But the R-sq is much less (6.5% vs. 22%).
- So other two variables do a better job of prediction.
- With those variables in the regression (phyheal and stress), don't need menheal as well.

Investigating via correlation

Leave out first column (subjno):

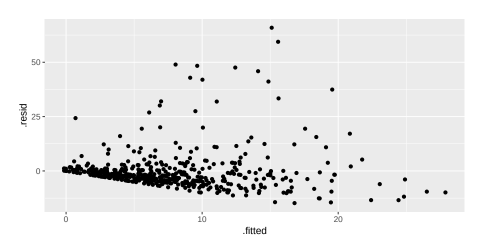
```
visits %>% select(-subjno) %>% cor()
```

```
## timedrs phyheal menheal stress
## timedrs 1.0000000 0.4395293 0.2555703 0.2865951
## phyheal 0.4395293 1.0000000 0.5049464 0.3055517
## menheal 0.2555703 0.5049464 1.0000000 0.3697911
## stress 0.2865951 0.3055517 0.3697911 1.0000000
```

- phyheal most strongly correlated with timedrs.
- Not much to choose between other two.
- But menheal has higher correlation with phyheal, so not as much to add to prediction as stress.
- Goes to show things more complicated in multiple regression.

Residual plot (from timedrs on all)

ggplot(visits.1, aes(x = .fitted, y = .resid)) + geom_point()

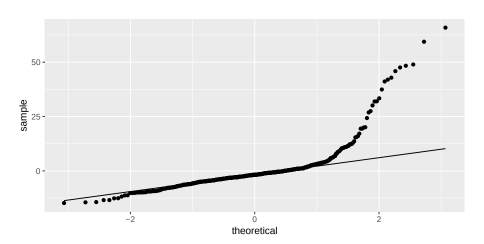


Comment

Apparently random. But...

Normal quantile plot of residuals

ggplot(visits.1, aes(sample = .resid)) + stat_qq() + stat_qq__

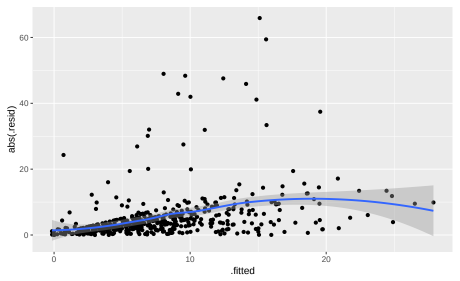


Absolute residuals

Is there trend in *size* of residuals (fan-out)? Plot *absolute value* of residual against fitted value (graph next page):

```
g <- ggplot(visits.1, aes(x = .fitted, y = abs(.resid))) +
  geom_point() + geom_smooth()</pre>
```

The plot



Comments

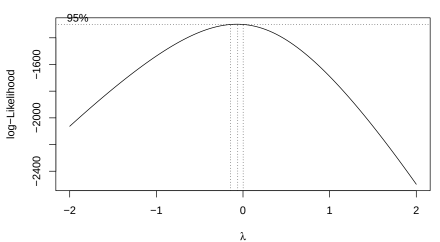
- On the normal quantile plot:
 - highest (most positive) residuals are way too high
 - distribution of residuals skewed to right (not normal at all)
- On plot of absolute residuals:
 - size of residuals getting bigger as fitted values increase
 - predictions getting more variable as fitted values increase
 - that is, predictions getting *less accurate* as fitted values increase, but predictions should be equally accurate all way along.
- Both indicate problems with regression, of kind that transformation of response often fixes: that is, predict function of response timedrs instead of timedrs itself.

Box-Cox transformations

- Taking log of timedrs and having it work: lucky guess. How to find good transformation?
- Box-Cox again.
- Extra problem: some of timedrs values are 0, but Box-Cox expects all
 Note response for boxcox:

```
boxcox(timedrs + 1 ~ phyheal + menheal + stress, data = visits
```

Try 1



Comments on try 1

- Best: λ just less than zero.
- Hard to see scale.
- Focus on λ in (-0.3, 0.1):

```
my.lambda <- seq(-0.3, 0.1, 0.01)
my.lambda
```

```
## [1] -0.30 -0.29 -0.28 -0.27 -0.26 -0.25 -0.24 -0.23 -0.22

## [10] -0.21 -0.20 -0.19 -0.18 -0.17 -0.16 -0.15 -0.14 -0.13

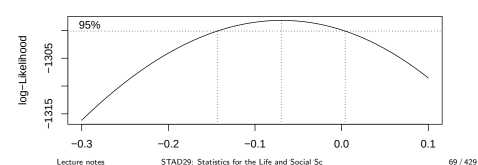
## [19] -0.12 -0.11 -0.10 -0.09 -0.08 -0.07 -0.06 -0.05 -0.04

## [28] -0.03 -0.02 -0.01 0.00 0.01 0.02 0.03 0.04 0.05

## [37] 0.06 0.07 0.08 0.09 0.10
```

Try 2

```
boxcox(timedrs + 1 ~ phyheal + menheal + stress,
  lambda = my.lambda,
  data = visits
)
```



Comments

- Best: λ just about -0.07.
- CI for λ about (-0.14, 0.01).
- Only nearby round number: $\lambda = 0$, log transformation.

Fixing the problems

- Try regression again, with transformed response instead of original one.
- Then check residual plot to see that it is OK now.

```
visits.3 <- lm(log(timedrs + 1) ~ phyheal + menheal + stress,
  data = visits
)</pre>
```

- timedrs+1 because some timedrs values 0, can't take log of 0.
- Won't usually need to worry about this, but when response could be zero/negative, fix that before transformation.

Output

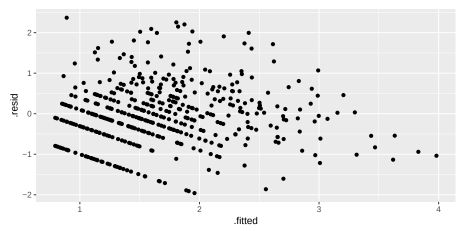
```
summary(visits.3)
##
## Call:
## lm(formula = log(timedrs + 1) ~ phyheal + menheal + stress, data = visits)
##
## Residuals:
##
       Min
              10 Median
                                 30
                                         Max
## -1.95865 -0.44076 -0.02331 0.42304 2.36797
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 0.3903862 0.0882908 4.422 1.22e-05 ***
## phyheal 0.2019361 0.0173624 11.631 < 2e-16 ***
## menheal 0.0071442 0.0101335 0.705 0.481
## stress 0.0013158 0.0002837 4.638 4.58e-06 ***
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.7625 on 461 degrees of freedom
## Multiple R-squared: 0.3682, Adjusted R-squared: 0.3641
## F-statistic: 89.56 on 3 and 461 DF. p-value: < 2.2e-16
```

Comments

- Model as a whole strongly significant again
- R-sq higher than before (37% vs. 22%) suggesting things more linear now
- Same conclusion re menheal: can take out of regression.
- Should look at residual plots (next pages). Have we fixed problems?

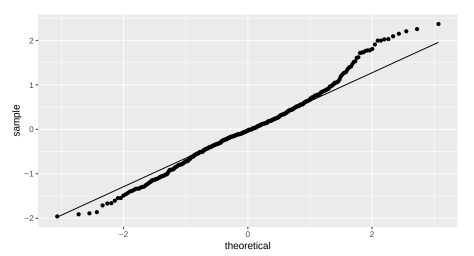
Residuals against fitted values

```
ggplot(visits.3, aes(x = .fitted, y = .resid)) +
  geom_point()
```



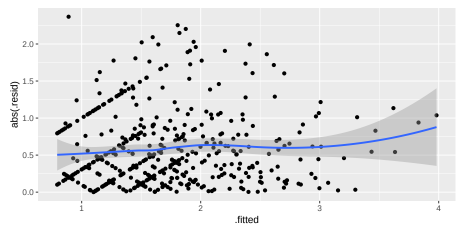
Normal quantile plot of residuals

ggplot(visits.3, aes(sample = .resid)) + stat_qq() + stat_qq__



Absolute residuals against fitted

```
ggplot(visits.3, aes(x = .fitted, y = abs(.resid))) +
  geom_point() + geom_smooth()
```



Comments

- Residuals vs. fitted looks a lot more random.
- Normal quantile plot looks a lot more normal (though still a little right-skewness)
- Absolute residuals: not so much trend (though still some).
- Not perfect, but much improved.

Testing more than one x at once

- The t-tests test only whether one variable could be taken out of the regression you're looking at.
- To test significance of more than one variable at once, fit model with and without variables
 - then use anova to compare fit of models:

Results of tests

```
anova(visits.6, visits.5)
```

```
## Analysis of Variance Table
##
## Model 1: log(timedrs + 1) ~ stress
## Model 2: log(timedrs + 1) ~ phyheal + menheal + stress
## Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 463 371.47
## 2 461 268.01 2 103.46 88.984 < 2.2e-16 ***
## ---
## Signif. codes:</pre>
```

- ## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

 Models don't fit equally well, so bigger one fits better.
 - Or "taking both variables out makes the fit worse, so don't do it".
 - Taking out those x's is a mistake. Or putting them in is a good idea.

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The punting data

Data set punting.txt contains 4 variables for 13 right-footed football kickers (punters): left leg and right leg strength (lbs), distance punted (ft), another variable called "fred". Predict punting distance from other variables:

| left | right | punt | fred |
|------|-------|--------|------|
| 170 | 170 | 162.50 | 171 |
| 130 | 140 | 144.0 | 136 |
| 170 | 180 | 174.50 | 174 |
| 160 | 160 | 163.50 | 161 |
| 150 | 170 | 192.0 | 159 |
| 150 | 150 | 171.75 | 151 |
| 180 | 170 | 162.0 | 174 |
| 110 | 110 | 104.83 | 111 |
| 110 | 120 | 105.67 | 114 |
| 120 | 130 | 117.58 | 126 |
| 140 | 120 | 140.25 | 129 |
| 130 | 140 | 150.17 | 136 |
| 150 | 160 | 165.17 | 154 |

Reading in

• Separated by multiple spaces with columns lined up:

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/punting.txt
punting <- read_table(my_url)

## Parsed with column specification:
## cols(
## left = col_double(),
## right = col_double(),
## punt = col_double(),
## fred = col_double()</pre>
```

)

The data

punting

```
A tibble: 13 x 4
##
        left right
                             fred
                     punt
       <dbl> <dbl> <dbl> <dbl>
##
         170
                170
                      162.
##
    1
                              171
         130
                      144
                              136
##
    2
                140
    3
         170
##
                180
                      174.
                              174
##
    4
         160
                160
                      164.
                              161
##
    5
         150
                170
                      192
                              159
##
    6
         150
                150
                      172.
                              151
##
    7
         180
                170
                      162
                              174
##
    8
         110
                110
                      105.
                              111
##
    9
         110
                120
                      106.
                              114
##
   10
         120
                130
                      118.
                              126
## 11
         140
                120
                      140.
                              129
## 12
         130
                140
                      150.
                              136
## 13
         150
                160
                      165.
                              154
```

Regression and output

```
punting.1 <- lm(punt ~ left + right + fred, data = punting)
glance(punting.1)

## # A tibble: 1 x 11

## r.squared adj.r.squared sigma statistic p.value df

## <dbl> <dbl> <dbl> <dbl> <dbl> <int>
## 1 0.778 0.704 14.7 10.5 0.00267 4

## # ... with 5 more variables: logLik <dbl>, AIC <dbl>,
## # BIC <dbl>, deviance <dbl>, df.residual <int>
```

```
## # A tibble: 4 x 5
```

tidy(punting.1)

STAD29: Statistics for the Life and Social Sc.

Comments

- Overall regression strongly significant, R-sq high.
- None of the x's significant! Why?
- *t*-tests only say that you could take any one of the *x*'s out without damaging the fit; doesn't matter which one.
- Explanation: look at correlations.

The correlations

cor(punting)

```
## left right punt fred
## left 1.0000000 0.8957224 0.8117368 0.9722632
## right 0.8957224 1.0000000 0.8805469 0.9728784
## punt 0.8117368 0.8805469 1.0000000 0.8679507
## fred 0.9722632 0.9728784 0.8679507 1.0000000
```

- All correlations are high: x's with punt (good) and with each other (bad, at least confusing).
- What to do? Probably do just as well to pick one variable, say right since kickers are right-footed.

Just right

```
punting.2 <- lm(punt ~ right, data = punting)
anova(punting.2, punting.1)</pre>
```

```
##
## Model 1: punt ~ right
## Model 2: punt ~ left + right + fred
## Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 11 1962.5
## 2 9 1938.2 2 24.263 0.0563 0.9456
```

Analysis of Variance Table

No significant loss by dropping other two variables.

Comparing R-squareds

```
summary(punting.1)$r.squared
```

```
## [1] 0.7781401
```

summary(punting.2)\$r.squared

```
## [1] 0.7753629
```

Basically no difference. In regression (over), right significant:

Regression results

tidy(punting.2)

```
## # A tibble: 2 x 5
               estimate std.error statistic
##
                                            p.value
    term
    <chr>>
                                              <dbl>
##
                  <dbl>
                           <dbl>
                                    <dbl>
  1 (Intercept)
               -3.69 25.3 -0.146 0.886
## 2 right
                   1.04
                          0.169 6.16 0.0000709
```

But...

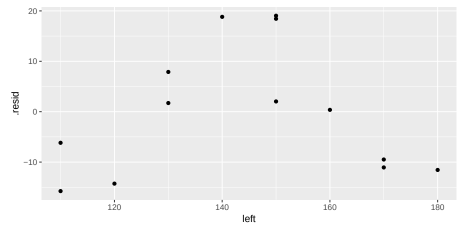
- Maybe we got the *form* of the relationship with left wrong.
- Check: plot *residuals* from previous regression (without left) against left.
- Residuals here are "punting distance adjusted for right leg strength".
- If there is some kind of relationship with left, we should include in model.
- Plot of residuals against original variable: augment from broom.

Augmenting punting.2

```
punting.2 %>% augment(punting) -> punting.2.aug
punting.2.aug %>% slice(1:8)
## # A tibble: 8 x 11
##
     left right punt fred .fitted .se.fit .resid
                                                .hat.
    <dbl> <dbl> <dbl> <dbl> <
                                         <dbl> <dbl>
##
                           <dbl>
                                  <dbl>
                     171 174. 5.29 -11.1
## 1
      170
           170
               162.
                                              0.157
## 2
     130
          140 144 136 142. 3.93 1.72
                                              0.0864
## 3
     170
          180 174. 174 184. 6.60 -9.49
                                              0.244
## 4
      160
           160 164. 161
                            163. 4.25 0.366 0.101
## 5
     150
          170
               192 159 174. 5.29 18.4
                                              0.157
## 6
      150
           150 172. 151
                           153. 3.73 19.0 0.0778
## 7
      180
           170
               162 174 174. 5.29 -11.6 0.157
      110
           110
               105.
                      111
                                   7.38 - 6.17
                                              0.305
## 8
                            111.
## # ...
     with 3 more variables: .sigma <dbl>, .cooksd <dbl>,
## #
      .std.resid <dbl>
```

Residuals against left

```
ggplot(punting.2.aug, aes(x = left, y = .resid)) +
  geom_point()
```



Comments

- There is a curved relationship with left.
- We should add left-squared to the regression (and therefore put left back in when we do that):

```
punting.3 <- lm(punt ~ left + I(left^2) + right,
  data = punting
)</pre>
```

Regression with left-squared

```
summary(punting.3)
##
## Call:
## lm(formula = punt ~ left + I(left^2) + right, data = punting)
##
## Residuals:
##
      Min 10 Median 30
                                        Max
## -11.3777 -5.3599 0.0459 4.5088 13.2669
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -4.623e+02 9.902e+01 -4.669 0.00117 **
## left 6.888e+00 1.462e+00 4.710 0.00110 **
## I(left^2) -2.302e-02 4.927e-03 -4.672 0.00117 **
## right 7.396e-01 2.292e-01 3.227 0.01038 *
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 7.931 on 9 degrees of freedom
## Multiple R-squared: 0.9352, Adjusted R-squared: 0.9136
## F-statistic: 43.3 on 3 and 9 DF. p-value: 1.13e-05
```

Comments

- This was definitely a good idea (R-squared has clearly increased).
- We would never have seen it without plotting residuals from punting.2 (without left) against left.
- Negative slope for leftsq means that increased left-leg strength only increases punting distance up to a point: beyond that, it decreases again.

Logistic regression (ordinal/nominal response)

Section 3

Logistic regression (ordinal/nominal response)

Logistic regression

- When response variable is measured/counted, regression can work well.
- But what if response is yes/no, lived/died, success/failure?
- Model probability of success.
- Probability must be between 0 and 1; need method that ensures this.
- Logistic regression does this. In R, is a generalized linear model with binomial "family":

```
glm(y ~ x, family="binomial")
```

Begin with simplest case.

Packages

```
library(MASS)
library(tidyverse)
library(broom)
library(nnet)
```

The rats, part 1

• Rats given dose of some poison; either live or die:

dose status

- 0 lived
- 1 died
- 2 lived
- 3 lived
- 4 died
- 5 died

rats <- read delim(my url, " ")

Parsed with column specification:

Read in:

cols(

```
## dose = col_double(),
## status = col_character()
## )
glimpse(rats)

## Observations: 6
## Variables: 2
## $ dose <dbl> 0, 1, 2, 3, 4, 5
```

\$ status <chr> "lived", "died", "lived", "lived", "died", ...

my url <- "http://www.utsc.utoronto.ca/~butler/d29/rat.txt"

Basic logistic regression

• Make response into a factor first:

```
rats2 <- rats %>% mutate(status = factor(status))
```

• then fit model:

```
status.1 <- glm(status ~ dose, family = "binomial", data = rats2)
```

Output

```
summary(status.1)
##
## Call:
## glm(formula = status ~ dose, family = "binomial", data = rats2)
##
## Deviance Residuals:
##
   0.5835 -1.6254 1.0381 1.3234 -0.7880 -0.5835
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) 1.6841 1.7979 0.937 0.349
## dose -0.6736 0.6140 -1.097 0.273
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 8.3178 on 5 degrees of freedom
##
## Residual deviance: 6.7728 on 4 degrees of freedom
## ATC: 10.773
##
## Number of Fisher Scoring iterations: 4
```

Interpreting the output

- ullet Like (multiple) regression, get tests of significance of individual x's
- Here not significant (only 6 observations).
- "Slope" for dose is negative, meaning that as dose increases, probability of event modelled (survival) decreases.

Output part 2: predicted survival probs

```
p <- predict(status.1, type = "response")
cbind(rats, p)</pre>
```

```
## 1 0 lived 0.8434490

## 2 1 died 0.7331122

## 3 2 lived 0.5834187

## 4 3 lived 0.4165813

## 5 4 died 0.2668878

## 6 5 died 0.1565510
```

dose status

##

The rats, more

- More realistic: more rats at each dose (say 10).
- Listing each rat on one line makes a big data file.
- Use format below: dose, number of survivals, number of deaths.

| dose | lived | died |
|------|-------|------|
| 0 | 10 | 0 |
| 1 | 7 | 3 |
| 2 | 6 | 4 |
| 3 | 4 | 6 |
| 4 | 2 | 8 |
| 5 | 1 | 9 |

- 6 lines of data correspond to 60 actual rats.
- Saved in rat2.txt.

These data

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/rat2.txt"
rat2 <- read_delim(my_url, " ")

## Parsed with column specification:
## cols(
## dose = col_double(),
## lived = col_double(),
## died = col_double()
## )
rat2</pre>
```

```
## # A tibble: 6 x 3
## dose lived died
## <dbl> <dbl> <dbl> <dbl> 
## 1 0 10 0
## 2 1 7 3
## 3 2 6 4
## 4 3 4 6
## 5 4 2 8
## 6 5 1 9
```

Create response matrix:

- Each row contains *multiple* observations.
- Create two-column response:
 - #survivals in first column,
 - #deaths in second.

```
response <- with(rat2, cbind(lived, died))
response</pre>
```

```
## lived died
## [1,] 10 0
## [2,] 7 3
## [3,] 6 4
## [4,] 4 6
## [5,] 2 8
## [6,] 1 9
```

Response is R matrix:

```
class(response)
```

Fit logistic regression

using response you just made:

```
rat2.1 <- glm(response ~ dose,
  family = "binomial",
  data = rat2
)</pre>
```

Output

```
summarv(rat2.1)
##
## Call:
## glm(formula = response ~ dose, family = "binomial", data = rat2)
##
## Deviance Residuals:
##
## 1.3421 -0.7916 -0.1034 0.1034 0.0389 0.1529
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) 2.3619 0.6719 3.515 0.000439 ***
## dose -0.9448 0.2351 -4.018 5.87e-05 ***
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 27.530 on 5 degrees of freedom
## Residual deviance: 2.474 on 4 degrees of freedom
## AIC: 18.94
```

Predicted survival probs

```
p <- predict(rat2.1, type = "response")
cbind(rat2, p)</pre>
```

```
dose lived died
##
        0
              10
                     0 0.9138762
## 1
                    3 0.8048905
## 3
               6
                     4 0.6159474
                     6 0.3840526
## 4
                    8 0.1951095
## 5
        4
        5
                     9 0.0861238
## 6
```

Comments

- Significant effect of dose.
- Effect of larger dose is to decrease survival probability ("slope" negative; also see in decreasing predictions.)

Multiple logistic regression

- With more than one x, works much like multiple regression.
- Example: study of patients with blood poisoning severe enough to warrant surgery. Relate survival to other potential risk factors.
- Variables, 1=present, 0=absent:
 - survival (death from sepsis=1), response
 - shock
 - malnutrition
 - alcoholism
 - age (as numerical variable)
 - bowel infarction
- See what relates to death.

Read in data

```
my url <-
  "http://www.utsc.utoronto.ca/~butler/d29/sepsis.txt"
sepsis <- read delim(my url, " ")
## Parsed with column specification:
## cols(
##
     death = col double(),
##
     shock = col double(),
     malnut = col double(),
##
##
     alcohol = col double(),
##
     age = col_double(),
##
     bowelinf = col double()
## )
```

The data

sepsis

```
A tibble: 106 x 6
##
      death shock malnut alcohol age bowelinf
##
      <dbl> <dbl> <dbl>
                              <dbl> <dbl>
                                                <dbl>
##
                                   0
                                         56
                                         80
##
    3
           0
                                         61
##
           0
                                         26
##
##
    5
           0
                                         53
                                         87
##
    6
           0
                                         21
##
                                         69
##
    8
                                         57
##
    9
   10
                                   0
                                         76
##
     ... with 96
                more rows
```

Fit model

```
sepsis.1 <- glm(death ~ shock + malnut + alcohol + age +
  bowelinf,
family = "binomial",
data = sepsis
)</pre>
```

Output part 1

tidy(sepsis.1)

```
## # A tibble: 6 x 5
##
     term
                 estimate std.error statistic p.value
     <chr>
                    <dbl>
                               <dbl>
                                         <dbl>
                                                  <dbl>
##
  1 (Intercept)
                  -9.75
                              2.54
                                         -3.84 0.000124
  2 shock
                   3.67
                             1.16
                                          3.15 0.00161
                   1.22
                             0.728
  3 malnut
                                          1.67 0.0948
                   3.35
                             0.982
                                          3.42 0.000635
## 4 alcohol
                   0.0922
                             0.0303
  5 age
                                          3.04 0.00237
                   2.80
                              1.16
                                          2.40 0.0162
## 6 bowelinf
```

- All P-values fairly small
- but malnut not significant: remove.

Removing malnut

```
sepsis.2 <- update(sepsis.1, . ~ . - malnut)
tidy(sepsis.2)</pre>
```

```
## # A tibble: 5 x 5
##
    term
                 estimate std.error statistic p.value
    <chr>>
                    <dbl>
                              <dbl>
                                        <dbl>
                                                 <dbl>
##
  1 (Intercept) -8.89
                             2.32
                                        -3.84 0.000124
  2 shock
                 3.70
                            1.10
                                         3.35 0.000797
##
                  3.19
                            0.917
                                         3.47 0.000514
  3 alcohol
                  0.0898
                            0.0292
                                         3.07 0.00211
  4 age
                             1.07
                                         2.23 0.0260
## 5 bowelinf
                  2.39
```

Everything significant now.

Comments

- Most of the original x's helped predict death. Only malnut seemed not to add anything.
- Removed malnut and tried again.
- Everything remaining is significant (though bowelinf actually became less significant).
- All coefficients are *positive*, so having any of the risk factors (or being older) *increases* risk of death.

Predictions from model without "malnut"

A few chosen at random:

```
sepsis.pred <- predict(sepsis.2, type = "response")
d <- data.frame(sepsis, sepsis.pred)
myrows <- c(4, 1, 2, 11, 32)
slice(d, myrows)</pre>
```

```
##
     death shock malnut alcohol age bowelinf sepsis.pred
                                              0 0.001415347
## 1
         0
                0
                                   26
                                              0.020552383
## 2
                0
                                   56
## 3
                0
                                   80
                                              0 0.153416834
                                   66
                                              1 0.931290137
## 4
                0
                                   49
                                              0.0.213000997
## 5
                0
```

Comments

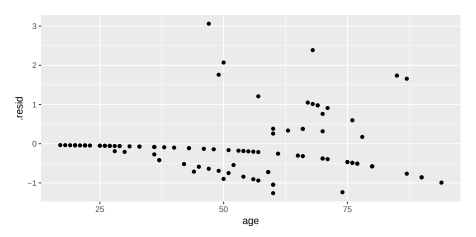
- Survival chances pretty good if no risk factors, though decreasing with age.
- Having more than one risk factor reduces survival chances dramatically.
- Usually good job of predicting survival; sometimes death predicted to survive.

Assessing proportionality of odds for age

- An assumption we made is that log-odds of survival depends linearly on age.
- Hard to get your head around, but basic idea is that survival chances go continuously up (or down) with age, instead of (for example) going up and then down.
- In this case, seems reasonable, but should check:

Residuals vs. age

```
ggplot(augment(sepsis.2), aes(x = age, y = .resid)) +
  geom_point()
```



Comments

- No apparent problems overall.
- Confusing "line" across: no risk factors, survived.

Probability and odds

• For probability p, odds is p/(1-p):

| Prob. | Odds | log-odds | in words |
|-------|------------------------|----------|--------------|
| 0.5 | 0.5/0.5 = 1/1 = 1.00 | 0.00 | "even money" |
| 0.1 | 0.1/0.9 = 1/9 = 0.11 | -2.20 | "9 to 1" |
| 0.4 | 0.4/0.6 = 1/1.5 = 0.67 | -0.41 | "1.5 to 1" |
| 0.8 | 0.8/0.2 = 4/1 = 4.00 | 1.39 | "4 to 1 on" |

- Gamblers use odds: if you win at 9 to 1 odds, get original stake back plus 9 times the stake.
- Probability has to be between 0 and 1
- Odds between 0 and infinity
- Log-odds can be anything: any log-odds corresponds to valid probability.

Odds ratio

- Suppose 90 of 100 men drank wine last week, but only 20 of 100 women.
- Prob of man drinking wine 90/100 = 0.9, woman 20/100 = 0.2.
- Odds of man drinking wine 0.9/0.1 = 9, woman 0.2/0.8 = 0.25.
- Ratio of odds is 9/0.25 = 36.
- Way of quantifying difference between men and women: "odds of drinking wine 36 times larger for males than females".

Sepsis data again

• Recall prediction of probability of death from risk factors:

```
sepsis.2.tidy <- tidy(sepsis.2)
sepsis.2.tidy</pre>
```

```
## # A tibble: 5 \times 5
##
                 estimate std.error statistic p.value
    term
                                                 <dhl>
##
    <chr>
                    <dbl>
                              <dbl>
                                        <dbl>
                             2.32
                                        -3.840.000124
## 1 (Intercept) -8.89
## 2 shock
                  3.70
                             1 10
                                         3.35 0.000797
                             0.917
                                         3.47 0.000514
## 3 alcohol
                  3.19
                   0.0898
                             0.0292
                                         3.07 0.00211
## 4 age
## 5 bowelinf
                   2.39
                             1.07
                                         2.23 0.0260
```

Slopes in column estimate.

Multiplying the odds

Can interpret slopes by taking "exp" of them. We ignore intercept.

```
sepsis.2.tidy %>%
mutate(exp_coeff=exp(estimate)) %>%
select(term, exp_coeff)
```

Interpretation

- These say "how much do you multiply odds of death by for increase of 1 in corresponding risk factor?" Or, what is odds ratio for that factor being 1 (present) vs. 0 (absent)?
- Eg. being alcoholic vs. not increases odds of death by 24 times
- One year older multiplies odds by about 1.1 times. Over 40 years, about $1.09^{40}=31$ times.

Odds ratio and relative risk

- Relative risk is ratio of probabilities.
- Above: 90 of 100 men (0.9) drank wine, 20 of 100 women (0.2).
- Relative risk 0.9/0.2=4.5. (odds ratio was 36).
- When probabilities small, relative risk and odds ratio similar.
- Eg. prob of man having disease 0.02, woman 0.01.
- Relative risk 0.02/0.01 = 2.

Odds ratio vs. relative risk

Odds for men and for women:

```
(od1 <- 0.02 / 0.98) # men
```

```
## [1] 0.02040816
```

```
(od2 <- 0.01 / 0.99) # women
```

```
## [1] 0.01010101
```

Odds ratio

```
od1 / od2
```

```
## [1] 2.020408
```

• Very close to relative risk of 2.

More than 2 response categories

- With 2 response categories, model the probability of one, and prob of other is one minus that. So doesn't matter which category you model.
- With more than 2 categories, have to think more carefully about the categories: are they
- ordered: you can put them in a natural order (like low, medium, high)
- nominal: ordering the categories doesn't make sense (like red, green, blue).
- R handles both kinds of response; learn how.

Ordinal response: the miners

- Model probability of being in given category or lower.
- Example: coal-miners often suffer disease pneumoconiosis. Likelihood of disease believed to be greater among miners who have worked longer.
- Severity of disease measured on categorical scale: none, moderate, 3 severe.

Miners data

• Data are frequencies:

| Exposure | ${\tt None}$ | Moderate | Severe |
|----------|--------------|----------|--------|
| 5.8 | 98 | 0 | 0 |
| 15.0 | 51 | 2 | 1 |
| 21.5 | 34 | 6 | 3 |
| 27.5 | 35 | 5 | 8 |
| 33.5 | 32 | 10 | 9 |
| 39.5 | 23 | 7 | 8 |
| 46.0 | 12 | 6 | 10 |
| 51.5 | 4 | 2 | 5 |

Reading the data

Data in aligned columns with more than one space between, so:

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/miners-tab.txt"
freqs <- read_table(my_url)
## Parsed with column specification:
## cols(</pre>
```

```
## Cols(
## Exposure = col_double(),
## None = col_double(),
## Moderate = col_double(),
## Severe = col_double()
## )
```

The data

freqs

```
A tibble: 8 x 4
##
     Exposure None Moderate Severe
##
         <dbl> <dbl>
                          <dbl>
                                  <dbl>
## 1
           5.8
                   98
                   51
## 2
          15
## 3
          21.5
                   34
                              6
                                      3
          27.5
                   35
                              5
                                      8
## 4
## 5
          33.5
                   32
                             10
          39.5
                   23
          46
                   12
                              6
                                     10
          51.5
                                      5
## 8
                    4
```

Tidying and row proportions

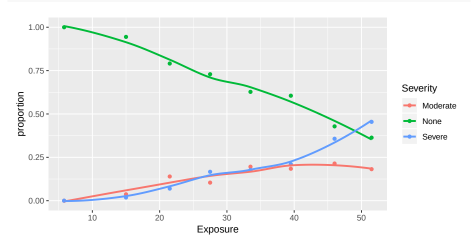
```
freqs %>%
  gather(Severity, Freq, None:Severe) %>%
  group_by(Exposure) %>%
  mutate(proportion = Freq / sum(Freq)) -> miners
```

Result

miners

```
## # A tibble: 24 \times 4
     Groups:
                Exposure [8]
      Exposure Severity Freq proportion
##
##
          <dbl> <chr>
                           <dbl>
                                       <dbl>
##
            5.8 None
                              98
                                      1
    1
##
           15
                None
                              51
                                      0.944
##
    3
           21.5 None
                              34
                                      0.791
##
           27.5 None
                              35
                                      0.729
##
           33.5 None
                              32
                                      0.627
##
    6
           39.5 None
                              23
                                      0.605
                              12
                                      0.429
##
           46
                None
##
    8
           51.5 None
                                      0.364
##
    9
            5.8 Moderate
                               0
                                      0
                Moderate
                                      0.0370
##
   10
           15
   # ... with 14 more rows
```

Plot proportions against exposure



Reminder of data setup

miners

```
# A tibble: 24 \times 4
   # Groups:
               Exposure [8]
##
      Exposure Severity Freq proportion
          <dbl> <chr>
                          <dbl>
                                      <dbl>
##
            5.8 None
                             98
##
                             51
                                     0.944
##
           15
                None
           21.5 None
                             34
                                     0.791
##
##
          27.5 None
                             35
                                     0.729
          33.5 None
                             32
                                     0.627
##
##
          39.5 None
                             23
                                     0.605
                             12
##
          46
                None
                                     0.429
           51.5 None
                                     0.364
##
##
            5.8 Moderate
                                     0
           15
                Moderate
                                     0.0370
     ... with 14 more rows
```

Creating an ordered factor

- Problem: on plot, Severity categories in wrong order.
- In the data frame, categories in correct order.
- Package forcats (in tidyverse) has functions for creating factors to specifications.
- fct_inorder takes levels in order they appear in data:

```
miners %>%
mutate(sev_ord = fct_inorder(Severity)) -> miners
```

To check:

```
levels(miners$sev_ord)
```

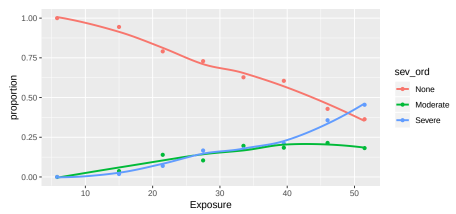
```
## [1] "None" "Moderate" "Severe"
```

New data frame

miners

```
## # A tibble: 24 \times 5
     Groups:
             Exposure
                          [8]
      Exposure Severity Freq proportion sev_ord
##
##
         <dbl> <chr>
                          <dbl>
                                      <dbl> <fct>
##
           5.8 None
                             98
    1
                                     1
                                             None
##
          15
                None
                             51
                                     0.944
                                            None
##
    3
          21.5 None
                             34
                                     0.791
                                            None
##
          27.5 None
                             35
                                     0.729
                                            None
                                     0.627
##
          33.5 None
                             32
                                            None
##
    6
          39.5 None
                             23
                                     0.605
                                            None
                             12
##
          46
                None
                                     0.429
                                            None
##
    8
          51.5 None
                                     0.364
                                            None
##
    9
           5.8 Moderate
                              0
                                     0
                                            Moderate
##
   10
           15
                Moderate
                                     0.0370 Moderate
   # ... with 14 more rows
```

Improved plot



Fitting ordered logistic model

Use function polr from package MASS. Like glm.

```
sev.1 <- polr(sev_ord ~ Exposure,
  weights = Freq,
  data = miners
)</pre>
```

Output: not very illuminating

```
summary(sev.1)
##
## Re-fitting to get Hessian
## Call:
## polr(formula = sev_ord ~ Exposure, data = miners, weights = Freq)
##
## Coefficients:
##
            Value Std. Error t value
## Exposure 0.0959 0.01194
                               8.034
##
  Intercepts:
##
                  Value Std. Error t value
## None | Moderate 3.9558 0.4097 9.6558
## Moderate|Severe 4.8690 0.4411 11.0383
##
## Residual Deviance: 416.9188
## AIC: 422.9188
```

Does exposure have an effect?

Fit model without Exposure, and compare using anova. Note 1 for model with just intercept:

```
sev.0 <- polr(sev_ord ~ 1, weights = Freq, data = miners)
anova(sev.0, sev.1)
## Likelihood ratio tests of ordinal regression models</pre>
```

Exposure definitely has effect on severity of disease.

##

Another way

• What (if anything) can we drop from model with exposure?

```
drop1(sev.1, test = "Chisq")
```

```
## Single term deletions
##
## Model:
## sev_ord ~ Exposure
##
           Df
              AIC LRT Pr(>Chi)
## <none> 422.92
## Exposure 1 509.16 88.243 < 2.2e-16 ***
## ---
## Signif. codes:
    0 '***' 0.001 '**' 0.01 '*' 0.05
##
## '.' 0.1 ' ' 1
```

Nothing. Exposure definitely has effect.

Predicted probabilities

Make new data frame out of all the exposure values (from original data frame), and predict from that:

```
sev.new <- tibble(Exposure = freqs$Exposure)</pre>
pr <- predict(sev.1, sev.new, type = "p")</pre>
miners.pred <- cbind(sev.new, pr)
miners.pred
```

```
##
                    None Moderate
                                           Severe
     Exposure
           5.8 0.9676920 0.01908912 0.01321885
## 1
          15.0 0.9253445 0.04329931 0.03135614
## 2
## 3
         21.5 0.8692003 0.07385858 0.05694115
## 4
         27.5 0.7889290 0.11413004 0.09694093
         33.5 0.6776641 0.16207145 0.16026444
## 5
## 6
         39.5 0.5418105 0.20484198 0.25334756
## 7
         46.0 0.3879962 0.22441555 0.38758828
          51.5 0.2722543 0.21025011 0.51749563
## 8
      Lecture notes
                     STAD29: Statistics for the Life and Social Sc.
```

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Comments

- Model appears to match data: as exposure goes up, prob of None goes down, Severe goes up (sharply for high exposure).
- Like original data frame, this one nice to look at but *not tidy*. We want to make graph, so tidy it.
- Also want the severity values in right order.
- Usual gather, plus a bit:

```
miners.pred %>%
gather(Severity, probability, -Exposure) %>%
mutate(sev_ord = fct_inorder(Severity)) -> preds
```

Some of the gathered predictions

```
preds %>% slice(1:15)
```

```
Exposure Severity probability
##
                                       sev\_ord
           5.8
                    None
                          0.96769203
                                          None
## 1
## 2
          15.0
                    None
                          0.92534455
                                          None
## 3
          21.5
                    None
                          0.86920028
                                          None
##
          27.5
                    None
                          0.78892903
                                          None
                    None
## 5
          33.5
                          0.67766411
                                          None
                          0.54181046
## 6
          39.5
                   None
                                          None
##
          46.0
                    None
                          0.38799618
                                          None
                          0.27225426
##
  8
          51.5
                    None
                                          None
                          0.01908912 Moderate
##
  9
           5.8 Moderate
## 10
          15.0 Moderate
                          0.04329931 Moderate
                          0.07385858 Moderate
##
  11
          21.5 Moderate
          27.5 Moderate
                          0.11413004 Moderate
##
  12
## 13
          33.5 Moderate
                          0.16207145 Moderate
          39.5 Moderate
                          0.20484198 Moderate
## 14
## 15
          46.0 Moderate
                          0.22441555 Moderate
```

Plotting predicted and observed proportions

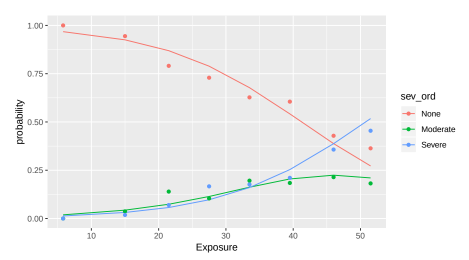
- Plot:
 - predicted probabilities, lines (shown) joining points (not shown)
 - data, just the points.
- Unfamiliar process: data from two different data frames:

```
g <- ggplot(preds, aes(
    x = Exposure, y = probability,
    colour = sev_ord
)) + geom_line() +
    geom_point(data = miners, aes(y = proportion))</pre>
```

Idea: final geom_point uses data in miners rather than preds,
 y-variable for plot is proportion from that data frame, but
 x-coordinate is Exposure, as it was before, and colour is Severity as before. The final geom_point "inherits" from the first aes as needed.

The plot: data match model

g



Unordered responses

- With unordered (nominal) responses, can use generalized logit.
- Example: 735 people, record age and sex (male 0, female 1), which of 3 brands of some product preferred.
- Data in mlogit.csv separated by commas (so read_csv will work):

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/mlogit.csv"
brandpref <- read_csv(my_url)

## Parsed with column specification:
## cols(
## brand = col_double(),
## sex = col_double(),
## age = col_double()</pre>
```

)

The data

brandpref

```
##
       brand
                sex
                        age
##
       <dbl> <dbl> <dbl>
##
                         24
                         26
##
    3
                         26
##
                         27
##
##
    5
                         27
           3
                         27
##
    6
                         27
##
                         27
##
    8
                         27
##
##
   10
                         27
     ... with 725 more rows
```

A tibble: 735 x 3

Lecture notes

Bashing into shape, and fitting model

• sex and brand not meaningful as numbers, so turn into factors:

```
brandpref <- brandpref %>%
  mutate(sex = factor(sex)) %>%
  mutate(brand = factor(brand))
```

• We use multinom from package nnet. Works like polr.

```
brands.1 <- multinom(brand ~ age + sex, data = brandpref)</pre>
```

```
## # weights: 12 (6 variable)
## initial value 807.480032
## iter 10 value 702.976983
## final value 702.970704
## converged
```

Can we drop anything?

• Unfortunately drop1 seems not to work:

```
drop1(brands.1, test = "Chisq", trace = 0)
```

```
## trying - age
```

Error in if (trace) $\{: argument is not interpretable as log$

 so fall back on fitting model without what you want to test, and comparing using anova.

Do age/sex help predict brand? 1/2

```
Fit models without each of age and sex:
```

```
brands.2 <- multinom(brand ~ age, data = brandpref)

## # weights: 9 (4 variable)

## initial value 807.480032

## iter 10 value 706.796323

## iter 10 value 706.796322

## final value 706.796322

## converged

brands.3 <- multinom(brand ~ sex, data = brandpref)</pre>
```

weights: 9 (4 variable)
initial value 807.480032
final value 791.861266
converged

Do age/sex help predict brand? 2/2

```
## Likelihood ratio tests of Multinomial Models
##
## Response: brand
## Model Resid. df Resid. Dev Test Df LR stat. Pr(Chi)
## 1 age 1466 1413.593
## 2 age + sex 1464 1405.941 1 vs 2 2 7.651236 0.02180495
anova(brands.3, brands.1)
## Likelihood ratio tests of Multinomial Models
##
## Response: brand
```

Model Resid. df Resid. Dev Test Df LR stat. Pr(Chi)

1583.723

2 age + sex 1464 1405.941 1 vs 2 2 177.7811

1466

sex

##

1

0

Do age/sex help predict brand? 3/3

- age definitely significant (second anova)
- sex seems significant also (first anova)
- Keep both.

Another way to build model

Start from model with everything and run step:

```
step(brands.1, trace = 0)
## trying - age
## trying - sex
## Call:
## multinom(formula = brand ~ age + sex, data = brandpref)
##
## Coefficients:
##
     (Intercept)
                       age
                                sex1
## 2 -11.77469 0.3682075 0.5238197
## 3 -22.72141 0.6859087 0.4659488
##
  Residual Deviance: 1405.941
## AIC: 1417.941
```

• Final model contains both age and sex so neither could be removed.

Predictions: all possible combinations

Create data frame with various age and sex:

```
ages <- c(24, 28, 32, 35, 38)
sexes <- factor(0:1)
new <- crossing(age = ages, sex = sexes)
new

## # A tibble: 10 x 2
## age sex
## <dbl> <fct>
## 1 24 0
## 2 24 1
```

10

Making predictions

```
p <- predict(brands.1, new, type = "probs")
probs <- cbind(new, p)

or
p %>% as_tibble() %>%
```

bind_cols(new) -> probs

The predictions

probs

```
A tibble: 10 x 5
         `1`
##
                              age sex
##
       <dbl>
              <dbl>
                    <dbl> <dbl> <fct>
##
    1 0.948
             0.0502 0.00181
                               24 0
##
    2 0.915
             0.0819 0.00279
                               24 1
    3 0.793 0.183
                    0.0236
##
                               28 0
    4 0.696 0.271 0.0329
                               28 1
##
    5 0.405 0.408
                    0.187
                               32 0
##
    6 0.291 0.495
                    0.214
                               32 1
##
##
   7 0.131 0.397
                    0.472
                               35 0
##
    8 0.0840 0.432
                    0.484
                               35 1
##
    9 0.0260 0.239
                    0.735
                               38 0
   10 0.0162 0.252
                    0.732
                               38 1
```

- Young males (sex=0) prefer brand 1, but older males prefer brand 3.
- Females similar, but like brand 1 less and brand 2 more.

Making a plot

- Plot fitted probability against age, distinguishing brand by colour and gender by plotting symbol.
- Also join points by lines, and distinguish lines by gender.
- I thought about facetting, but this seems to come out clearer.
- First need tidy data frame, by familiar process:

```
probs %>%
  gather(brand, probability, -(age:sex)) -> probs.long
```

The tidy data (random sample of rows)

```
probs.long %>% sample_n(10)
   # A tibble: 10 \times 4
##
         age sex brand probability
      <dbl> <fct> <chr>
##
                                  <dbl>
                               0.0162
##
          38 1
    1
##
          35 1
                               0.484
##
    3
          24 0
                               0.0502
##
    4
          24 1
                    3
                               0.00279
##
    5
          32 1
                    3
                               0.214
    6
          35 1
                               0.432
##
##
          28 1
                               0.271
##
          38 0
                               0.239
    8
##
    9
          38 0
                               0.0260
```

38 1

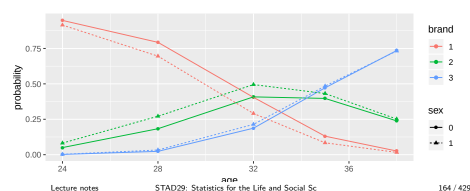
10

0.252

The plot

Lecture notes

```
ggplot(probs.long, aes(
  x = age, y = probability,
  colour = brand, shape = sex
  geom_point() + geom_line(aes(linetype = sex))
```



Digesting the plot

- Brand vs. age: younger people (of both genders) prefer brand 1, but older people (of both genders) prefer brand 3. (Explains significant age effect.)
- Brand vs. sex: females (dashed) like brand 1 less than males (solid), like brand 2 more (for all ages).
- Not much brand difference between genders (solid and dashed lines of same colours close), but enough to be significant.
- Model didn't include interaction, so modelled effect of gender on brand same for each age, modelled effect of age same for each gender.

Alternative data format

Summarize all people of same brand preference, same sex, same age on one line of data file with frequency on end:

```
1 0 24 1
```

1 0 26 2

1 0 27 4

1 0 28 4

1 0 29 7

1 0 30 3

. .

Whole data set in 65 lines not 735! But how?

Getting alternative data format

```
brandpref %>%
  group_by(age, sex, brand) %>%
  summarize(Freq = n()) %>%
  ungroup() -> b
## [conflicted] `summarize` found in 2 packages.
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict prefer()`
## * conflict prefer("summarize", "plyr")
## * conflict prefer("summarize", "dplyr")
b %>% slice(1:6)
```

Error in eval(lhs, parent, parent): object 'b' not found

Fitting models, almost the same

- Just have to remember weights to incorporate frequencies.
- Otherwise multinom assumes you have just 1 obs on each line!
- Again turn (numerical) sex and brand into factors:

Error in is.data.frame(data): object 'bf' not found

```
b %>%
  mutate(sex = factor(sex)) %>%
  mutate(brand = factor(brand)) -> bf

## Error in eval(lhs, parent, parent): object 'b' not found
b.1 <- multinom(brand ~ age + sex, data = bf, weights = Freq)

## Error in is.data.frame(data): object 'bf' not found
b.2 <- multinom(brand ~ age, data = bf, weights = Freq)</pre>
```

P-value for sex identical

```
anova(b.2, b.1)
```

Error in anova(b.2, b.1): object 'b.2' not found

Same P-value as before, so we haven't changed anything important.

Including data on plot

 Everyone's age given as whole number, so maybe not too many different ages with sensible amount of data at each:

```
b %>%
group_by(age) %>%
summarize(total = sum(Freq))
```

```
## Error in eval(lhs, parent, parent): object 'b' not found
```

Comments and next

- Not great (especially at low end), but live with it.
- Need proportions of frequencies in each brand for each age-gender combination. Mimic what we did for miners:

```
b %>%
group_by(age, sex) %>%
mutate(proportion = Freq / sum(Freq)) -> brands
```

```
## Error in eval(lhs, parent, parent): object 'b' not found
```

Checking proportions for age 32

```
brands %>% filter(age == 32)
```

- ## Error in eval(lhs, parent, parent): object 'brands' not found
 - First three proportions (males) add up to 1.
 - Last three proportions (females) add up to 1.
 - So looks like proportions of right thing.

Attempting plot

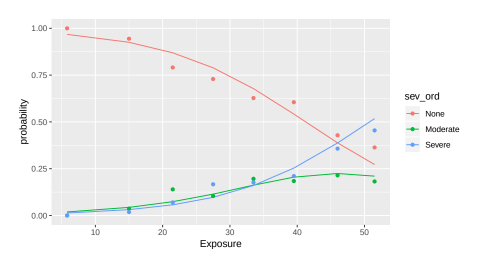
- Take code from previous plot and:
- remove geom_point for fitted values
- add geom_point with correct data= and aes to plot data.

```
g <- ggplot(probs.long, aes(
    x = age, y = probability,
    colour = brand, shape = sex
)) +
    geom_line(aes(linetype = sex)) +
    geom_point(data = brands, aes(y = proportion))</pre>
```

- ## Error in fortify(data): object 'brands' not found
 - Data seem to correspond more or less to fitted curves:

The plot

g



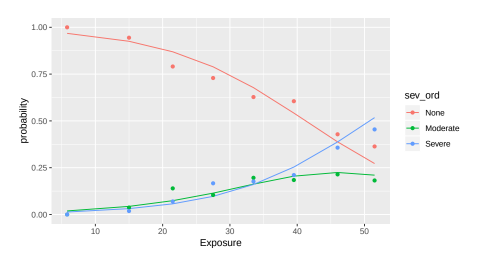
But...

- Some of the plotted points based on a lot of people, and some only a few.
- Idea: make the *size* of plotted point bigger if point based on a lot of people (in Freq).
- Hope that larger points then closer to predictions.
- Code:

```
g <- ggplot(probs.long, aes(
    x = age, y = probability,
    colour = brand, shape = sex
)) +
    geom_line(aes(linetype = sex)) +
    geom_point(
        data = brands,
        aes(y = proportion, size = Freq)
)</pre>
```

The plot

g



Trying interaction between age and gender

```
b.4 <- update(b.1, . ~ . + age:sex)
## Error in update(b.1, . ~ . + age:sex): object 'b.1' not found
anova(b.1, b.4)</pre>
```

- ## Error in anova(b.1, b.4): object 'b.1' not found
 - No evidence that effect of age on brand preference differs for the two genders.

Section 4

Survival analysis

Survival analysis

- So far, have seen:
 - response variable counted or measured (regression)
 - response variable categorized (logistic regression)

and have predicted response from explanatory variables.

- But what if response is time until event (eg. time of survival after surgery)?
- Additional complication: event might not have happened at end of study (eg. patient still alive). But knowing that patient has "not died yet" presumably informative. Such data called *censored*.
- Enter survival analysis, in particular the "Cox proportional hazards model".
- Explanatory variables in this context often called covariates.

Example: still dancing?

- 12 women who have just started taking dancing lessons are followed for up to a year, to see whether they are still taking dancing lessons, or have quit. The "event" here is "quit".
- This might depend on:
 - a treatment (visit to a dance competition)
 - woman's age (at start of study).

Data

| Months | Quit | Treatment | Age |
|--------|------|-----------|-----|
| 1 | 1 | 0 | 16 |
| 2 | 1 | 0 | 24 |
| 2 | 1 | 0 | 18 |
| 3 | 0 | 0 | 27 |
| 4 | 1 | 0 | 25 |
| 7 | 1 | 1 | 26 |
| 8 | 1 | 1 | 36 |
| 10 | 1 | 1 | 38 |
| 10 | 0 | 1 | 45 |
| 12 | 1 | 1 | 47 |

About the data

- months and quit are kind of combined response:
 - Months is number of months a woman was actually observed dancing
 - quit is 1 if woman quit, 0 if still dancing at end of study.
- Treatment is 1 if woman went to dance competition, 0 otherwise.
- Fit model and see whether Age or Treatment have effect on survival.
- Want to do predictions for probabilities of still dancing as they depend on whatever is significant, and draw plot.

Packages (for this section)

- Install packages survival and survminer if not done.
- Load survival, survminer, broom and tidyverse:

```
library(tidyverse)
library(survival)
library(survminer)
library(broom)
```

Read data

)

Column-aligned:

```
url <- "http://www.utsc.utoronto.ca/~butler/d29/dancing.txt"
dance <- read_table(url)

## Parsed with column specification:
## cols(
## Months = col_double(),
## Quit = col_double(),
## Treatment = col double(),</pre>
```

Age = col_double()

The data

dance

```
## # A tibble: 12 x 4
##
      Months Quit Treatment
                                   Age
##
        <dbl> <dbl>
                          <dbl> <dbl>
##
                                    16
##
    2
                                    24
    3
            2
##
                                    18
    4
            3
                                    27
##
##
    5
            4
                                    25
            5
##
                                    21
##
    7
           11
                                    55
##
    8
                                    26
            8
                                    36
##
## 10
           10
                                    38
           10
                                    45
## 11
## 12
           12
                                    47
```

Examine response and fit model

• Response variable (has to be outside data frame):

```
mth <- with(dance, Surv(Months, Quit))
mth</pre>
```

```
## [1] 1 2 2 3+ 4 5 11 7 8 10 10+ 12
```

• Then fit model, predicting mth from explanatories:

```
dance.1 <- coxph(mth ~ Treatment + Age, data = dance)</pre>
```

Output looks a lot like regression

```
summary(dance.1)
```

```
## Call:
## coxph(formula = mth ~ Treatment + Age, data = dance)
##
##
   n= 12, number of events= 10
##
##
              coef exp(coef) se(coef) z Pr(>|z|)
## Treatment -4.44915 0.01169 2.60929 -1.705 0.0882 .
     -0.36619 0.69337 0.15381 -2.381 0.0173 *
## Age
## ---
## Signif. codes:
## 0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' ' 1
##
           exp(coef) exp(-coef) lower .95 upper .95
##
## Treatment 0.01169 85.554 7.026e-05 1.9444
## Age 0.69337 1.442 5.129e-01 0.9373
##
## Concordance= 0.964 (se = 0.039)
## Likelihood ratio test= 21.68 on 2 df. p=2e-05
## Wald test = 5.67 on 2 df. p=0.06
## Score (logrank) test = 14.75 on 2 df, p=6e-04
```

Conclusions

- Use $\alpha = 0.10$ here since not much data.
- Three tests at bottom like global F-test. Consensus that something predicts survival time (whether or not dancer quit and how long it took).
- Age (definitely), Treatment (marginally) both predict survival time.

Model checking

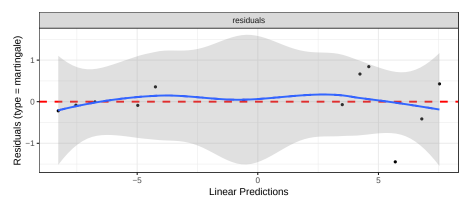
- With regression, usually plot residuals against fitted values.
- Not quite same here (nonlinear model), but "martingale residuals" should have no pattern vs. "linear predictor".
- ggcoxdiagnostics from package survminer makes plot, to which we add smooth. If smooth trend more or less straight across, model OK.
- Martingale residuals can go very negative, so won't always look normal.

Martingale residual plot for dance data

This looks good (with only 12 points):

```
ggcoxdiagnostics(dance.1) + geom_smooth(se = F)
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x



Predicted survival probs

- The function we use is called survfit, though actually works rather like predict.
- First create a data frame of values to predict from. We'll do all combos of ages 20 and 40, treatment and not, using crossing to get all the combos:

```
treatments <- c(0, 1)
ages <- c(20, 40)
dance.new <- crossing(Treatment = treatments, Age = ages)
dance.new</pre>
```

```
## # A tibble: 4 x 2
## Treatment Age
## (dbl> <dbl>
## 1 0 20
## 2 0 40
## 3 1 20
## 4 1 40
```

The predictions

One prediction *for each time* for each combo of age and treatment in dance.new:

```
s <- survfit(dance.1, newdata = dance.new, data = dance)
summary(s)
## Call: survfit(formula = dance.1, newdata = dance.new, data = dance)
##
##
   time n.risk n.event survival1 survival2 survival3 survival4
##
      1
            12
                        8.76e-01 1.00e+00
                                           9.98e-01
                                                        1.000
                                                        1.000
##
            11
                     2 3.99e-01 9.99e-01
                                           9.89e-01
      4
             8
                       1.24e-01 9.99e-01
                                           9.76e-01
                                                       1.000
##
      5
                        2.93e-02 9.98e-01
                                           9.60e-01
                                                        1.000
##
             6
                     1 2.96e-323 6.13e-01
                                           1.70e-04
                                                        0.994
##
                                           1.35e-98
##
      8
             5
                        0.00e+00 2.99e-06
                                                        0.862
##
     10
             4
                        0.00e+00 3.61e-20
                                           0.00e+00
                                                        0.593
     11
                        0.00e+00 0.00e+00
                                           0.00e+00
                                                        0.000
##
##
     12
                        0.00e+00
                                 0.00e+00
                                           0.00e+00
                                                        0.000
```

Conclusions from predicted probs

- Older women more likely to be still dancing than younger women (compare "profiles" for same treatment group).
- Effect of treatment seems to be to increase prob of still dancing (compare "profiles" for same age for treatment group vs. not)
- Would be nice to see this on a graph. This is ggsurvplot from package survminer:

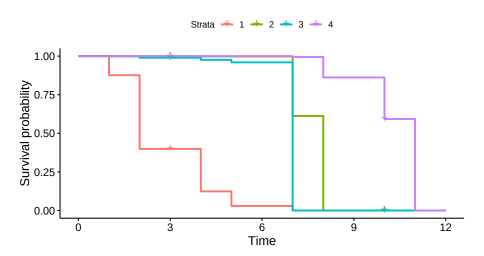
```
g <- ggsurvplot(s, conf.int = F)
```

uses "strata" thus (dance.new):

```
## # A tibble: 4 x 2
## Treatment Age
## <dbl> <dbl> ## 1 0 20
## 2 0 40
## 3 1 20
## 4 1 40
Lecture notes
```

Plotting survival probabilities

g



Discussion

- Survivor curve farther to the right is better (better chance of surviving longer).
- Best is age 40 with treatment, worst age 20 without.
- Appears to be:
 - age effect (40 better than 20)
 - treatment effect (treatment better than not)
 - In analysis, treatment effect only marginally significant.

A more realistic example: lung cancer

- When you load in an R package, get data sets to illustrate functions in the package.
- One such is lung. Data set measuring survival in patients with advanced lung cancer.
- Along with survival time, number of "performance scores" included, measuring how well patients can perform daily activities.
- Sometimes high good, but sometimes bad!
- Variables below, from the data set help file (?lung).

The variables

Format

inst: Institution code

time: Survival time in days

status: censoring status 1=censored, 2=dead

age: Age in years

sex: Male=1 Female=2

ph.ecog: ECOG performance score (0=good 5=dead)

ph.karno: Karnofsky performance score (bad=0-good=100) rated by physician

pat.karno: Karnofsky performance score as rated by patient

meal.cal: Calories consumed at meals

wt.loss: Weight loss in last six months

Uh oh, missing values

```
lung %>% slice(1:16)
```

```
##
       inst time status age sex ph.ecog ph.karno pat.karno meal.cal wt.loss
## 1
          3
              306
                            74
                                                      90
                                                                100
                                                                          1175
                                                                                      NA
## 2
              455
                            68
                                                      90
                                                                 90
                                                                          1225
                                                                                      15
                                  1
                                            0
## 3
          3 1010
                            56
                                                      90
                                                                            NA
                                                                                      15
                                  1
                                                                  90
              210
                            57
                                                                          1150
## 4
          5
                                  1
                                                      90
                                                                 60
                                                                                      11
## 5
              883
                            60
                                            0
                                                    100
                                                                            NA
                                  1
                                                                 90
                                                                                       0
## 6
         12 1022
                            74
                                  1
                                                      50
                                                                 80
                                                                           513
                                                                                       0
                                            1
## 7
          7
              310
                            68
                                  2
                                            2
                                                      70
                                                                 60
                                                                           384
                                                                                      10
## 8
              361
                            71
                                  2
                                                                           538
         11
                                                      60
                                                                 80
                                                                                       1
##
   9
          1
              218
                            53
                                  1
                                            1
                                                      70
                                                                 80
                                                                           825
                                                                                      16
## 10
              166
                            61
                                                      70
                                                                 70
                                                                           271
                                                                                      34
                                  1
## 11
          6
              170
                            57
                                  1
                                                      80
                                                                 80
                                                                          1025
                                                                                      27
## 12
         16
              654
                         2
                            68
                                  2
                                            2
                                                      70
                                                                  70
                                                                            NΑ
                                                                                      23
## 13
         11
              728
                            68
                                  2
                                                      90
                                                                 90
                                                                            NA
                                                                                       5
## 14
         21
               71
                            60
                                  1
                                           NA
                                                      60
                                                                 70
                                                                          1225
                                                                                      32
## 15
         12
              567
                            57
                                            1
                                                      80
                                                                  70
                                                                          2600
                                                                                      60
##
  16
          1
              144
                            67
                                  1
                                                      80
                                                                 90
                                                                            NA
                                                                                      15
```

A closer look

summary(lung)

```
##
        inst
                        time
                                         status
                                                          age
                                                                          sex
          : 1.00
                              5.0
                                            :1.000
                                                            :39.00
                                                                            :1.000
   Min.
                   Min.
                                    Min.
                                                     Min.
                                                                     Min.
   1st Qu.: 3.00
                   1st Qu.: 166.8
                                    1st Qu.:1.000
                                                     1st Qu.:56.00
                                                                     1st Qu.:1.000
   Median :11.00
                   Median : 255.5
                                    Median :2.000
                                                     Median :63.00
                                                                     Median :1.000
                          : 305.2
   Mean
          :11.09
                   Mean
                                    Mean
                                            :1.724
                                                     Mean
                                                            :62.45
                                                                     Mean
                                                                            :1.395
   3rd Qu.:16.00
                   3rd Qu.: 396.5
                                    3rd Qu.:2.000
                                                     3rd Qu.:69.00
                                                                     3rd Qu.:2.000
   Max
          :33.00
                   Max.
                          .1022.0
                                     Max
                                            .2 000
                                                     Max
                                                            .82.00
                                                                     Max.
                                                                            :2.000
   NA's
         : 1
                                                                           wt.loss
      ph.ecog
                       ph.karno
                                       pat.karno
                                                          meal.cal
   Min.
          :0.0000
                    Min.
                           : 50.00
                                     Min.
                                            : 30.00
                                                      Min.
                                                            . 96.0
                                                                        Min.
                                                                               :-24.000
   1st Qu.:0.0000
                    1st Qu.: 75.00
                                     1st Qu.: 70.00
                                                      1st Qu.: 635.0
                                                                        1st Qu.: 0.000
   Median :1.0000
                    Median: 80.00
                                     Median : 80.00
                                                      Median : 975.0
                                                                        Median: 7.000
   Mean
          :0.9515
                           : 81.94
                                            : 79.96
                                                             : 928.8
                                                                             : 9.832
                    Mean
                                     Mean
                                                       Mean
                                                                        Mean
   3rd Qu.:1.0000
                    3rd Qu.: 90.00
                                     3rd Qu.: 90.00
                                                       3rd Qu.:1150.0
                                                                        3rd Qu.: 15.750
          :3.0000
                           :100.00
                                            :100.00
                                                              :2600.0
                                                                               : 68.000
   Max.
                    Max.
                                     Max.
                                                       Max.
                                                                        Max.
   NA's
          :1
                    NA's
                           :1
                                     NA's
                                            :3
                                                       NA's
                                                              :47
                                                                        NA's
                                                                               :14
```

Remove obs with any missing values

15

-5

```
lung %>% drop_na() -> lung.complete
lung.complete %>%
   select(meal.cal:wt.loss) %>%
   slice(1:10)
## meal.cal wt.loss
```

```
## 2
           1150
                      11
## 3
            513
                      10
## 4
            384
            538
## 5
            825
                      16
## 6
            271
## 7
                      34
## 8
          1025
                      27
           2600
                      60
## 9
```

1150

1225

1

10

Missing values seem to be gone.

Check!

summary(lung.complete)

```
inst
                          time
                                           status
                                                            age
                                                                             sex
    Min.
           : 1.00
                                5.0
                                              :1.000
                                                               :39.00
                                                                               :1.000
                     Min.
                                      Min.
                                                       Min.
                                                                        Min.
    1st Qu.: 3.00
                     1st Qu.: 174.5
                                      1st Qu.:1.000
                                                       1st Qu.:57.00
                                                                        1st Qu.:1.000
   Median :11.00
                    Median : 268.0
                                      Median :2.000
                                                       Median :64.00
                                                                        Median :1.000
    Mean
           :10.71
                     Mean
                            : 309.9
                                      Mean
                                              :1.719
                                                       Mean
                                                               :62.57
                                                                        Mean
                                                                               :1.383
    3rd Qu.:15.00
                    3rd Qu.: 419.5
                                      3rd Qu.:2.000
                                                       3rd Qu.:70.00
                                                                        3rd Qu.:2.000
   Max.
           :32.00
                            :1022.0
                                              :2.000
                    Max.
                                      Max.
                                                       Max.
                                                               :82.00
                                                                        Max.
                                                                               :2.000
##
       ph.ecog
                        ph.karno
                                         pat.karno
                                                            meal.cal
                                                                              wt.loss
   Min.
           :0.0000
                      Min.
                             : 50.00
                                       Min.
                                              : 30.00
                                                         Min.
                                                                           Min.
                                                               : 96.0
                                                                                  :-24.000
   1st Qu.:0.0000
                      1st Qu.: 70.00
                                       1st Qu.: 70.00
                                                         1st Qu.: 619.0
                                                                           1st Qu.:
                                                                                     0.000
   Median :1.0000
                      Median : 80.00
                                       Median: 80.00
                                                         Median: 975.0
                                                                           Median :
                                                                                     7.000
   Mean
           :0.9581
                      Mean
                             : 82.04
                                       Mean
                                              : 79.58
                                                         Mean
                                                                : 929.1
                                                                           Mean
                                                                                 : 9.719
   3rd Qu.:1.0000
                     3rd Qu.: 90.00
                                       3rd Qu.: 90.00
                                                         3rd Qu.:1162.5
                                                                           3rd Qu.: 15.000
   Max.
           :3.0000
                      Max.
                             :100.00
                                       Max.
                                              :100.00
                                                         Max.
                                                                :2600.0
                                                                           Max.
                                                                                  : 68.000
```

No missing values left.

Model 1: use everything except inst

```
names(lung.complete)

## [1] "inst" "time" "status" "age" "sex" "ph.ecc
## [8] "pat.karno" "meal.cal" "wt.loss"

• Event was death, goes with status of 2:

resp <- with(lung.complete, Surv(time, status == 2))
lung.1 <- coxph(resp ~ . - inst - time - status,
    data = lung.complete
)</pre>
```

"Dot" means "all the other variables".

summary of model 1: too tiny to see!

```
## Call:
## coxph(formula = resp ~ . - inst - time - status, data = lung.complete)
##
##
    n= 167, number of events= 120
##
##
                 coef exp(coef) se(coef)
                                              z Pr(>|z|)
## age
           1.080e-02 1.011e+00 1.160e-02 0.931 0.35168
          -5.536e-01 5.749e-01 2.016e-01 -2.746 0.00603 **
## sex
## ph.ecog 7.395e-01 2.095e+00 2.250e-01 3.287 0.00101 **
## ph.karno 2.244e-02 1.023e+00 1.123e-02 1.998 0.04575 *
## pat.karno -1.207e-02 9.880e-01 8.116e-03 -1.488 0.13685
## meal.cal 2.835e-05 1.000e+00 2.594e-04 0.109 0.91298
## wt.loss -1.420e-02 9.859e-01 7.766e-03 -1.828 0.06748 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
           exp(coef) exp(-coef) lower .95 upper .95
## age
             1.0109
                        0.9893
                                 0.9881 1.0341
## sex
            0.5749
                        1.7395 0.3872
                                          0.8534
## ph.ecog
            2.0950
                        0.4773 1.3479
                                        3.2560
## ph.karno 1.0227
                        0.9778 1.0004 1.0455
## pat.karno 0.9880 1.0121 0.9724
                                        1.0038
                     1.0000 0.9995
## meal.cal
          1.0000
                                         1.0005
                                0.9710
## wt.loss
              0.9859
                        1.0143
                                          1.0010
##
## Concordance= 0.653 (se = 0.029 )
## Likelihood ratio test= 28.16 on 7 df. p=2e-04
                     = 27.5 on 7 df, p=3e-04
## Wald test
## Score (logrank) test = 28.31 on 7 df, p=2e-04
```

summary(lung.1)

Overall significance

The three tests of overall significance:

```
glance(lung.1) %>% select(starts_with("p.value"))
## # A tibble: 1 x 3
```

```
## p.value.log p.value.sc p.value.wald
## <dbl> <dbl> <dbl> <dbl> ## 1 0.000205 0.000193 0.000271
```

All strongly significant. Something predicts survival.

Coefficients for model 1

```
tidy(lung.1) %>% select(term, p.value) %>% arrange(p.value)
```

A tibble: 7×2

- sex and ph.ecog definitely significant here
- age, pat.karno and meal.cal definitely not
- Take out definitely non-sig variables, and try again.

Model 2

```
lung.2 <- update(lung.1, . ~ . - age - pat.karno - meal.cal)
tidy(lung.2) %>% select(term, p.value)

## # A tibble: 4 x 2

## term p.value
## <chr> <dbl>
```

1 sex 0.00409 ## 2 ph.ecog 0.000112 ## 3 ph.karno 0.101 ## 4 wt.loss 0.108

Compare with first model:

anova(lung.2, lung.1)

```
## Analysis of Deviance Table
## Cox model: response is resp
## Model 1: ~ sex + ph.ecog + ph.karno + wt.loss
## Model 2: ~ (inst + time + status + age + sex + ph.ecog + p
```

No harm in taking out those variables.

Model 3

```
Take out ph.karno and wt.loss as well.
lung.3 <- update(lung.2, . ~ . - ph.karno - wt.loss)</pre>
```

tidy(lung.3) %>% select(term, estimate, p.value)

Lecture notes

2 ph.ecog 0.483 0.000266

Check whether that was OK

anova(lung.3, lung.2)

```
Analysis of Deviance Table
   Cox model: response is resp
##
##
   Model 1: ~ sex + ph.ecog
##
  Model 2: ~ sex + ph.ecog + ph.karno + wt.loss
     loglik Chisq Df P(>|Chi|)
##
## 1 -498.38
## 2 -495.67 5.4135 2 0.06675 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 '
```

Just OK.

Commentary

- OK (just) to take out those two covariates.
- Both remaining variables strongly significant.
- Nature of effect on survival time? Consider later.
- Picture?

Plotting survival probabilities

• Create new data frame of values to predict for, then predict:

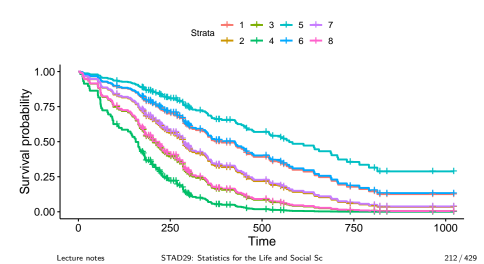
```
sexes <- c(1, 2)
ph.ecogs <- 0:3
lung.new <- crossing(sex = sexes, ph.ecog = ph.ecogs)
lung.new</pre>
```

```
## # A tibble: 8 x 2
## sex ph.ecog
## 1 1 0
## 2 1 1
## 3 1 2
## 4 1 3
## 5 2 0
## 6 2 1
## 7 2 2
## 8 2 3
```

s <- survfit(lung.3, data = lung.complete, newdata = lung.new)

The plot

ggsurvplot(s, conf.int = F)



Discussion of survival curves

- Best survival is teal-blue curve, stratum 5, females with (ph.ecog) score 0.
- Next best: blue, stratum 6, females with score 1, and red, stratum 1, males score 0.
- Worst: green, stratum 4, males score 3.
- For any given ph.ecog score, females have better predicted survival than males.
- For both genders, a lower score associated with better survival.

The coefficients in model 3

A tibble: 2×3

```
tidy(lung.3) %>% select(term, estimate, p.value)
```

```
## term estimate p.value

## <chr> <dbl> <dbl> <dbl>

## 1 sex -0.510 0.00958

## 2 ph.ecog 0.483 0.000266
```

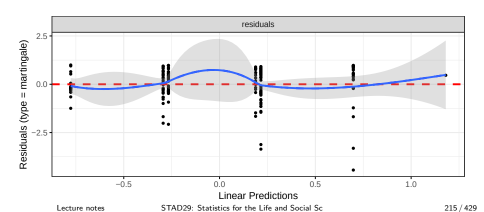
- sex coeff negative, so being higher sex value (female) goes with less hazard of dying.
- ph.ecog coeff positive, so higher ph.ecog score goes with more hazard of dying
- Two coeffs about same size, so being male rather than female corresponds to 1-point increase in ph.ecog score. Note how survival curves come in 3 pairs plus 2 odd.

Martingale residuals for this model

No problems here:

```
ggcoxdiagnostics(lung.3) + geom_smooth(se = F)
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x



When the Cox model fails

 Invent some data where survival is best at middling age, and worse at high and low age:

```
age <- seq(20, 60, 5)
survtime <- c(10, 12, 11, 21, 15, 20, 8, 9, 11)
stat <- c(1, 1, 1, 1, 0, 1, 1, 1, 1)
d <- tibble(age, survtime, stat)
y <- with(d, Surv(survtime, stat))</pre>
```

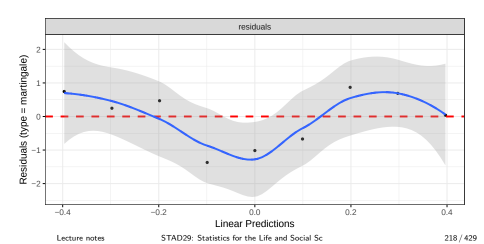
• Small survival time 15 in middle was actually censored, so would have been longer if observed.

Fit Cox model

```
y.1 \leftarrow coxph(y \sim age, data = d)
summary(y.1)
## Call:
## coxph(formula = y ~ age, data = d)
##
## n= 9, number of events= 8
##
         coef exp(coef) se(coef) z Pr(>|z|)
##
## age 0.01984   1.02003   0.03446   0.576
                                         0.565
##
      exp(coef) exp(-coef) lower .95 upper .95
##
## age
           1.02 0.9804 0.9534 1.091
##
## Concordance= 0.545 (se = 0.105)
## Likelihood ratio test= 0.33 on 1 df, p=0.6
## Wald test
                = 0.33 on 1 df, p=0.6
## Score (logrank) test = 0.33 on 1 df,
                                         p = 0.6
```

Martingale residuals

Down-and-up indicates incorrect relationship between age and survival:



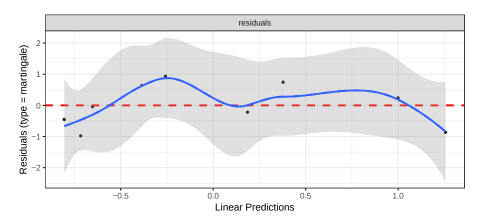
Attempt 2

Add squared term in age:

```
y.2 <- coxph(y ~ age + I(age^2), data = d)
tidy(y.2) %>% select(term, estimate, p.value)
```

Martingale residuals this time

Not great, but less problematic than before:



Section 5

Analysis of variance

Analysis of variance

- Analysis of variance used with:
 - counted/measured response
 - categorical explanatory variable(s)
 - that is, data divided into groups, and see if response significantly different among groups
 - $\, \bullet \,$ or, see whether knowing group membership helps to predict response.
- Typically two stages:
 - F-test to detect any differences among/due to groups
 - if *F*-test significant, do *multiple comparisons* to see which groups significantly different from which.
- Need special multiple comparisons method because just doing (say) two-sample t-tests on each pair of groups gives too big a chance of finding "significant" differences by accident.

STAD29: Statistics for the Life and Social Sc.

Packages

These:

```
library(tidyverse)
library(broom)
library(car) # for Levene's text
```

Example: Pain threshold and hair colour

- Do people with different hair colour have different abilities to deal with pain?
- Men and women of various ages divided into 4 groups by hair colour: light and dark blond, light and dark brown.
- Each subject given a pain sensitivity test resulting in pain threshold score: higher score is higher pain tolerance.
- 19 subjects altogether.

The data

In hairpain.txt:

| hair pain | darkblond 43 |
|---------------|---------------|
| lightblond 62 | lightbrown 42 |
| lightblond 60 | lightbrown 50 |
| lightblond 71 | lightbrown 41 |
| lightblond 55 | lightbrown 37 |
| lightblond 48 | darkbrown 32 |
| darkblond 63 | darkbrown 39 |
| darkblond 57 | darkbrown 51 |
| darkblond 52 | darkbrown 30 |
| darkblond 41 | darkbrown 35 |

Summarizing the groups

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/hairpain.txt"
hairpain <- read_delim(my_url, " ")
hairpain %>%
  group_by(hair) %>%
  summarize(
   n = n(),
   xbar = mean(pain),
   s = sd(pain)
)
```

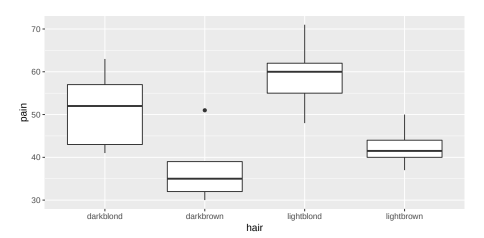
```
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict_prefer()`
## * conflict_prefer("summarize", "plyr")
## * conflict_prefer("summarize", "dplyr")
```

[conflicted] `summarize` found in 2 packages.

Brown-haired people seem to have lower pain tolerance.

Boxplot

ggplot(hairpain, aes(x = hair, y = pain)) + geom_boxplot()



Assumptions

- Data should be:
 - normally distributed within each group
 - same spread for each group
- darkbrown group has upper outlier (suggests not normal)
- darkblond group has smaller IQR than other groups.
- But, groups small.
- Shrug shoulders and continue for moment.

Testing equality of SDs

• via Levene's test in package car:

```
leveneTest(pain ~ hair, data = hairpain)
## Warning in leveneTest.default(y = y, group = group, ...): group
```

```
## Levene's Test for Homogeneity of Variance (center = median)
## Df F value Pr(>F)
## group 3 0.3927 0.76
## 15
```

- No evidence (at all) of difference among group SDs.
- Possibly because groups small.

Analysis of variance

```
hairpain.1 <- aov(pain ~ hair, data = hairpain)
summary(hairpain.1)</pre>
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## hair 3 1361 453.6 6.791 0.00411 **
## Residuals 15 1002 66.8
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

- P-value small: the mean pain tolerances for the four groups are *not* all the same.
- Which groups differ from which, and how?

Multiple comparisons

- Which groups differ from which? Multiple comparisons method. Lots.
- ullet Problem: by comparing all the groups with each other, doing many tests, have large chance to (possibly incorrectly) reject H_0 : groups have equal means.
- 4 groups: 6 comparisons (1 vs 2, 1 vs 3, ..., 3 vs 4). 5 groups: 10 comparisons. Thus 6 (or 10) chances to make mistake.
- Get "familywise error rate" of 0.05 (whatever), no matter how many comparisons you're doing.
- My favourite: Tukey, or "honestly significant differences": how far apart might largest, smallest group means be (if actually no differences). Group means more different: significantly different.

Tukey

• TukeyHSD:

```
TukeyHSD(hairpain.1)
```

```
##
    Tukey multiple comparisons of means
      95% family-wise confidence level
##
##
## Fit: aov(formula = pain ~ hair, data = hairpain)
##
##
  $hair
                         diff
                                     lwr
##
                                               upr
                                                       p adi
  darkbrown-darkblond
                        -13.8 -28.696741 1.0967407 0.0740679
                       8.0 -6.896741 22.8967407 0.4355768
## lightblond-darkblond
## lightbrown-darkblond -8.7 -24.500380 7.1003795 0.4147283
## lightblond-darkbrown 21.8
                                6.903259 36.6967407 0.0037079
## lightbrown-darkbrown 5.1 -10.700380 20.9003795 0.7893211
## lightbrown-lightblond -16.7 -32.500380 -0.8996205 0.0366467
```

The old-fashioned way

- List group means in order
- Draw lines connecting groups that are *not* significantly different:

```
darkbrown lightbrown darkblond lightblond 37.4 42.5 51.2 59.2
```

- lightblond significantly higher than everything except darkblond (at $\alpha=0.05$).
- darkblond in middle ground: not significantly less than lightblond, not significantly greater than darkbrown and lightbrown.
- More data might resolve this.
- Looks as if blond-haired people do have higher pain tolerance, but not completely clear.

Some other multiple-comparison methods

- Work any time you do k tests at once (not just ANOVA).
 - **Bonferroni**: multiply all P-values by k.
 - **Holm**: multiply smallest P-value by k, next-smallest by k-1, etc.
 - False discovery rate: multiply smallest P-value by k/1, 2nd-smallest by k/2, ..., i-th smallest by k/i.
- Stop after non-rejection.

Example

- \bullet P-values 0.005, 0.015, 0.03, 0.06 (4 tests all done at once) Use $\alpha=0.05.$
- Bonferroni:
 - Multiply all P-values by 4 (4 tests).
 - Reject only 1st null.
- Holm:
 - Times smallest P-value by 4: 0.005 * 4 = 0.020 < 0.05, reject.
 - Times next smallest by 3: 0.015 * 3 = 0.045 < 0.05, reject.
 - \bullet Times next smallest by 2: 0.03*2=0.06>0.05, do not reject. Stop.

...Continued

- With P-values 0.005, 0.015, 0.03, 0.06:
- False discovery rate:
 - Times smallest P-value by 4: 0.005 * 4 = 0.02 < 0.05: reject.
 - Times second smallest by 4/2: 0.015 * 4/2 = 0.03 < 0.05, reject.
 - Times third smallest by 4/3: 0.03 * 4/3 = 0.04 < 0.05, reject.
 - Times fourth smallest by 4/4: 0.06*4/4=0.06>0.05, do not reject. Stop.

pairwise.t.test

```
attach(hairpain)
pairwise.t.test(pain, hair, p.adj = "none")
##
   Pairwise comparisons using t tests with pooled SD
##
## data: pain and hair
##
             darkblond darkbrown lightblond
## darkbrown 0.01748
## lightblond 0.14251 0.00075
## lightbrown 0.13337 0.36695
                                0.00817
##
## P value adjustment method: none
pairwise.t.test(pain, hair, p.adj = "holm")
   Pairwise comparisons using t tests with pooled SD
## data: pain and hair
```

P value adjustment method: holm

darkbrown 0.0699 -## lightblond 0.4001 0.0045 ## lightbrown 0.4001 0.4001

##

darkblond darkbrown lightblond

0.0408

pairwise.t.test part 2

darkblond darkbrown lightblond

0.0490

0.0045

1.0000

```
pairwise.t.test(pain, hair, p.adj = "fdr")
##
   Pairwise comparisons using t tests with pooled SD
##
## data: pain and hair
              darkblond darkbrown lightblond
## darkbrown 0.0350
## lightblond 0.1710
                     0.0045
## lightbrown 0.1710 0.3670
                                 0.0245
##
## P value adjustment method: fdr
pairwise.t.test(pain, hair, p.adi = "bon")
##
   Pairwise comparisons using t tests with pooled SD
##
## data: pain and hair
##
```

P value adjustment method: bonferroni

darkbrown 0.1049

##

lightblond 0.8550 ## lightbrown 0.8002

Comments

- P-values all adjusted upwards from "none".
- Required because 6 tests at once.
- Highest P-values for Bonferroni: most "conservative".
- Prefer Tukey or FDR or Holm.
- Tukey only applies to ANOVA, not to other cases of multiple testing.

Rats and vitamin B

- What is the effect of dietary vitamin B on the kidney?
- A number of rats were randomized to receive either a B-supplemented diet or a regular diet.
- Desired to control for initial size of rats, so classified into size classes lean and obese.
- After 20 weeks, rats' kidneys weighed.
- Variables:
 - Response: kidneyweight (grams).
 - Explanatory: diet, ratsize.
- Read in data:

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/vitaminb.tx
vitaminb <- read delim(my url, " ")</pre>
```

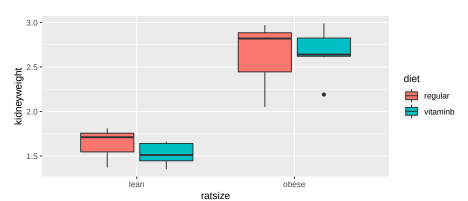
The data

vitaminb

```
## # A tibble: 28 x 3
##
      ratsize diet
                       kidneyweight
##
      <chr>
              <chr>
                              <dbl>
              regular
##
    1 lean
                                1.62
    2 lean
              regular
                                1.8
##
                                1.71
##
    3 lean
              regular
##
    4 lean
              regular
                                1.81
##
    5 lean
              regular
                                1.47
              regular
                                1.37
##
    6 lean
##
   7 lean
              regular
                                1.71
              vitaminb
##
    8 lean
                                1.51
##
    9 lean
             vitaminb
                                1.65
## 10 lean
              vitaminb
                                1.45
## # ... with 18 more rows
```

Grouped boxplot

```
ggplot(vitaminb, aes(
   x = ratsize, y = kidneyweight,
   fill = diet
)) + geom_boxplot()
```



What's going on?

Calculate group means:

function (object, ...)
UseMethod("summary")
<bytecode: 0x561d8d5b3e20>
<environment: namespace:base>

```
summary <- vitaminb %>%
  group_by(ratsize, diet) %>%
  summarize(mean = mean(kidneyweight))
## [conflicted] `summarize` found in 2 packages.
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict_prefer()`
## * conflict_prefer("summarize", "plyr")
## * conflict_prefer("summarize", "dplyr")
summary
```

vitaminb.1 <- aov(kidneyweight ~ ratsize * diet,</pre>

Significance/nonsignificance as we expected.Note no significant interaction (can be removed).

ANOVA with interaction

```
data = vitaminb
summary(vitaminb.1)
                                         Pr(>F)
##
               Df Sum Sq Mean Sq F value
                          8.068 141.179 1.53e-11 ***
## ratsize
                  8.068
                  0.012 0.012 0.218
## diet
                                          0.645
## ratsize:diet 1 0.036 0.036 0.638
                                          0.432
## Residuals 24 1.372 0.057
##
                 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '
## Signif. codes:
```

Interaction plot

 Plot mean of response variable against one of the explanatory, using other one as groups. Start from summary:

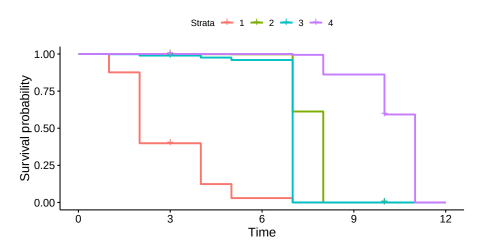
```
g <- ggplot(summary, aes(</pre>
  x = ratsize, y = mean,
  colour = diet, group = diet
)) +
  geom point() + geom line()
```

```
## Error: You're passing a function as global data.
## Have you misspelled the `data` argument in `ggplot()`
```

For this, have to give both group and colour.

The interaction plot

g



Take out interaction

```
vitaminb.2 <- update(vitaminb.1, . ~ . - ratsize:diet)
summary(vitaminb.2)</pre>
```

- No Tukey for diet: not significant.
- No Tukey for ratsize: only two sizes, and already know that obese rats have larger kidneys than lean ones.
- Bottom line: diet has no effect on kidney size once you control for size of rat.

The auto noise data

In 1973, the President of Texaco cited an automobile filter developed by Associated Octel Company as effective in reducing pollution. However, questions had been raised about the effects of filter silencing. He referred to the data included in the report (and below) as evidence that the silencing properties of the Octel filter were at least equal to those of standard silencers.

```
u <- "http://www.utsc.utoronto.ca/~butler/d29/autonoise.txt"
autonoise <- read_table(u)</pre>
```

```
## Parsed with column specification:
## cols(
## noise = col_double(),
## size = col_character(),
## type = col_character(),
## side = col_character()
##
```

The data

autonoise

```
A tibble: 36 \times 4
##
      noise size type
                          side
##
      <dbl> <chr> <chr> <chr>
##
        840 M
                   Std
                       R
        770 L
                   Octel L
##
    3
##
        820 M
                   Octel R
##
        775 L
                   Octel R
    5
##
        825 M
                   Octel L
        840 M
##
    6
                   Std
                        R.
        845 M
##
    7
                   Std
##
    8
        825 M
                   Octel L
##
        815 M
                   Octel L
## 10
        845 M
                   Std
                          R
## # ... with 26 more rows
```

Making boxplot

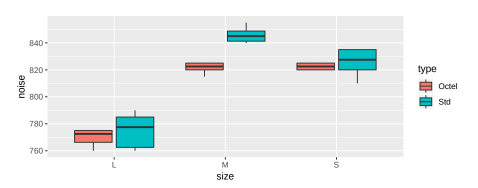
- Make a boxplot, but have combinations of filter type and engine size.
- Use grouped boxplot again, thus:

```
g <- autonoise %>%
  ggplot(aes(x = size, y = noise, fill = type)) +
  geom_boxplot()
```

The boxplot

- See difference in engine noise between Octel and standard is larger for medium engine size than for large or small.
- Some evidence of differences in spreads (ignore for now):

g



ANOVA

```
autonoise.1 <- aov(noise ~ size * type, data = autonoise)
summary(autonoise.1)</pre>
```

- The interaction is significant, as we suspected from the boxplots.
- The within-group spreads don't look very equal, but only based on 6 obs each.

Tukey: ouch!

```
autonoise.2 <- TukeyHSD(autonoise.1)
autonoise.2$`size:type`</pre>
```

```
##
                          diff
                                      lwr
                                                 upr
                                                            p adj
                    51.6666667
                                37.463511
                                           65.869823 6.033496e-11
## M:Octel-L:Octel
## S:Octel-L:Octel
                    52.5000000
                                38.296844
                                           66.703156 4.089762e-11
## I.:Std-I.:Octel
                     5.0000000
                                -9.203156
                                           19.203156 8.890358e-01
## M:Std-L:Octel
                    75.8333333
                               61.630177
                                           90.036489 4.962697e-14
## S:Std-L:Octel
                    55.8333333
                               41.630177
                                           70.036489 9.002910e-12
## S:Octel-M:Octel
                     0.8333333 -13.369823
                                           15.036489 9.999720e-01
## L:Std-M:Octel
                   -46.6666667 -60.869823 -32.463511 6.766649e-10
## M:Std-M:Octel
                    24.1666667
                                 9.963511
                                           38.369823 1.908995e-04
## S:Std-M:Octel
                     4.1666667 -10.036489
                                           18.369823 9.454142e-01
## L:Std-S:Octel
                   -47.5000000 -61.703156 -33.296844 4.477636e-10
## M:Std-S:Octel
                    23.3333333
                                 9.130177
                                           37.536489 3.129974e-04
## S:Std-S:Octel
                     3.3333333 -10.869823
                                           17.536489 9.787622e-01
## M:Std-L:Std
                    70.8333333 56.630177
                                           85.036489 6.583623e-14
## S:Std-L:Std
                    50.8333333
                                36.630177
                                           65.036489 8.937329e-11
## S:Std-M:Std
                   -20.0000000 -34.203156
                                           -5.796844 2.203265e-03
```

Interaction plot

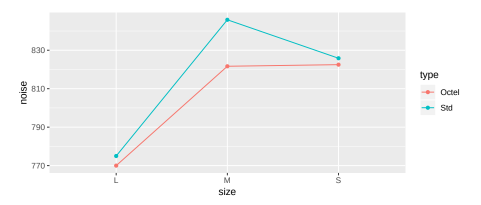
- This time, don't have summary of mean noise for each size-type combination.
- One way is to compute summaries (means) first, and feed into ggplot as in vitamin B example.
- Or, have ggplot compute them for us, thus:

```
g <- ggplot(autonoise, aes(
    x = size, y = noise,
    colour = type, group = type
)) +
    stat_summary(fun.y = mean, geom = "point") +
    stat_summary(fun.y = mean, geom = "line")</pre>
```

Interaction plot

The lines are definitely *not* parallel, showing that the effect of type is different for medium-sized engines than for others:

g



If you don't like that...

...then compute the means first, in a pipeline:

```
autonoise %>%
group_by(size, type) %>%
summarize(mean_noise = mean(noise)) %>%
ggplot(aes(
    x = size, y = mean_noise, group = type,
    colour = type
)) + geom_point() + geom_line()
```

```
## [conflicted] `summarize` found in 2 packages.
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict_prefer()`
## * conflict_prefer("summarize", "plyr")
## * conflict_prefer("summarize", "dplyr")
```

Simple effects for auto noise example

- In auto noise example, weren't interested in all comparisons between car size and filter type combinations.
- Wanted to demonstrate (lack of) difference between filter types for each car type.
- These are called **simple effects** of one variable (filter type) conditional on other variable (car type).
- To do this, pull out just the data for small cars, compare noise for the two filter types. Then repeat for medium and large cars. (Three one-way ANOVAs.)

Do it using dplyr tools

• Small cars:

```
autonoise %>%
  filter(size == "S") %>%
  aov(noise ~ type, data = .) %>%
  summary()
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## type 1 33.3 33.33 0.548 0.476
## Residuals 10 608.3 60.83
```

- No filter difference for small cars.
- For Medium, change S to M and repeat.

Simple effect of filter type for medium cars

```
autonoise %>%
 filter(size == "M") %>%
 aov(noise ~ type, data = .) %>%
 summary()
##
             Df Sum Sq Mean Sq F value Pr(>F)
        1 1752.1 1752.1 68.93 8.49e-06 ***
## type
## Residuals 10 254.2 25.4
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

• There *is* an effect of filter type for medium cars. Look at means to investigate (over).

Mean noise for each filter type

...for medium engine size:

```
autonoise %>%
  filter(size == "M") %>%
  group by(type) %>%
  summarize(m = mean(noise))
## [conflicted] `summarize` found in 2 packages.
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict prefer()`
## * conflict prefer("summarize", "plyr")
## * conflict_prefer("summarize", "dplyr")
```

Octel filters produce less noise for medium cars.

Large cars

Large cars:

```
autonoise %>%
  filter(size == "L") %>%
  aov(noise ~ type, data = .) %>%
  summary()
```

```
## Df Sum Sq Mean Sq F value Pr(>F)
## type 1 75 75 0.682 0.428
## Residuals 10 1100 110
```

• No significant difference again.

Or...

#

use glance from broom:

```
autonoise %>%
 filter(size == "L") %>%
 aov(noise ~ type, data = .) %>%
 glance()
## # A tibble: 1 x 11
##
    r.squared adj.r.squared sigma statistic p.value
                                                      df
##
        <dbl>
                      <dbl> <dbl>
                                     <dbl> <dbl> <int>
                   -0.0298 10.5 0.682 0.428
## 1
     0.0638
## # ... with 5 more variables: logLik <dbl>, AIC <dbl>,
```

BIC <dbl>, deviance <dbl>, df.residual <int>

P-value same as from summary output.

All at once, using split/apply/combine

```
The "split" part:
```

```
autonoise %>%
group_by(size) %>%
nest()
```

```
## size data
## <chr> tist>
## 1 M <tibble [12 × 3]>
## 2 L <tibble [12 × 3]>
## 3 S <tibble [12 × 3]>
```

A tibble: 3 x 2

Now have *three* rows, with the data frame for each size encoded as *one element* of this data frame.

Apply

 Write function to do aov on a data frame with columns noise and type, returning P-value:

```
aov_pval <- function(x) {
  noise.1 <- aov(noise ~ type, data = x)
  gg <- glance(noise.1)
  gg$p.value
}</pre>
```

Test it:

```
autonoise %>%
  filter(size == "L") %>%
  aov_pval()
```

[1] 0.428221

Check.

Combine

 Apply this function to each of the nested data frames (one per engine size):

```
autonoise %>%
  group_by(size) %>%
  nest() %>%
  mutate(p_val = map_dbl(data, ~ aov_pval(.)))
```

map_dbl because aov_pval returns a decimal number (a dbl).
 Investigate what happens if you use map instead.

A tibble: 3 x 3

Tidy up

 The data column was stepping-stone to getting answer. Don't need it any more:

```
simple_effects <- autonoise %>%
  group_by(size) %>%
  nest() %>%
  mutate(p_val = map_dbl(data, ~ aov_pval(.))) %>%
  select(-data)
simple_effects
```

Simultaneous tests

A tibble: 3 x 4

- When testing simple effects, doing several tests at once. (In this case,
 3.)
- Have to adjust P-values for this. Eg. Holm:

```
simple_effects %>%
  arrange(p_val) %>%
  mutate(multiplier = 4 - row_number()) %>%
  mutate(p_val_adj = p_val * multiplier)
```

* No change in rejection decisions.

^{*} Octel filters sig. better in terms of noise for medium cars, and not sig. different for other

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Confidence intervals

- Perhaps better way of assessing simple effects: look at confidence intervals rather than tests.
- Gives us sense of accuracy of estimation, and thus whether non-significance might be lack of power: "absence of evidence is not evidence of absence".
- Works here because two filter types, using t.test for each engine type.
- Want to show that the Octel filter is equivalent to or better than the standard filter, in terms of engine noise.

Equivalence and noninferiority

- Known as "equivalence testing" in medical world. A good read: link. Basic idea: decide on size of difference δ that would be considered "equivalent", and if CI entirely inside $\pm \delta$, have evidence in favour of equivalence.
- We really want to show that the Octel filters are "no worse" than the standard one: that is, equivalent *or better* than standard filters.
- Such a "noninferiority test" done by checking that upper limit of CI, new minus old, is *less* than δ . (This requires careful thinking about (i) which way around the difference is and (ii) whether a higher or lower value is better.)

CI for small cars

Same idea as for simple effect test:

```
autonoise %>%
  filter(size == "S") %>%
  t.test(noise ~ type, data = .) %>%
  .[["conf.int"]]
```

```
## [1] -14.517462 7.850795
## attr(,"conf.level")
## [1] 0.95
```

CI for medium cars

```
autonoise %>%
  filter(size == "M") %>%
  t.test(noise ~ type, data = .) %>%
  .[["conf.int"]]
```

```
## [1] -30.75784 -17.57549
## attr(,"conf.level")
## [1] 0.95
```

CI for large cars

```
autonoise %>%
  filter(size == "L") %>%
  t.test(noise ~ type, data = .) %>%
  .[["conf.int"]]
## [1] -19.270673 9.270673
```

attr(,"conf.level")

[1] 0.95

Or, all at once: split/apply/combine

```
ci_func <- function(x) {
  tt <- t.test(noise ~ type, data = x)
  tt$conf.int
}
autonoise %>%
  group_by(size) %>%
  nest() %>%
  mutate(ci = map(data, ~ ci_func(.))) %>%
  unnest(ci) -> cis
```

Results

cis

```
## # A tibble: 6 x 2
##
     size
                сi
##
     <chr> <dbl>
## 1 M
           -30.8
## 2 M
           -17.6
## 3 L
           -19.3
## 4 L
             9.27
## 5 S
           -14.5
## 6 S
             7.85
```

Procedure

- Function to get CI of difference in noise means for types of filter on input data frame
- Group by size, nest (mini-df per size)
- Calculate CI for each thing in data (ie. each size). map: CI is two numbers long
- unnest ci column to see two numbers in each CI.

Cls and noninferiority test

- Suppose we decide that a 20 dB difference would be considered equivalent. (I have no idea whether that is reasonable.)
- Intervals:

```
cis %>%
  mutate(hilo = rep(c("lower", "upper"), 3)) %>%
  spread(hilo, ci)
```

Comments

- In all cases, upper limit of CI is less than 20 dB. The Octel filters are "noninferior" to the standard ones.
- Caution: we did 3 procedures at once again. The true confidence level is not 95%. (Won't worry about that here.)

Contrasts in ANOVA

- Sometimes, don't want to compare all groups, only some of them.
- Might be able to specify these comparisons ahead of time; other comparisons of no interest.
- Wasteful to do ANOVA and Tukey.

Example: chainsaw kickback

- From link.
- Forest manager concerned about safety of chainsaws issued to field crew. 4 models of chainsaws, measure "kickback" (degrees of deflection) for 5 of each:

So far, standard 1-way ANOVA: what differences are there among models?

chainsaw kickback (2)

- But: models A and D are designed to be used at home, while models B and C are industrial models.
- Suggests these comparisons of interest:
- home vs. industrial
- the two home models A vs. D
- the two industrial models B vs. C.
- Don't need to compare all the pairs of models.

What is a contrast?

- Contrast is a linear combination of group means.
- Notation: μ_A for (population) mean of group A, and so on.
- In example, compare two home models: $H_0: \mu_A \mu_D = 0$.
- Compare two industrial models: $H_0: \mu_B \mu_C = 0$.
- Compare average of two home models vs. average of two industrial models: $H_0:\frac{1}{2}(\mu_A+\mu_D)-\frac{1}{2}(\mu_B+\mu_C)=0$ or $H_0:0.5\mu_A-0.5\mu_B-0.5\mu_C+0.5\mu_D=0$.
- Note that coefficients of contrasts add to 0, and right-hand side is 0.

Contrasts in R

• Comparing two home models A and D ($\mu_A - \mu_D = 0$):

$$c.home <- c(1, 0, 0, -1)$$

• Comparing two industrial models B and C ($\mu_B - \mu_C = 0$):

c.industrial
$$\leftarrow c(0, 1, -1, 0)$$

• Comparing home average vs. industrial average $(0.5\mu_A - 0.5\mu_B - 0.5\mu_C + 0.5\mu_D = 0)$:

c.home.ind
$$\leftarrow$$
 c(0.5, -0.5, -0.5, 0.5)

Orthogonal contrasts

• What happens if we multiply the contrast coefficients one by one?

```
c.home * c.industrial
```

```
## [1] 0 0 0 0
```

c.home * c.home.ind

```
## [1] 0.5 0.0 0.0 -0.5
```

c.industrial * c.home.ind

```
## [1] 0.0 -0.5 0.5 0.0
```

 in each case, the results add up to zero. Such contrasts are called orthogonal.

Orthogonal contrasts (2)

Compare these:

```
c1 <- c(1, -1, 0)
c2 <- c(0, 1, -1)
sum(c1 * c2)
```

```
## [1] -1
```

Not zero, so c1 and c2 are not orthogonal.

- Orthogonal contrasts are much easier to deal with.
- Can use non-orthogonal contrasts, but more trouble (beyond us).

Read in data

```
url <- "http://www.utsc.utoronto.ca/~butler/d29/chainsaw.txt"
chain.wide <- read_table(url)
chain.wide</pre>
```

```
## # A tibble: 5 x 4
##
        Α
             В
##
    <dbl> <dbl> <dbl> <dbl>
## 1
       42
            28
                 57
                       29
    17
         50 45
                       29
    24 44 48
                       22
                       34
    39 32 41
## 5
       43
            61
                 54
                       30
```

Tidying

Need all the kickbacks in one column:

```
chain <- gather(chain.wide, model, kickback, A:D,
  factor_key = T
)</pre>
```

Starting the analysis (2)

The proper data frame (tiny):

chain

```
## # A tibble: 20 x 2
      model kickback
      <fct>
                <dbl>
                   42
                   17
                   24
                   39
                   43
                   28
                   50
                   44
                   32
## 10 B
                   61
                   57
## 12 C
                   45
## 13 C
                   48
## 14 C
                   41
## 15 C
                   54
## 16 D
                   29
                   22
## 19 D
                   34
## 20 D
                   30
```

Setting up contrasts

-0.5

-0.5

0.5

[2,]

[3,]

[4,]

ANOVA as if regression

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```
chain.1 <- lm(kickback ~ model, data = chain)</pre>
summary(chain.1)
##
## Call:
## lm(formula = kickback ~ model, data = chain)
##
## Residuals:
##
     Min 10 Median 30 Max
## -16.00 -7.10 0.60 6.25 18.00
##
## Coefficients:
##
                   Estimate Std. Error t value Pr(>|t|)
                     38.450 2.179 17.649 6.52e-12 ***
## (Intercept)
## modelc.home
                  2.100 3.081 0.682 0.50524
## modelc.industrial -3.000 3.081 -0.974 0.34469
## modelc.home.ind -15.100 4.357 -3.466 0.00319 **
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 9.743 on 16 degrees of freedom
## Multiple R-squared: 0.4562, Adjusted R-squared: 0.3542
```

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Conclusions

```
tidy(chain.1) %>% select(term, p.value)
```

A tibble 4×2

- Two home models not sig. diff. (P-value 0.51)
- Two industrial models not sig. diff. (P-value 0.34)
- Home, industrial models are sig. diff. (P-value 0.0032).

Means by model

The means:

```
chain %>%
  group_by(model) %>%
  summarize(mean.kick = mean(kickback)) %>%
  arrange(desc(mean.kick))
```

```
## [conflicted] `summarize` found in 2 packages.
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict_prefer()`
## * conflict_prefer("summarize", "plyr")
## * conflict_prefer("summarize", "dplyr")
```

- Home models A & D have less kickback than industrial ones B & C.
- Makes sense because industrial users should get training to cope with additional kickback.

Section 6

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
- u 10 0
- b 7 19
- b 12 26
- b 27 33
- b 24 35
- b 18 30
- b 22 31
- b 26 34
- b 21 28
- b 14 23
- b 9 22
- ט פ ט

Packages

tidyverse and broom:

library(tidyverse)

library(broom)

Read in data

```
url <- "http://www.utsc.utoronto.ca/~butler/d29/ancova.txt"
prepost <- read delim(url, " ")</pre>
prepost %>% sample n(9) # randomly chosen rows
## # A tibble: 9 x 3
    drug before after
##
## <chr> <dbl> <dbl>
## 1 b
             24 35
            7 19
## 2 b
           18 30
## 3 b
          21 40
## 4 a
          12 26
## 5 b
## 6 a
           14
                27
           22 31
## 7 b
           14 23
## 8 b
```

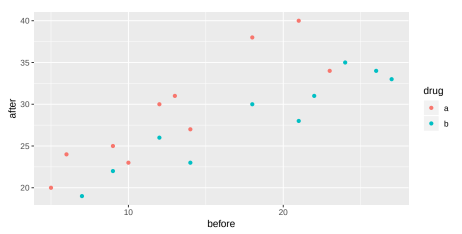
9 a

12

30

Making a plot

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



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.

The means

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

## [conflicted] `summarize` found in 2 packages.
```

```
## Either pick the one you want with `::`
## * plyr::summarize
## * dplyr::summarize
## Or declare a preference with `conflict_prefer()`
## * conflict_prefer("summarize", "plyr")
## * conflict_prefer("summarize", "dplyr")
```

Mean "after" score slightly higher for treatment A.

```
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```

Testing for interaction

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

```
## Analysis of Variance Table
##
## Response: after
              Df Sum Sq Mean Sq F value Pr(>F)
##
## before 1 430.92 430.92 62.6894 6.34e-07 ***
## drug
         1 115.31 115.31 16.7743 0.0008442 ***
## before:drug 1 12.34 12.34 1.7948 0.1990662
## Residuals 16 109.98 6.87
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

• Interaction not significant. Will remove later.

Predictions, with interaction included

Make combinations of before score and drug:

```
before = c(5, 15, 25),
  drug = c("a", "b")
new
## # A tibble: 6 \times 2
##
     before drug
      <dbl> <chr>
##
          5 a
## 1
## 2
          5 b
     15 a
     15 b
## 4
     25 a
         25 b
```

new <- crossing(</pre>

Do predictions:

```
pred <- predict(prepost.1, new)
preds <- bind_cols(new, pred = pred)
preds</pre>
```

```
## # A tibble: 6 \times 3
##
    before drug
                  pred
     <dbl> <chr> <dbl>
##
                  21.3
## 1
         5 a
## 2
         5 b
                  18.7
       15 a
                  31.1
## 3
## 4
     15 b
               25.9
## 5
    25 a
               40.8
## 6
        25 b
                  33.2
```

Making a plot with lines for each drug

```
g <- ggplot(prepost,
  aes(x = before, y = after, colour = drug)) +
  geom_point() + geom_line(data = preds, aes(y = pred))</pre>
```

- Here, final line:
 - joins points by lines for different data set (preds rather than prepost),
 - different y (pred rather than after),
 - but same x (x=before inherited from first aes).
- Last line could (more easily) be

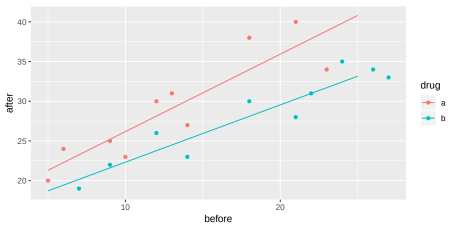
```
geom_smooth(method = "lm", se = F)
```

which would work here, but not for later plot.

The plot

Lecture notes

- Lines almost parallel, but not quite.
- Non-parallelism (interaction) not significant:



Taking out interaction

```
prepost.2 <- update(prepost.1, . ~ . - before:drug)</pre>
anova(prepost.2)
## Analysis of Variance Table
##
## Response: after
##
            Df Sum Sq Mean Sq F value Pr(>F)
## before 1 430.92 430.92 59.890 5.718e-07 ***
## drug 1 115.31 115.31 16.025 0.0009209 ***
## Residuals 17 122.32 7.20
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

- Take out non-significant interaction.
- before and drug strongly significant.
- Do predictions again and plot them.

Predicted values again (no-interaction model)

```
pred <- predict(prepost.2, new)
preds <- bind_cols(new, pred = pred)
preds</pre>
```

```
##
   before drug
               pred
##
    <dbl> <chr> <dbl>
        5 a 22.5
## 1
## 2
        5 b
           17.3
   15 a
               30.8
## 3
    15 b 25.6
## 4
   25 a 39.0
## 5
## 6
       25 b
               33.9
```

A tibble: 6×3

Each increase of 10 in before score results in 8.3 in predicted after score, the same for both drugs.

Making a plot, again

```
g <- ggplot(
  prepost,
  aes(x = before, y = after, colour = drug)
) +
  geom_point() +
  geom_line(data = preds, aes(y = pred))</pre>
```

Exactly same as before, but using new predictions.

The no-interaction plot of predicted values

g

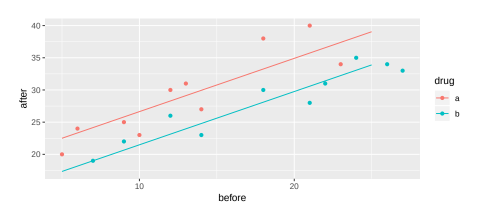


Figure 39: plot of chunk cabazzo

Different look at model output

summary(prepost.2)

- anova(prepost.2) tests for significant effect of before score and of drug, but doesn't help with interpretation.
- summary(prepost.2) views as regression with slopes:

```
##
## 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:
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```

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.

Section 7

Multivariate ANOVA

Multivariate analysis of variance

- Standard ANOVA has just one response variable.
- What if you have more than one response?
- Try an ANOVA on each response separately.
- But might miss some kinds of interesting dependence between the responses that distinguish the groups.

Packages

```
library(car)
library(tidyverse)
```

Small example

- Measure yield and seed weight of plants grown under 2 conditions: low and high amounts of fertilizer.
- Data (fertilizer, yield, seed weight):

```
url <- "http://www.utsc.utoronto.ca/~butler/d29/manova1.txt"
hilo <- read_delim(url, " ")

## Parsed with column specification:
## cols(</pre>
```

```
## cols(
## fertilizer = col_character(),
## yield = col_double(),
## weight = col_double()
## )
```

2 responses, yield and seed weight.

The data

hilo

```
## # A tibble: 8 x 3
##
    fertilizer yield weight
    <chr>
             <dbl> <dbl>
##
                        10
## 1 low
                 34
                        14
## 2 low
                 29
## 3 low
                 35
                        11
## 4 low
                 32
                       13
## 5 high
                 33
                        14
                        12
## 6 high
                 38
## 7 high
                 34
                        13
                        14
## 8 high
                 35
```

Boxplot for yield for each fertilizer group

ggplot(hilo, aes(x = fertilizer, y = yield)) + geom_boxplot()



Figure 40: plot of chunk ferto

Boxplot for weight for each fertilizer group

ggplot(hilo, aes(x = fertilizer, y = weight)) + geom_boxplot()

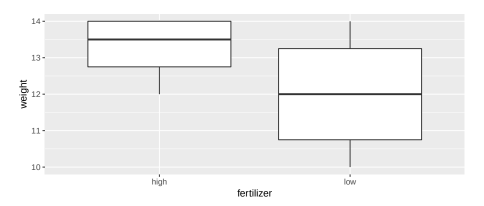


Figure 41: plot of chunk casteldisangro

ANOVAs for yield and weight

Residuals 6 12.750 2.125

```
hilo.y <- aov(yield ~ fertilizer, data = hilo)
summary(hilo.y)

## Df Sum Sq Mean Sq F value Pr(>F)
## fertilizer 1 12.5 12.500 2.143 0.194
## Residuals 6 35.0 5.833
hilo.w <- aov(weight ~ fertilizer, data = hilo)
summary(hilo.w)
```

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

Neither response depends significantly on fertilizer. But...

fertilizer 1 3.125 3.125 1.471 0.271

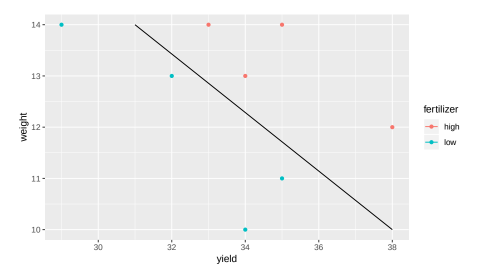
##

Plotting both responses at once

- Have two response variables (not more), so can plot the response variables against each other, labelling points by which fertilizer group they're from.
- \bullet First, create data frame with points (31,14) and (38,10) (why? Later):

• Then plot data as points, and add line through points in d:

The plot



Comments

- Graph construction:
 - Joining points in d by line.
 - geom_line inherits colour from aes in ggplot.
 - Data frame d has no fertilizer (previous colour), so have to unset.
- Results:
 - High-fertilizer plants have both yield and weight high.
 - True even though no sig difference in yield or weight individually.
 - Drew line separating highs from lows on plot.

MANOVA finds multivariate differences

• Is difference found by diagonal line significant? MANOVA finds out.

```
response <- with(hilo, cbind(yield, weight))
hilo.1 <- manova(response ~ fertilizer, data = hilo)
summary(hilo.1)</pre>
```

```
## Df Pillai approx F num Df den Df Pr(>F)
## fertilizer 1 0.80154 10.097 2 5 0.01755 *
## Residuals 6
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

 Yes! Difference between groups is diagonally, not just up/down (weight) or left-right (yield). The yield-weight combination matters.

Strategy

- Create new response variable by gluing together columns of responses, using cbind.
- Use manova with new response, looks like 1m otherwise.
- With more than 2 responses, cannot draw graph. What then?
- If MANOVA test significant, cannot use Tukey. What then?
- Use discriminant analysis (of which more later).

Another way to do MANOVA

Install (once) and load package car:

library(car)

Another way...

```
hilo.2.lm <- lm(response ~ fertilizer, data = hilo)
hilo.2 <- Manova(hilo.2.lm)
hilo.2
##
## Type II MANOVA Tests: Pillai test statistic
##
             Df test stat approx F num Df den Df Pr(>F)
## fertilizer 1 0.80154 10.097 2 5 0.01755 *
## Signif. codes:
## 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

- Same result as small-m manova.
- Manova will also do repeated measures, coming up later.

Another example: peanuts

- Three different varieties of peanuts (mysteriously, 5, 6 and 8) planted in two different locations.
- Three response variables: y, smk and w.

```
u <- "http://www.utsc.utoronto.ca/~butler/d29/peanuts.txt"
peanuts.orig <- read_delim(u, " ")</pre>
```

```
## Parsed with column specification:
## cols(
## obs = col_double(),
## location = col_double(),
## variety = col_double(),
## y = col_double(),
## smk = col_double(),
## w = col_double()
## "
```

The data

peanuts.orig

```
## # A tibble: 12 x 6
##
         obs location variety
                                          smk
##
      <dbl>
                 <dbl>
                          <dbl> <dbl> <dbl> <dbl> <dbl>
##
           1
                               5
                                  195.
                                         153.
                                                51.4
##
    2
                               5
                                  194.
                                         168.
                                                53.7
##
    3
           3
                               5
                                  190.
                                       140.
                                                55.5
           4
                              5
                                  180.
##
                                         121.
                                                44.4
    5
           5
                                  203
                                         157.
                                                49.8
##
                              6
    6
           6
                                  196.
                                         166
                                                45.8
##
                              6
                                  203.
##
                              6
                                         166.
                                                60.4
##
    8
           8
                              6
                                  198.
                                         162.
                                                54.1
           9
                                  194.
                                         164.
                                                57.8
##
                              8
                                  187
##
   10
          10
                              8
                                         165.
                                                58.6
## 11
          11
                              8
                                  202.
                                         167.
                                                65
## 12
          12
                               8
                                  200
                                         174.
                                                67.2
```

Setup for analysis

```
peanuts <- peanuts.orig %>%
  mutate(
    location = factor(location),
    variety = factor(variety)
response <- with(peanuts, cbind(y, smk, w))
head(response)
##
                smk
   [1.] 195.3 153.1 51.4
   [2,] 194.3 167.7 53.7
   [3,] 189.7 139.5 55.5
  [4,] 180.4 121.1 44.4
## [5,] 203.0 156.8 49.8
## [6,] 195.9 166.0 45.8
```

Analysis (using Manova)

1 0.89348 11.1843

2 1.70911 9.7924 6 10

3

10

```
## location:variety 2 1.29086 3.0339 6

## Pr(>F)

## location 0.020502 *

## variety 0.001056 **

## location:variety 0.058708 .

## ---
```

Signif. codes:

location

variety

0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Comments

- Interaction not quite significant, but main effects are.
- Combined response variable (y,smk,w) definitely depends on location and on variety
- Weak dependence of (y,smk,w) on the location-variety combination.
- Understanding that dependence beyond our scope right now.

Section 8

Repeated measures by profile analysis

Repeated measures by profile analysis

- More than one response measurement for each subject. Might be
- measurements of the same thing at different times
- measurements of different but related things
- Generalization of matched pairs ("matched triples", etc.).
- Variation: each subject does several different treatments at different times (called *crossover design*).
- Expect measurements on same subject to be correlated, so assumptions of independence will fail.
- Called *repeated measures*. Different approaches, but *profile analysis* uses Manova (set up right way).
- Another approach uses mixed models (random effects).

Packages

```
library(car)
library(tidyverse)
```

Example: histamine in dogs

- 8 dogs take part in experiment.
- Dogs randomized to one of 2 different drugs.
- Response: log of blood concentration of histamine 0, 1, 3 and 5 minutes after taking drug. (Repeated measures.)
- Data in dogs.txt, column-aligned.

Read in data

```
my url <- "http://www.utsc.utoronto.ca/~butler/d29/dogs.txt"
dogs <- read table(my url)</pre>
## Parsed with column specification:
## cols(
##
     dog = col_character(),
##
     drug = col character(),
     x = col character(),
##
##
     lh0 = col double(),
     lh1 = col double(),
##
##
     lh3 = col double(),
     lh5 = col double()
##
```

)

dogs.1 <- lm(response ~ drug, data = dogs)</pre>

Setting things up

dogs

```
## # A tibble: 8 x 7
##
                               1h0
                                     lh1
                                           1h3
                                                 1h5
    dog
          drug
                       X
##
    <chr> <chr>
                       <chr> <dbl> <dbl> <dbl> <dbl> <dbl>
                             -3.22 - 1.61 - 2.3 - 2.53
## 1 A
          Morphine
                       N
## 2 B
          Morphine
                             -3.91 -2.81 -3.91 -3.91
## 3 C
          Morphine N -2.66 \ 0.34 \ -0.73 \ -1.43
                         -1.77 -0.56 -1.05 -1.43
## 4 D
          Morphine
          Trimethaphan N
                             -3.51 - 0.48 - 1.17 - 1.51
## 5 E
## 6 F
          Trimethaphan N
                             -3.51 0.05 -0.31 -0.51
                             -2.66 -0.19 0.07 -0.22
## 7 G
          Trimethaphan N
## 8 H
          Trimethaphan N
                             -2.41 1.14 0.72 0.21
response <- with(dogs, cbind(lh0, lh1, lh3, lh5))
```

The repeated measures MANOVA

Get list of response variable names; we call them times. Save in data frame.

```
times <- colnames(response)
times.df <- data.frame(times)
dogs.2 <- Manova(dogs.1,
   idata = times.df,
   idesign = ~times
)
dogs.2</pre>
```

```
##
## Type II Repeated Measures MANOVA Tests: Pillai test statistic
## Df test stat approx F num Df den Df Pr(>F)
## (Intercept) 1 0.76347 19.3664 1 6 0.004565 **
## drug 1 0.34263 3.1272 1 6 0.127406
## times 1 0.94988 25.2690 3 4 0.004631 **
## drug:times 1 0.89476 11.3362 3 4 0.020023 *
## ---
## Signif. codes:
```

Wide and long format

- Interaction significant. Pattern of response over time different for the two drugs.
- Want to investigate interaction.

The wrong shape

dogs %>% slice(1:6)

But data frame has several observations per line ("wide format"):

```
## # A tibble: 6 x 7
##
    dog
          drug
                       x
                               1h0
                                     lh1
                                           1h3
                                                 1h5
    <chr> <chr>
                       <chr> <dbl> <dbl> <dbl> <dbl> <dbl>
##
                             -3.22 -1.61 -2.3 -2.53
## 1 A
          Morphine
          Morphine
                       N -3.91 -2.81 -3.91 -3.91
## 2 B
          Morphine
                       N -2.66 0.34 -0.73 -1.43
## 3 C
          Morphine
                         -1.77 -0.56 -1.05 -1.43
## 4 D
## 5 E
          Trimethaphan N
                         -3.51 -0.48 -1.17 -1.51
          Trimethaphan N
## 6 F
                             -3.51 0.05 -0.31 -0.51
```

- Plotting works with data in "long format": one response per line.
- The responses are log-histamine at different times, labelled 1h-something. Call them all 1h and put them in one column, with the time they belong to labelled.

Running gather, try 1

```
dogs %>% gather(time, lh, lh0:lh5)
    A tibble: 32 x 5
                                         1h
##
      dog
            drug
                               time
                         х
      <chr> <chr>
##
                         <chr> <chr> <dbl>
                               1h0
##
    1 A
            Morphine
                                      -3.22
##
   2 B
            Morphine
                               lh0
                                      -3.91
##
   3 C
            Morphine
                               1h0
                                      -2.66
   4 D
            Morphine
                               1h0
                                      -1.77
##
##
   5 E
            Trimethaphan N
                               lh0
                                      -3.51
##
   6 F
            Trimethaphan N
                               1h0
                                      -3.51
##
   7 G
            Trimethaphan N
                               1h0
                                      -2.66
##
   8 H
            Trimethaphan N
                               1h0
                                      -2.41
   9 A
            Morphine
                               lh1
                                      -1.61
##
##
   10 B
            Morphine
                               lh1
                                      -2.81
   # ... with 22 more rows
```

Getting the times

gather(timex, lh, lh0:lh5) %>%

dogs %>%

Not quite right: for the times, we want just the numbers, not the letters 1h every time. Want new variable containing just number in time: parse_number.

```
mutate(time = parse number(timex))
    A tibble: 32 x 6
##
      dog
                               timex
                                         1h
                                             time
            drug
                         х
##
      <chr> <chr>
                         <chr> <chr> <dbl> <dbl>
##
    1 A
            Morphine
                         N
                               1h0
                                      -3.22
##
   2 B
            Morphine
                         N
                               lh0 -3.91
                                                0
                         N
##
   3 C
            Morphine
                               lh0
                                      -2.66
                                                0
            Morphine
                         N
                               1h0 -1.77
##
   4 D
                                                0
##
   5 F.
            Trimethaphan N
                               lh0 -3.51
                                                0
            Trimethaphan N
                               1h0
##
   6 F
                                     -3.51
                                                0
   7 G
            Trimethaphan N
                               1h0
                                      -2.66
##
                                                0
   8 H
            Trimethaphan N
                               lh0
                                      -2.41
                                                0
##
            Morphine
                                lh1
                                      -1.61
##
   9 A
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```

What I did differently

- I realized that gather was going to produce something like 1h1, which
 I needed to do something further with, so this time I gave it a
 temporary name timex.
- This enabled me to use the name time for the actual numeric time.
- This works now, so next save into a new data frame dogs.long.

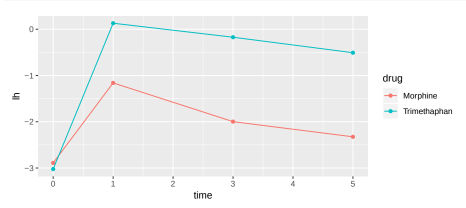
Saving the pipelined results

```
dogs %>%
  gather(timex, lh, lh0:lh5) %>%
  mutate(time = parse_number(timex)) -> dogs.long
```

This says:

- Take data frame dogs, and then:
- Combine the columns 1h0 through 1h5 into one column called 1h, with the column that each 1h value originally came from labelled by timex, and then:
- Pull out numeric values in timex, saving in time and then:
- save the result in a data frame dogs.long.

Interaction plot



Comments

- Plot mean 1h value at each time, joining points on same drug by lines.
- drugs same at time 0
- after that, Trimethaphan higher than Morphine.
- Effect of drug not consistent over time: significant interaction.

Take out time zero

- Lines on interaction plot would then be parallel, and so interaction should no longer be significant.
- Go back to original "wide" dogs data frame.

```
response <- with(dogs, cbind(lh1, lh3, lh5)) # excl time 0
dogs.1 <- lm(response ~ drug, data = dogs)
times <- colnames(response)
times.df <- data.frame(times)
dogs.2 <- Manova(dogs.1,
   idata = times.df,
   idesign = ~times
)</pre>
```

Results and comments

dogs.2

```
##
## Type II Repeated Measures MANOVA Tests: Pillai test statistic
## Df test stat approx F num Df den Df Pr(>F)
## (Intercept) 1 0.54582 7.2106 1 6 0.036281 *
## drug 1 0.44551 4.8207 1 6 0.070527 .
## times 1 0.85429 14.6569 2 5 0.008105 **
## drug:times 1 0.43553 1.9289 2 5 0.239390
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

- Correct: interaction no longer significant.
- Significant effect of time.
- Drug effect not quite significant (some variety among dogs within drug).

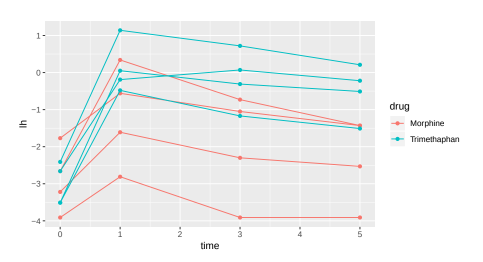
Is the non-significant drug effect reasonable?

- Plot actual data: 1h against days, labelling observations by drug: "spaghetti plot".
- Uses long data frame (confusing, yes I know):
- Plot (time,lh) points coloured by drug
- and connecting measurements for each dog by lines.
- This time, we want group=dog (want the measurements for each dog joined by lines), but colour=drug:

```
g <- ggplot(dogs.long, aes(
   x = time, y = lh,
   colour = drug, group = dog
)) +
   geom_point() + geom_line()</pre>
```

The spaghetti plot

g



Comments

- For each dog over time, there is a strong increase and gradual decrease in log-histamine. This explains the significant time effect.
- The pattern is more or less the same for each dog, regardless of drug. This explains the non-significant interaction.
- Most of the trimethaphan dogs (blue) have higher log-histamine throughout (time 1 and after), and some of the morphine dogs have lower.
- But two of the morphine dogs have log-histamine profiles like the trimethaphan dogs. This ambiguity is probably why the drug effect is not quite significant.

The exercise data

- 30 people took part in an exercise study.
- Each subject was randomly assigned to one of two diets ("low fat" or "non-low fat") and to one of three exercise programs ("at rest", "walking", "running").
- There are $2 \times 3 = 6$ experimental treatments, and thus each one is replicated 30/6 = 5 times.
- Nothing unusual so far.
- However, each subject had their pulse rate measured at three different times (1, 15 and 30 minutes after starting their exercise), so have repeated measures.

Reading the data

```
Separated by tabs:
url <- "http://www.utsc.utoronto.ca/~butler/d29/exercise.txt"
exercise.long <- read_tsv(url)

## Parsed with column specification:
## cols(
## id = col_double(),
## diet = col_character(),
## exertype = col_character(),
## pulse = col_double(),</pre>
```

time = col character()

##

)

The data

```
exercise.long %>% slice(1:8)
```

```
## # A tibble: 8 x 5
       id diet exertype pulse time
##
##
    <dbl> <chr> <chr>
                          <dbl> <chr>
       1 nonlowfat atrest
## 1
                            85 min01
## 2
       1 nonlowfat atrest
                            85 min15
## 3
       1 nonlowfat atrest
                            88 min30
## 4
       2 nonlowfat atrest 90 min01
     2 nonlowfat atrest 92 min15
## 5
     2 nonlowfat atrest 93 min30
## 6
       3 nonlowfat atrest
## 7
                            97 min01
## 8
       3 nonlowfat atrest
                            97 min15
```

- This is "long format", which is usually what we want.
- But for repeated measures analysis, we want wide format!
- "undo" gather: spread.

Making wide format

 spread needs: a column that is going to be split, and the column to make the values out of:

```
exercise.long %>% spread(time, pulse) -> exercise.wide
exercise.wide %>% sample_n(5)

## # A tibble: 5 x 6
## id diet exertype min01 min15 min30
```

```
<chr>
##
   <dbl> <chr>
                        <dbl> <dbl> <dbl>
      25 nonlowfat running
## 1
                          94
                              110
                                   116
      15 nonlowfat walking
                               96 95
## 2
                          89
## 3
      19 lowfat
                walking
                          97
                               98 100
## 4 8 lowfat.
                atrest
                          92
                               94
                                  95
## 5
       2 nonlowfat atrest
                          90
                               92
                                    93
```

 Normally gather min01, min15, min30 into one column called pulse labelled by the number of minutes. But Manova needs it the other way.

Setting up the repeated-measures analysis

Make a response variable consisting of min01, min15, min30:

```
response <- with(exercise.wide, cbind(min01, min15, min30))
```

• Predict that from diet and exertype and interaction using lm:

```
exercise.1 <- lm(response ~ diet * exertype,
  data = exercise.wide
)</pre>
```

• Run this through Manova:

Results

exercise.2

```
##
  Type II Repeated Measures MANOVA Tests: Pillai test statistic
##
                      Df test stat approx F num Df den Df
                           0.99767
                                   10296.7
                                                       24 < 2.2e-16 ***
   (Intercept)
## diet
                           0.37701
                                       14.5
                                                       24 0.0008483 ***
                           0.79972
                                       47.9
                                                 2
                                                       24 4.166e-09 ***
## exertype
                        2 0.28120 4.7
## diet:exertype
                                                       24 0.0190230 *
## times
                           0.78182
                                   41.2
                                                       23 2.491e-08 ***
                           0.25153
                                        3.9
                                                       23 0.0357258 *
## diet:times
                       2 0.83557
                                        8.6
                                                 4
                                                       48 2.538e-05 ***
## exertype:times
                           0.51750
                                        4.2
                                                 4
                                                       48 0.0054586 **
## diet:exertype:times
## ---
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

- Three-way interaction significant, so cannot remove anything.
- Pulse rate depends on diet and exercise type *combination*, and *that* is different for each time.

Making some graphs

- Three-way interactions are difficult to understand. To make an attempt, look at some graphs.
- Plot time trace of pulse rates for each individual, joined by lines, and make separate plots for each diet-exertype combo.
- ggplot again. Using long data frame:

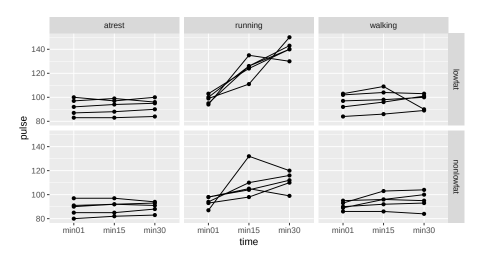
```
g <- ggplot(exercise.long, aes(
   x = time, y = pulse,
   group = id
)) + geom_point() + geom_line() +
   facet_grid(diet ~ exertype)</pre>
```

• facet_grid(diet~exertype): do a separate plot for each combination of diet and exercise type, with diets going down the page and exercise types going across. (Graphs are usually landscape, so have the factor exertype with more levels going across.)

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The graph(s)

g



Comments on graphs

- For subjects who were at rest, no change in pulse rate over time, for both diet groups.
- For walking subjects, not much change in pulse rates over time. Maybe a small increase on average between 1 and 15 minutes.
- For both running groups, an overall increase in pulse rate over time, but the increase is stronger for the lowfat group.
- No consistent effect of diet over all exercise groups.
- No consistent effect of exercise type over both diet groups.
- No consistent effect of time over all diet-exercise type combos.

"Simple effects" of diet for the subjects who ran

- Looks as if there is only any substantial time effect for the runners. For them, does diet have an effect?
- Pull out only the runners from the wide data:

```
exercise.wide %>%
filter(exertype == "running") -> runners.wide
```

 Create response variable and do MANOVA. Some of this looks like before, but I have different data now:

```
response <- with(runners.wide, cbind(min01, min15, min30))
runners.1 <- lm(response ~ diet, data = runners.wide)
times <- colnames(response)
times.df <- data.frame(times)
runners.2 <- Manova(runners.1,
   idata = times.df,
   idesign = ~times
)</pre>
```

Results

runners.2

```
##
## Type II Repeated Measures MANOVA Tests: Pillai test statistic
            Df test stat approx F num Df den Df
##
                         9045.3
                                         8 1.668e-13 ***
## (Intercept) 1
                0.99912
## diet
             1 0.84986 45.3
                                   1 8 0.0001482 ***
## times 1 0.92493 43.1 2 7 0.0001159 ***
## diet:times 1 0.68950 7.8 2
                                         7 0.0166807 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

text under

- The diet by time interaction is still significant (at $\alpha=0.05$): the effect of time on pulse rates is different for the two diets.
- At $\alpha=0.01$, the interaction is not significant, and then we have only two (very) significant main effects of diet and time.

How is the effect of diet different over time?

 Table of means. Only I need long data for this, so make it (in a pipeline):

```
runners.wide %>%
  gather(time, pulse, min01:min30) %>%
  group_by(time, diet) %>%
  summarize(
   mean = mean(pulse),
   sd = sd(pulse)
) -> summ
```

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Understanding diet-time interaction

• The summary:

summ

```
## Error in eval(expr, envir, enclos): object 'summ' not found
```

- Pulse rates at any given time higher for lowfat (diet effect),
- Pulse rates increase over time of exercise (time effect),
- but the amount by which pulse rate higher for a diet depends on time: diet by time interaction.

Interaction plot

 We went to trouble of finding means by group, so making interaction plot is now mainly easy:

```
## Error in ggplot(summ, aes(x = time, y = mean, colour = die
```

Comment on interaction plot

- The lines are not parallel, so there is interaction between diet and time for the runners.
- The effect of time on pulse rate is different for the two diets, even though all the subjects here were running.

Section 9

Discriminant analysis

Discriminant analysis

- ANOVA and MANOVA: predict a (counted/measured) response from group membership.
- Discriminant analysis: predict group membership based on counted/measured variables.
- Covers same ground as logistic regression (and its variations), but emphasis on classifying observed data into correct groups.
- Does so by searching for linear combination of original variables that best separates data into groups (canonical variables).
- Assumption here that groups are known (for data we have). If trying to "best separate" data into unknown groups, see cluster analysis xxx.
- Examples: revisit seed yield and weight data, peanut data, professions/activities data; remote-sensing data.

Packages xxx

```
library(MASS)
library(tidyverse)
library(ggrepel)
library(ggbiplot)
```

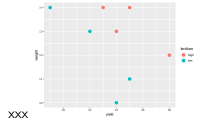
ggrepel allows labelling points on a plot so they don't overwrite each other.

About select

- Both dplyr (in tidyverse) and MASS have a function called select, and they do different things.
- How do you know which select is going to get called?
- With library, the one loaded last is visible, and others are not.
- Thus we can access the select in dplyr but not the one in MASS. If we wanted that one, we'd have to say MASS::select.
- I loaded MASS before tidyverse. If I had done it the other way around, the tidyverse select, which I want to use, would have been the invisible one.
- xxx Alternative: load conflicted package. Any time you load two
 packages containing functions with same name, you get error and have
 to choose between them.

Example 1: seed yields and weights xxx

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/manova1.txt"
hilo <- read_delim(my_url, " ")
g <- ggplot(hilo, aes(
    x = yield, y = weight,
    colour = fertilizer
)) + geom_point(size = 4)</pre>
```



Recall data from MANOVA: needed a multivariate analy-

Basic discriminant analysis

```
hilo.1 <- lda(fertilizer ~ yield + weight, data = hilo)
```

- Uses 1da from package MASS.
- "Predicting" group membership from measured variables.

Output

```
hilo.1
## Call:
## lda(fertilizer ~ yield + weight, data = hilo)
##
## Prior probabilities of groups:
## high low
## 0.5 0.5
##
## Group means:
## yield weight
## high 35.0 13.25
## low 32.5 12.00
##
## Coefficients of linear discriminants:
##
                 I.D1
## yield -0.7666761
## weight -1.2513563
```

Things to take from output

- Group means: high-fertilizer plants have (slightly) higher mean yield and weight than low-fertilizer plants.
- "Coefficients of linear discriminants": LD1, LD2,...are scores constructed from observed variables that best separate the groups.
- For any plant, get LD1 score by taking -0.76 times yield plus -1.25 times weight, add up, standardize.
- the LD1 coefficients are like slopes:
 - if yield higher, LD1 score for a plant lower
 - if weight higher, LD1 score for a plant lower
- High-fertilizer plants have higher yield and weight, thus low (negative)
 LD1 score. Low-fertilizer plants have low yield and weight, thus high (positive)
 LD1 score. xxx
- One LD1 score for each observation. Plot with actual groups.

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How many linear discriminants?

- Smaller of these: xxx
 - Number of variables
 - Number of groups minus 1
- Seed yield and weight: 2 variables, 2 groups, $\min(2, 2-1) = 1$.

Getting LD scores xxx

Feed output from LDA into predict:

```
hilo.pred <- predict(hilo.1)
```

Component x contains LD score(s), here in descending order:

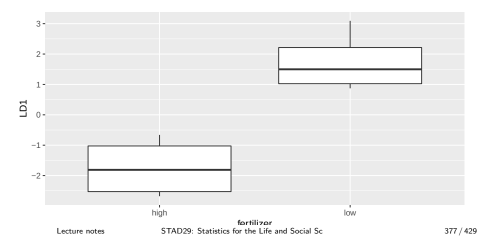
```
d <- cbind(hilo, hilo.pred$x) %>% arrange(desc(LD1))
d
```

```
LD1
##
    fertilizer yield weight
## 1
           low
                  34
                         10 3.0931414
                         14 1.9210963
## 2
           low
               29
               35 11 1.0751090
## 3
           low
## 4
           low
                  32
                         1.3
                             0.8724245
                  34
## 5
          high
                         13 -0.6609276
                  33
## 6
          high
                         14 -1.1456079
## 7
          high
                  38
                         12 - 2.4762756
                  35
                         14 -2.6789600
## 8
          high
```

xxx High fertilizer have yield and weight high, negative LD1 scores.

Plotting LD1 scores

With one LD score, plot against (true) groups, eg. boxplot: xxx



Potentially misleading

These are like regression slopes: xxx

hilo.1\$scaling

```
## LD1
## yield -0.7666761
## weight -1.2513563
```

- Reflect change in LD1 score for 1-unit change in variables.
- But one-unit change in variables might not be comparable:

summary(hilo)

```
fertilizer
                            vield
##
                                             weight
                                                :10.00
    Length:8
                        Min.
                               :29.00
                                        Min.
##
    Class :character
                        1st Qu.:32.75
                                         1st Qu.:11.75
##
##
    Mode :character
                        Median :34.00
                                         Median :13.00
                                         Mean :12.62
##
                        Mean :33.75
                        3rd Qu.:35.00
                                         3rd Qu.:14.00
##
##
                        Max. :38.00
                                         Max. :14.00
       Lecture notes
                        STAD29: Statistics for the Life and Social Sc.
```

What else is in hilo.pred?

```
names(hilo.pred)
```

```
## [1] "class" "posterior" "x"
```

- class: predicted fertilizer level (based on values of yield and weight).
- posterior: predicted probability of being low or high fertilizer given yield and weight.

Predictions and predicted groups

```
...based on yield and weight: xxx
cbind(hilo, predicted = hilo.pred$class)
```

```
fertilizer yield weight predicted
##
                           10
## 1
            low
                    34
                                     low
## 2
            low
                   29
                           14
                                     low
## 3
            low
                   35
                           11
                                     low
## 4
            low
                   32
                           13
                                    low
                   33
## 5
           high
                           14
                                   high
                   38
                       12
## 6
           high
                                   high
## 7
           high
                   34
                           13
                                    high
## 8
           high
                    35
                           14
                                    high
```

```
table(obs = hilo$fertilizer, pred = hilo.pred$class)
```

```
## pred
## obs high low
## high 4 0
## low 0 4
```

Understanding the predicted groups

- Each predicted fertilizer level is exactly same as observed one (perfect prediction).
- Table shows no errors: all values on top-left to bottom-right diagonal.

Posterior probabilities

show how clear-cut the classification decisions were:

```
pp <- round(hilo.pred$posterior, 4)
d <- cbind(hilo, hilo.pred$x, pp)
d</pre>
```

```
##
     fertilizer yield weight
                                    LD1
                                          high
                                                  low
## 1
            IOW
                  34
                          10 3.0931414 0.0000 1.0000
## 2
           low
                29
                          14 1.9210963 0.0012 0.9988
                35
## 3
           low
                          11 1.0751090 0.0232 0.9768
           low
                32
                          13
                             0.8724245 0.0458 0.9542
## 4
## 5
                  33
                          14 -1.1456079 0.9818 0.0182
          high
## 6
          high
                  38
                          12 -2.4762756 0.9998 0.0002
          high
                  34
                          13 -0.6609276 0.9089 0.0911
## 7
                  35
                          14 -2.6789600 0.9999 0.0001
## 8
          high
```

Only obs. 7 has any doubt: yield low for a high-fertilizer, but high weight makes up for it.

Example 2: the peanuts xxx

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/peanuts.txt"
peanuts <- read_delim(my_url, " ")
peanuts</pre>
```

```
A tibble: 12 x 6
##
       obs location variety
                                    smk
     <dbl>
              <dbl>
                      <dbl> <dbl> <dbl> <dbl> <
##
##
                             195.
                                   153.
                                         51.4
                            194.
                                   168. 53.7
##
##
                          5 190.
                                   140.
                                        55.5
         4
                                   121. 44.4
##
                            180.
##
                             203
                                   157. 49.8
                            196.
                                   166
                                        45.8
##
##
                          6 203.
                                   166. 60.4
##
         8
                          6 198.
                                   162. 54.1
   8
##
                          8 194.
                                   164.
                                         57.8
        10
                            187 165. 58.6
##
  10
        11
                          8 202.
                                   167.
                                         65
                                         67.2
## 12
        12
                             200
                                   174.
```

 Recall: location and variety both significant in MANOVA. Make combo of them (over):

Location-variety combos xxx

```
peanuts %% unite(combo, c(variety, location)) ->
peanuts.combo
peanuts.combo
```

```
## # A tibble: 12 x 5
##
       obs combo
                       smk
##
     <dbl> <chr> <dbl> <dbl> <dbl> <dbl>
         1 5_1
                 195.
                      153. 51.4
##
  1
   2
         2 5_1 194. 168. 53.7
##
         3 5 2 190. 140. 55.5
##
   4
         4 5 2 180. 121. 44.4
##
##
   5
         5 6 1
                 203 157.
                            49.8
##
   6
         6 6 1
              196. 166 45.8
                 203. 166. 60.4
##
         7 6 2
         8 6 2
                 198. 162. 54.1
##
   8
         9 8 1
              194.
                      164.
                           57.8
##
## 10
        10 8_1
                 187 165. 58.6
                 202. 167. 65
## 11
        11 8_2
## 12
        12 8 2
                 200
                       174.
                            67.2
```

١ -

Discriminant analysis

```
peanuts.1 <- lda(combo ~ y + smk + w, data = peanuts.combo)
peanuts.1$scaling</pre>
```

```
## LD1 LD2 LD3

## y -0.4027356 -0.02967881 0.18839237

## smk -0.1727459 0.06794271 -0.09386294

## w 0.5792456 0.16300221 0.07341123
```

peanuts.1\$svd

```
## [1] 6.141323 2.428396 1.075589
```

• Now 3 LDs (3 variables, 6 groups, min(3, 6-1) = 3).

Comments xxx

- First: relationship of LDs to original variables. Look for coeffs far from zero: here,
- high LD1 mainly high w or low y.
- high LD2 mainly high w.
- svd values show relative importance of LDs: LD1 much more important than LD2.

Group means by variable xxx

peanuts.1\$means

```
## y smk w
## 5_1 194.80 160.40 52.55
## 5_2 185.05 130.30 49.95
## 6_1 199.45 161.40 47.80
## 6_2 200.15 163.95 57.25
## 8_1 190.25 164.80 58.20
## 8 2 200.75 170.30 66.10
```

- 5_2 clearly smallest on y, smk, near smallest on w
- 8_2 clearly biggest on smk, w, also largest on y
- 8_1 large on w, small on y.

The predictions and misclassification

```
peanuts.pred <- predict(peanuts.1)
table(
  obs = peanuts.combo$combo,
  pred = peanuts.pred$class
)</pre>
```

```
##
      pred
## obs
       5_1 5_2 6_1 6_2 8_1 8_2
##
    5 1
             0
                    0
    5 2 0 2 0
##
                           0
    6 1 0
                    0
##
             0
                           0
    6_2 1
##
             0
##
    8_1 0
             0
                    0
                           0
    8 2
             0
##
```

Actually classified very well. Only one 6_2 classified as a 5_1, rest all correct.

Posterior probabilities xxx

```
pp <- round(peanuts.pred$posterior, 2)</pre>
peanuts.combo %>%
 select(-c(y, smk, w)) \%
 cbind(., pred = peanuts.pred$class, pp)
##
     obs combo pred 5 1 5 2 6 1 6 2 8 1 8 2
          5_1 5_1 0.69 0 0 0.31 0.00 0.00
## 1
## 2
       2 5 1 5 1 0.73 0 0 0.27 0.00 0.00
## 3
       3 5_2 5_2 0.00 1 0 0.00 0.00 0.00
## 4
      4 5 2 5 2 0.00 1 0 0.00 0.00 0.00
       5 6 1 6 1 0.00
                         0 1 0.00 0.00 0.00
## 5
## 6
      6 6_1 6_1 0.00 0 1 0.00 0.00 0.00
      7 6 2 6 2 0.13
                         0 0 0.87 0.00 0.00
## 7
       8 6 2 5 1 0.53
                         0 0 0.47 0.00 0.00
## 8
                         0 0 0.02 0.75 0.21
## 9
      9 8 1 8 1 0.02
      10 8 1 8 1 0.00
## 10
                         0 0 0.00 0.99 0.01
## 11
      11 8 2 8 2 0.00
                         0 0 0.00 0.03 0.97
## 12
      12
          8 2 8 2 0.00
                            0 0.00 0.06 0.94
```

Some doubt about which combo each plant belongs in, but not too much. The one misclassified plant was a close call.

Discriminant scores, again

- How are discriminant scores related to original variables?
- Construct data frame with original data and discriminant scores side by side: xxx

- xxx LD1 positive if w large and/or y small.
- LD2 positive if w large.

peanuts.1\$scaling

Discriminant scores for data

```
{\tt mm}
```

```
##
     combo
             v smk w LD1 LD2 LD3
## 1
       5_1 195.3 153.1 51.4 -1.42 -1.01 0.26
       5 1 194.3 167.7 53.7 -2.20 0.38 -1.13
## 2
## 3
       5_2 189.7 139.5 55.5 5.56 -1.10 0.79
       5 2 180.4 121.1 44.4 6.06 -3.89 -0.05
## 4
       6 1 203.0 156.8 49.8 -6.08 -1.25 1.25
## 5
## 6
       6_1 195.9 166.0 45.8 -7.13 -1.07 -1.24
## 7
       6 2 202.7 166.1 60.4 -1.43 1.12 1.10
## 8
       6_2 197.6 161.8 54.1 -2.28 -0.05 0.08
       8_1 193.5 164.5 57.8 1.05 0.86 -0.67
## 9
     8 1 187.0 165.1 58.6 4.02 1.22 -1.90
## 10
     8_2 201.5 166.8 65.0 1.60 1.95 1.15
## 11
       8 2 200.0 173.8 67.2 2.27 2.83 0.37
## 12
```

- Obs. 5 and 6 have most negative LD1: large y, small w.
- Obs. 4 has most negative LD2: small w.

Predict typical LD1 scores

First and third quartiles for three response variables:

```
quartiles <- peanuts %>%
  select(y:w) %>%
  map df(quantile, c(0.25, 0.75))
quartiles
## # A tibble: 2 x 3
##
             \mathtt{smk}
##
   <dbl> <dbl> <dbl>
## 1 193. 156. 51
## 2 200. 166. 59.0
new <- with(quartiles, crossing(y, smk, w))</pre>
```

The combinations

```
new
```

```
# A tibble: 8 x 3
##
            smk
##
    <dbl> <dbl> <dbl>
    193. 156.
## 1
                51
## 2
    193. 156. 59.0
## 3 193. 166. 51
    193. 166. 59.0
## 4
    200.
           156.
                 51
## 5
     200.
           156.
                 59.0
## 6
    200. 166.
                 51
## 7
## 8
     200.
           166.
                 59.0
```

pp <- predict(peanuts.1, new)</pre>

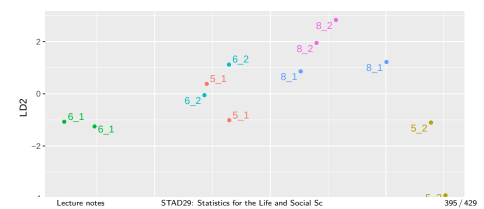
Predicted typical LD1 scores

```
cbind(new, pp$x) %>% arrange(LD1)
```

```
## y smk w LD1 LD2 LD3
## 1 200.375 166.275 51.00 -5.9688625 -0.3330095 -0.04523828
## 2 200.375 155.875 51.00 -4.1723048 -1.0396138 0.93093630
## 3 192.550 166.275 51.00 -2.8174566 -0.1007728 -1.51940856
## 4 200.375 166.275 59.05 -1.3059358 0.9791583 0.54572212
## 5 192.550 155.875 51.00 -1.0208989 -0.8073770 -0.54323399
## 6 200.375 155.875 59.05 0.4906219 0.2725540 1.52189670
## 7 192.550 166.275 59.05 1.8454701 1.2113950 -0.92844817
## 8 192.550 155.875 59.05 3.6420278 0.5047907 0.04772641
```

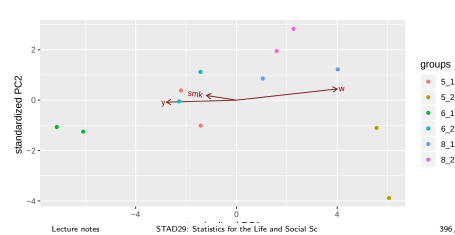
- Very negative LD1 score with large y and small w
- smk doesn't contribute much to LD1
- Very positive LD1 score with small y and large w.
- Same as we saw from Coefficients of Linear Discriminants.

Plot LD1 vs. LD2, labelling by combo



"Bi-plot" from ggbiplot xxx

```
ggbiplot(peanuts.1,
  groups = factor(peanuts.combo$combo)
)
```



Installing ggbiplot

- ggbiplot not on CRAN, so usual install.packages will not work.
- Install package devtools first (once):

```
install.packages("devtools")
```

Then install ggbiplot (once):

```
library(devtools)
install_github("vqv/ggbiplot")
```

Cross-validation

- So far, have predicted group membership from same data used to form the groups — dishonest!
- Better: *cross-validation*: form groups from all observations *except one*, then predict group membership for that left-out observation.
- No longer cheating!
- Illustrate with peanuts data again.

Misclassifications

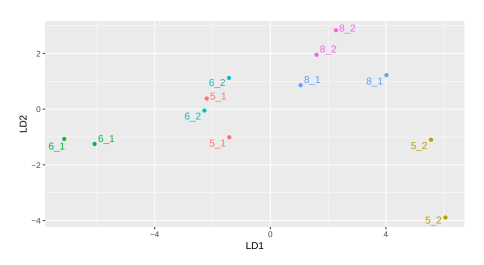
Fitting and prediction all in one go: xxx text under

```
peanuts.cv <- lda(combo ~ y + smk + w,
   data = peanuts.combo, CV = T
)
table(
  obs = peanuts.combo$combo,
  pred = peanuts.cv$class
)</pre>
```

```
##
      pred
       5 1 5 2 6 1 6 2 8 1 8 2
## obs
##
   5_1
        0
##
   5 2 0 1 0 0 1
   6 1 0
##
   6 2 1
##
##
   8 1
        0 1 0 0
##
   8 2
        0
```

Repeat of LD plot xxx

g



Posterior probabilities

data.frame(

pp <- round(peanuts.cv\$posterior, 3)</pre>

```
obs = peanuts.combo$combo,
 pred = peanuts.cv$class, pp
##
     obs pred X5_1 X5_2 X6_1 X6_2 X8_1 X8_2
     5 1 6 2 0.162 0.00 0.000 0.838 0.000 0.000
## 1
## 2 5_1 6_2 0.200 0.00 0.000 0.799 0.000 0.000
## 3 5 2 8 1 0.000 0.18 0.000 0.000 0.820 0.000
## 4 5 2 5 2 0.000 1.00 0.000 0.000 0.000 0.000
## 5 6_1 6_1 0.194 0.00 0.669 0.137 0.000 0.000
## 6 6 1 6 1 0.000 0.00 1.000 0.000 0.000 0.000
## 7 6_2 6_2 0.325 0.00 0.000 0.667 0.001 0.008
## 8 6 2 5 1 0.821 0.00 0.000 0.179 0.000 0.000
## 9 8 1 8 2 0.000 0.00 0.000 0.000 0.000 1.000
## 10 8_1 5_2 0.000 1.00 0.000 0.000 0.000 0.000
## 11 8 2 8 2 0.001 0.00 0.000 0.004 0.083 0.913
```

12 8 2 8 2 0.000 0.00 0.000 0.000 0.167 0.833

Why more misclassification?

- When predicting group membership for one observation, only uses the *other one* in that group.
- So if two in a pair are far apart, or if two groups overlap, great potential for misclassification.
- Groups 5_1 and 6_2 overlap.
- 5_2 closest to 8_1s looks more like an 8_1 than a 5_2 (other one far away).
- 8_1s relatively far apart and close to other things, so one appears to be a 5_2 and the other an 8_2.

Example 3: professions and leisure activities

- 15 individuals from three different professions (politicians, administrators and belly dancers) each participate in four different leisure activities: reading, dancing, TV watching and skiing. After each activity they rate it on a 0–10 scale.
- Some of the data: xxx

```
bellydancer 7 10 6 5
bellydancer 8 9 5 7
bellydancer 5 10 5 8
politician 5 5 5 6
politician 4 5 6 5
admin 4 2 2 5
admin 7 1 2 4
admin 6 3 3 3
```

• How can we best use the scores on the activities to predict a person's profession?

Discriminant analysis xxx

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/profile.txt"
active <- read_delim(my_url, " ")
active.1 <- lda(job ~ reading + dance + tv + ski, data = active)
active.1$svd</pre>
```

```
## [1] 9.856638 3.434555
```

```
active.1$scaling
```

```
## LD1 LD2

## reading -0.01297465 0.4748081

## dance -0.95212396 0.4614976

## tv -0.47417264 -1.2446327

## ski 0.04153684 0.2033122
```

- Two discriminants, first fair bit more important than second.
- LD1 depends (negatively) most on dance, a bit on tv. xxx

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LD2 depends mostly on tv.

5

Misclassification

admin

##

##

##

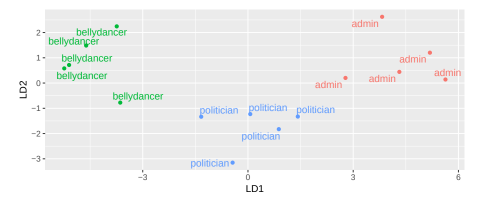
```
active.pred <- predict(active.1)
table(obs = active$job, pred = active.pred$class)

## pred
## obs admin bellydancer politician</pre>
```

Everyone correctly classified.

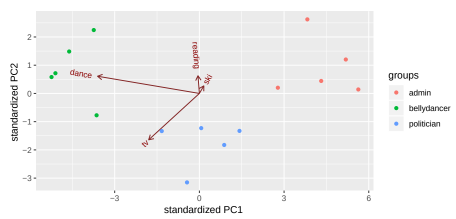
bellydancer politician

Plotting LDs xxx



Biplot xxx

ggbiplot(active.1, groups = active\$job)

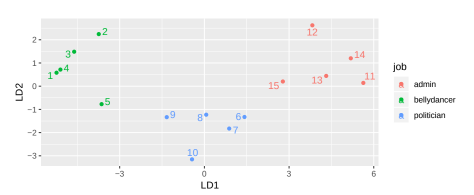


Comments on plot

- Groups well separated: bellydancers top left, administrators top right, politicians lower middle.
- Bellydancers most negative on LD1: like dancing most.
- Administrators most positive on LD1: like dancing least.
- Politicians most negative on LD2: like TV-watching most.

Plotting individual persons xxx

Make label be identifier of person. Now need legend:



Posterior probabilities

```
pp <- round(active.pred$posterior, 3)
data.frame(obs = active$job, pred = active.pred$class, pp)</pre>
```

```
##
              obs
                         pred admin bellydancer politician
## 1
      bellydancer bellydancer 0.000
                                           1.000
                                                       0.000
## 2
      bellydancer bellydancer 0.000
                                           1,000
                                                       0.000
## 3
      bellydancer bellydancer 0.000
                                           1.000
                                                       0.000
## 4
      bellvdancer bellvdancer 0.000
                                                       0.000
                                           1.000
## 5
      bellydancer bellydancer 0.000
                                                       0.003
                                           0.997
       politician politician 0.003
## 6
                                           0.000
                                                       0.997
       politician politician 0.000
                                           0.000
                                                       1.000
## 7
## 8
       politician politician 0.000
                                           0.000
                                                       1,000
## 9
       politician politician 0.000
                                           0.002
                                                       0.998
## 10
       politician politician 0.000
                                           0.000
                                                       1,000
## 11
            admin
                        admin 1.000
                                           0.000
                                                       0.000
## 12
            admin
                        admin 1.000
                                           0.000
                                                       0.000
## 13
            admin
                        admin 1.000
                                           0.000
                                                       0.000
## 14
            admin
                        admin 1.000
                                           0.000
                                                       0.000
## 15
                                           0.000
                                                       0.018
            admin
                       admin 0.982
```

Not much doubt.

Cross-validating the jobs-activities data

Recall: no need for predict. Just pull out class and make a table:

```
active.cv <- lda(job ~ reading + dance + tv + ski,
   data = active, CV = T
)
table(obs = active$job, pred = active.cv$class)</pre>
```

```
## pred
## obs admin bellydancer politician
## admin 5 0 0
## bellydancer 0 4 1
## politician 0 5
```

This time one of the bellydancers was classified as a politician.

and look at the posterior probabilities

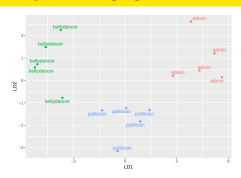
picking out the ones where things are not certain:

```
pp <- round(active.cv$posterior, 3)
data.frame(obs = active$job, pred = active.cv$class, pp) %>%
  mutate(max = pmax(admin, bellydancer, politician)) %>%
  filter(max < 0.9995)</pre>
```

```
##
                     pred admin bellydancer politician
  1 bellydancer politician 0.000
                                      0.001
                                                0.999 0.999
     politician politician 0.006
                                                0.994 0.994
##
                                      0.000
     politician politician 0.001
                                      0.000
                                                0.999 0.999
##
     politician politician 0.000
                                      0.009
                                                0.991 0.991
## 4
## 5
          admin
                     admin 0.819
                                      0.000
                                                0.181 0.819
```

- Bellydancer was "definitely" a politician!
- One of the administrators might have been a politician too.

Why did things get misclassified?



- * Go back to plot of discriminant scores:
- * one bellydancer much closer to the politicians,
- * one administrator a bit closer to the politicians.

Example 4: remote-sensing data xxx from here

- View 38 crops from air, measure 4 variables x1-x4.
- Go back and record what each crop was.
- Can we use the 4 variables to distinguish crops?

Reading in

```
my_url <- "http://www.utsc.utoronto.ca/~butler/d29/remote-sens
crops <- read_table(my_url)</pre>
## Parsed with column specification:
## cols(
     crop = col_character(),
##
##
     x1 = col double(),
     x2 = col double(),
##
     x3 = col double(),
##
##
     x4 = col double(),
```

cr = col character()

##

)

Starting off: number of LDs

```
crops.lda <- lda(crop ~ x1 + x2 + x3 + x4, data = crops)
crops.lda$svd</pre>
```

```
## [1] 2.2858251 1.1866352 0.6394041 0.2303634
```

- 4 LDs (four variables, six groups).
- 1st one important, maybe 2nd as well.

Connecting original variables and LDs

crops.lda\$means

```
## x1 x2 x3 x4
## Clover 46.36364 32.63636 34.18182 36.63636
## Corn 15.28571 22.71429 27.42857 33.14286
## Cotton 34.50000 32.66667 35.00000 39.16667
## Soybeans 21.00000 27.00000 23.50000 29.66667
## Sugarbeets 31.00000 32.16667 20.00000 40.50000
```

```
round(crops.lda$scaling, 3)
```

```
## LD1 LD2 LD3 LD4

## x1 -0.061 0.009 -0.030 -0.015

## x2 -0.025 0.043 0.046 0.055

## x3 0.016 -0.079 0.020 0.009

## x4 0.000 -0.014 0.054 -0.026
```

links grouns to original variables to I Ds Lecture notes STAD29: Statistics for the Life and Social Sc

LD1 and texttt{LD2}

round(crops.lda\$scaling, 3)

```
## LD1 LD2 LD3 LD4

## x1 -0.061 0.009 -0.030 -0.015

## x2 -0.025 0.043 0.046 0.055

## x3 0.016 -0.079 0.020 0.009

## x4 0.000 -0.014 0.054 -0.026

$
```

- LD1 mostly x1 (minus), so clover low on LD1, corn high.
- LD2 x3 (minus), x2 (plus), so sugarbeets should be high on LD2.

Predictions

Thus:

```
crops.pred <- predict(crops.lda)
table(obs = crops$crop, pred = crops.pred$class)</pre>
```

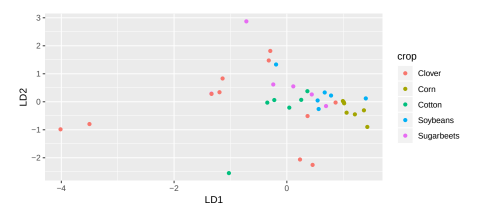
```
##
               pred
## obs
                Clover Corn Cotton Soybeans Sugarbeets
     Clover
##
                      6
                                                        2
##
     Corn
##
     Cotton
    Soybeans
##
##
     Sugarbeets
```

- Not very good, eg. only 6 of 11 Clover classified correctly.
- Set up for plot:

```
mm <- data.frame(crop = crops$crop, crops.pred$x)
```

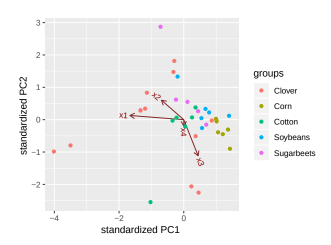
Plotting the LDs

```
ggplot(mm, aes(x = LD1, y = LD2, colour = crop)) +
  geom_point()
```



Biplot

ggbiplot(crops.lda, groups = crops\$crop)



Try removing Clover

• the dplyr way:

```
crops %>% filter(crop != "Clover") -> crops2
crops2.lda <- lda(crop ~ x1 + x2 + x3 + x4, data = crops2)</pre>
```

- LDs for crops2 will be different from before.
- Concentrate on plot and posterior probs.

```
crops2.pred <- predict(crops2.lda)
mm <- data.frame(crop = crops2$crop, crops2.pred$x)</pre>
```

lda output

Different from before:

```
crops2.lda$means
```

```
## x1 x2 x3 x4
## Corn 15.28571 22.71429 27.42857 33.14286
## Cotton 34.50000 32.66667 35.00000 39.16667
## Soybeans 21.00000 27.00000 23.50000 29.66667
## Sugarbeets 31.00000 32.16667 20.00000 40.50000
crops2.lda$svd
```

[1] 3.3639389 1.6054750 0.4180292

crops2.lda\$scaling

```
## LD1 LD2 LD3

## x1 0.14077479 0.007780184 -0.0312610362

## x2 0.03006972 0.007318386 0.0085401510

## x3 -0.06363974 -0.099520895 -0.0005309869

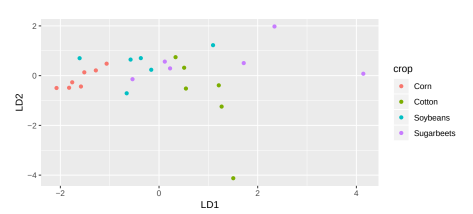
## x4 -0.00677414 -0.035612707 0.0577718649
```

STAD29: Statistics for the Life and Social Sc.

Plot

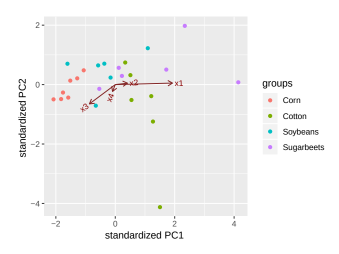
A bit more clustered:

```
ggplot(mm, aes(x = LD1, y = LD2, colour = crop)) +
geom_point()
```



Biplot

ggbiplot(crops2.lda, groups = crops2\$crop)



Quality of classification

```
table(obs = crops2$crop, pred = crops2.pred$class)
```

```
## obs Corn Cotton Soybeans Sugarbeets
## Corn 6 0 1 0
## Cotton 0 4 2 0
## Soybeans 2 0 3 1
## Sugarbeets 0 0 3 3
```

Better.

Posterior probs, the wrong ones

obs

```
\1 xxx {
post <- round(crops2.pred$posterior, 3)</pre>
data.frame(obs = crops2$crop, pred = crops2.pred$class, post) %>%
  filter(obs != pred)
```

```
pred Corn Cotton Soybeans Sugarbeets
## 1
          Corn
                Soybeans 0.443 0.034
                                       0.494
                                                 0.029
      Soybeans Sugarbeets 0.010
                                       0.299
## 2
                               0.107
                                                 0.584
##
      Soybeans
                    Corn 0.684 0.009
                                       0.296
                                                 0.011
## 4
      Soybeans
                    Corn 0.467 0.199
                                       0.287
                                                 0.047
## 5
        Cotton
                Soybeans 0.056 0.241
                                       0.379
                                                 0.324
##
        Cotton
                Soybeans 0.066 0.138
                                       0.489
                                                 0.306
                Soybeans 0.381 0.146
                                       0.395
                                                 0.078
##
    Sugarbeets
                Soybeans 0.106 0.144
    Sugarbeets
                                       0.518
                                                 0.232
  9 Sugarbeets
                Soybeans 0.088
                               0.207
                                       0.489
                                                 0.216
```

 These were the misclassified ones, but the posterior probability of being correct was not usually too low.

##

MANOVA

##

Began discriminant analysis as a followup to MANOVA. Do our variables significantly separate the crops (excluding Clover)?

```
response <- with(crops2, cbind(x1, x2, x3, x4))
crops2.manova <- manova(response ~ crop, data = crops2)
summary(crops2.manova)</pre>
```

Df Pillai approx F num Df den Df Pr(>F)

```
## crop 3 0.9113 2.1815 12 60 0.02416 *
## Residuals 21
## ---
## Signif. codes:
## 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Yes, at least one of the crops differs (in means) from the others. So it is worth doing this analysis.

We did this the wrong way around, though!

The right way around

- First, do a MANOVA to see whether any of the groups differ significantly on any of the variables.
- If the MANOVA is significant, do a discriminant analysis in the hopes of understanding how the groups are different.
- For remote-sensing data (without Clover):
- LD1 a fair bit more important than LD2 (definitely ignore LD3).
- LD1 depends mostly on x1, on which Cotton was high and Corn was low.
- Discriminant analysis in MANOVA plays the same kind of role that Tukey does in ANOVA.
- ## Error in FUN(X[[i]], ...): invalid 'name' argument