# Regression revisited

# 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, exclude = "select") # for Box-Cox, later
library(tidyverse)
library(broom)
library(marginaleffects)
```

## 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://ritsokiguess.site/datafiles/sleep.txt"
sleep <- read_delim(my_url, " ")</pre>
```

#### Check data

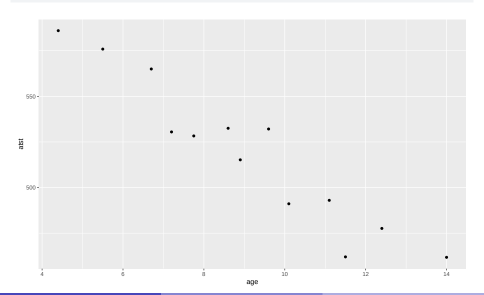
#### sleep

```
A tibble: 13 x 2
   atst
          age
  <dbl> <dbl>
   586
         4.4
   462. 14
   491. 10.1
   565
       6.7
   462 11.5
   532. 9.6
   478. 12.4
   515. 8.9
   493 11.1
10
   528. 7.75
11
   576. 5.5
12
   532. 8.6
13
   530. 7.2
```

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

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# 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)

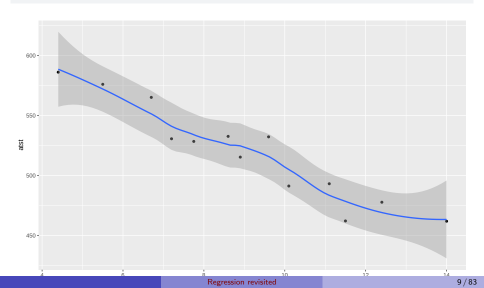
```
atst age
atst 1.0000000 -0.9515469
age -0.9515469 1.0000000
```

#### 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()
```



# 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 ***
      -14.041 1.368 -10.26 5.70e-07 ***
age
Signif. codes:
  '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Regression revisited

#### 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)

# also one-line summary of model:

```
glance(sleep.1)
```

### Broom part 2

#### sleep.1 %>% augment(sleep)

```
# A tibble: 13 x 8
   atst
         age .fitted .resid .hat .sigma .cooksd
  <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
   586
      4.4
               585. 1.30 0.312 13.8 0.00320
   462. 14 450. 11.8 0.341
                                 13.0 0.319
   491. 10.1 505. -13.6 0.0887
                                 13.0 0.0568
   565 6.7
            552. 12.6 0.137
                                 13.1 0.0844
   462 11.5
            485. -23.0 0.141
                                 11.3 0.294
   532. 9.6
               512. 20.4 0.0801
                                 12.0 0.114
   478. 12.4
               472. 5.23 0.198
                                 13.7 0.0243
8 515. 8.9
               522. -6.32 0.0772
                                 13.6 0.0105
9
   493 11.1
            491. 2.37 0.122
                                 13.8 0.00258
10
   528. 7.75
               538. -9.37 0.0954
                                 13.4 0.0296
11
   576. 5.5 569. 6.64 0.214
                                 13.6 0.0441
   532. 8.6
12
            526. 6.77 0.0792
                                 13.6 0.0124
13 530. 7.2
               545. -14.9 0.114
                                  12.9 0.0933
# i 1 more variable: .std.resid <dbl>
```

Useful for plotting residuals against an x-variable.

# 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.

# The marginal effects package 1/2

To get predictions for specific values, set up a dataframe with those values first:

```
new <- datagrid(model = sleep.1, age = c(10, 5))
new</pre>
```

```
age rowid
1 10 1
2 5 2
```

Any variables in the dataframe that you don't specify are set to their mean values (quantitative) or most common category (categorical).

# The marginal effects package 2/2

Then feed into newdata in predictions. This contains a lot of columns, so you probably want only to display the ones you care about:

```
cbind(predictions(sleep.1, newdata = new)) %>%
select(estimate, conf.low, conf.high, age)
```

```
estimate conf.low conf.high age
1 506.0729 498.4899 513.6558 10
2 576.2781 563.2588 589.2974 5
```

The confidence limits are a 95% confidence interval for the mean response at that age.

#### Prediction intervals

These are obtained (instead) with predict as below. Use the same dataframe new as before:

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

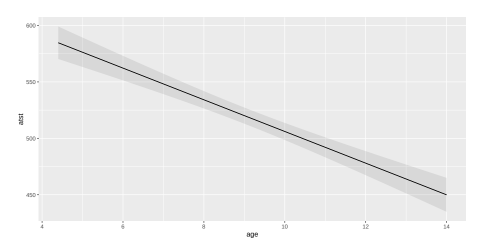
```
fit lwr upr
1 506.0729 475.8982 536.2475
2 576.2781 543.8474 608.7088
```

```
cbind(new, pp)
```

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

# Plotting the confidence intervals for mean response again:

```
plot_predictions(sleep.1, condition = "age")
```



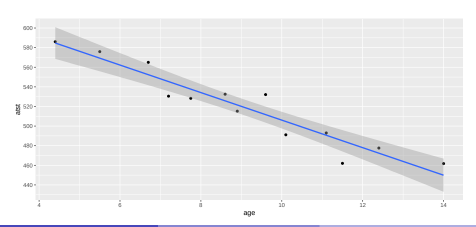
#### Comments

- 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

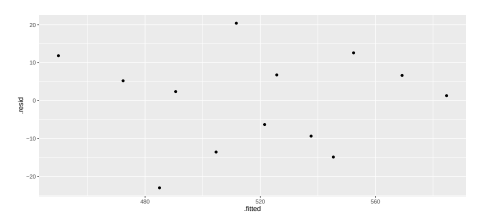
How to tell whether a straight-line regression is appropriate?

- 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.

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

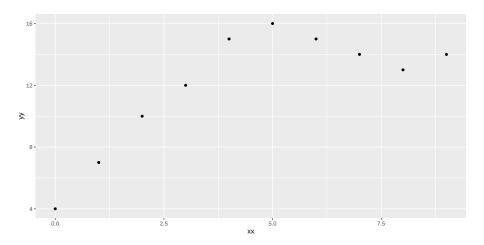


# An inappropriate regression

#### Different data:

```
my_url <- "http://ritsokiguess.site/datafiles/curvy.txt"
curvy <- read_delim(my_url, " ")</pre>
```

# Scatterplot



## Regression line, anyway

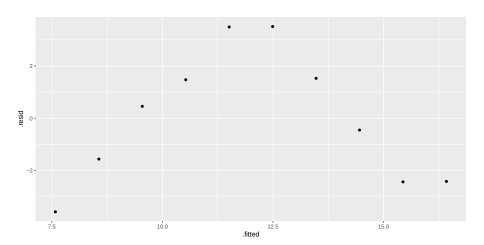
```
curvy.1 <- lm(yy ~ xx, data = curvy)
summary(curvy.1)</pre>
```

```
Call:
lm(formula = yy ~ xx, data = curvy)
Residuals:
  Min 10 Median 30 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 **
       0.9818 0.2925 3.356 0.00998 **
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
```

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# 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 \leftarrow lm(yy \sim xx + I(xx^2), data = curvy)
```

- 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))
```

### Regression 2

summary(curvy.2)

```
Call:
lm(formula = yy \sim xx + I(xx^2), data = curvy)
Residuals:
                                  Max
   Min 10 Median 30
-1.2091 -0.3602 -0.2364 0.8023 1.2636
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 3.90000 0.77312 5.045 0.001489 **
          3.74318    0.40006    9.357    3.31e-05 ***
хx
I(xx^2) -0.30682 0.04279 -7.170 0.000182 ***
Signif. codes:
0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.9833 on 7 degrees of freedom
Multiple R-squared: 0.9502, Adjusted R-squared:
```

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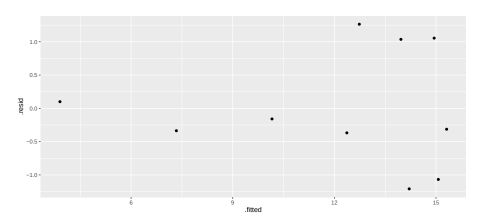
#### Comments

- 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 apparent 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  $\ln y = \ln a + bx$ .
- 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)

Regression revisited

## Some made-up data

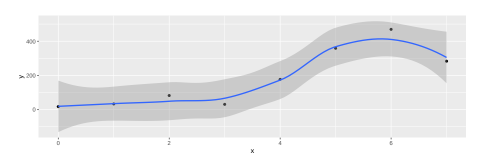
```
my_url <- "http://ritsokiguess.site/datafiles/madeup2.csv"
madeup <- read_csv(my_url)
madeup</pre>
```

```
# A tibble: 8 x 3
   ...1 x y
 <dbl> <dbl> <dbl>
           0 17.9
     2 1 33.6
3
          2 82.7
4
       3 31.2
5
          4 177.
6
           5 359.
           6 469.
8
           7 283.
```

Seems to be faster-than-linear growth, maybe exponential growth.

# Scatterplot: faster than linear growth

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

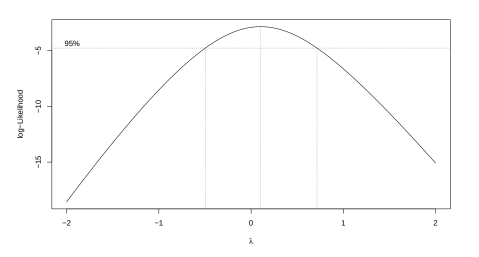


# Running Box-Cox

- 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

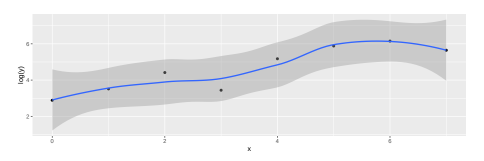
- $\lambda$  (lambda) is the power by which you should transform y to get the relationship straight (straighter). Power 0 is "take logs"
- Middle dotted line marks best single value of  $\lambda$  (here about 0.1).
- Outer dotted lines mark 95% CI for  $\lambda$ , here -0.3 to 0.7, approx. (Rather uncertain 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  $\lambda$  like 2,1,0.5,0,-0.5,-1. Here 0 and 0.5 good values to pick.

Regression revisited

# Did transformation straighten things?

ullet Plot transformed y against x. Here, log:

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



Looks much straighter.

# Regression with transformed y

```
madeup.1 \leftarrow lm(log(y) \sim x, data = madeup)
summary(madeup.1)
```

```
Call:
lm(formula = log(y) \sim x, data = madeup)
Residuals:
   Min 1Q Median 3Q Max
-0.9688 -0.2577 0.1663 0.3881 0.5534
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 3.02884 0.37935 7.984 0.000206 ***
x
 0.46006 0.09068 5.073 0.002281 **
Signif. codes:
0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.5877 on 6 degrees of freedom
```

Multiple R-squared: 0.811, Adjusted R-squared: 0.7794

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# 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://ritsokiguess.site/datafiles/regressx.txt"
visits <- read_delim(my_url, " ")</pre>
```

#### Check data

#### visits

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

## Fit multiple regression

visits.1 <- lm(timedrs ~ phyheal + menheal + stress,</pre>

```
data = visits)
summary(visits.1)
Call:
lm(formula = timedrs ~ phyheal + menheal + stress, data = visits)
Residuals:
   Min 1Q Median
                          30
                                 Max
-14.792 -4.353 -1.815 0.902 65.886
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) -3.704848 1.124195 -3.296 0.001058 **
phyheal 1.786948 0.221074 8.083 5.6e-15 ***
menheal -0.009666 0.129029 -0.075 0.940318
stress 0.013615 0.003612 3.769 0.000185 ***
Signif. codes:
0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 9.708 on 461 degrees of freedom
```

Multiple R-squared: 0.2188. Adjusted R-squared: 0.2137

Regression revisited

E-atatiatic: 42 03 on 3 and 461 DE newsluce

# 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.
- The physical health and stress variables definitely 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:

#### Just menheal

```
summary(visits.2)
Call:
lm(formula = timedrs ~ menheal, data = visits)
Residuals:
   Min 1Q Median 3Q Max
-13.826 -5.150 -2.818 1.177 72.513
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 3.8159 0.8702 4.385 1.44e-05 ***
menheal 0.6672 0.1173 5.688 2.28e-08 ***
Signif. codes:
0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 10.6 on 463 degrees of freedom
```

Multiple R-squared: 0.06532, Adjusted R-squared: 0.0633

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

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## 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):

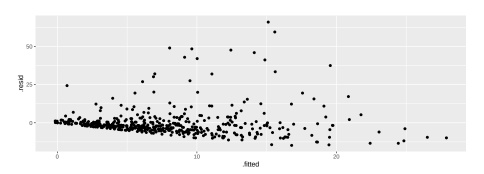
```
visits %>% select(-subjno) %>% cor()
```

```
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.

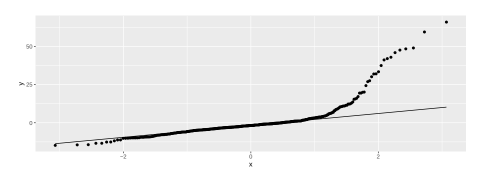
# Residual plot (from timedrs on all)

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



Apparently random. But...

# Normal quantile plot of residuals

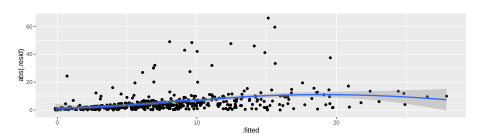


Not normal at all; upper tail is way too long.

### Absolute residuals

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

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



#### Comments

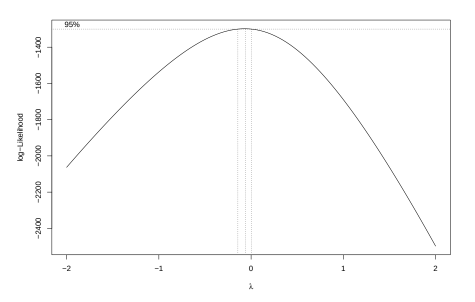
- On the normal quantile plot:
  - highest (most positive) residuals are way too high
  - distribution of residuals skewed to right (not normal at all)
- On plot of absolute residuals:
  - size of residuals getting bigger as fitted values increase
  - predictions getting more variable as fitted values 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:  $\lambda$  just less than zero.
- Hard to see scale.
- Focus on  $\lambda$  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

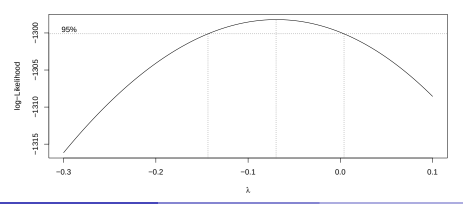
[19] -0.12 -0.11 -0.10 -0.09 -0.08 -0.07 -0.06 -0.05 -0.04

[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:  $\lambda$  just about -0.07.
- CI for  $\lambda$  about (-0.14, 0.01).
- ullet 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)

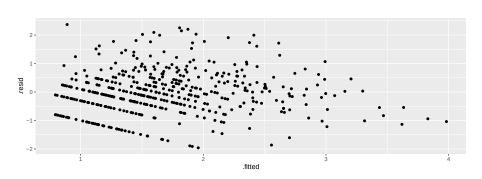
```
Call:
lm(formula = log(timedrs + 1) ~ phyheal + menheal + stress, data = visits)
Residuals:
    Min
         10 Median 30
                                      Max
-1.95865 -0.44076 -0.02331 0.42304 2.36797
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.3903862 0.0882908 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 '**' 0.01 '*' 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?

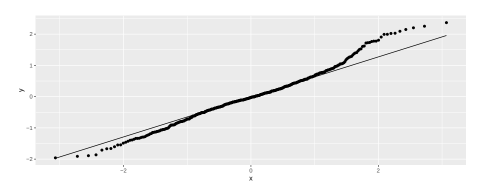
# Residuals against fitted values

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



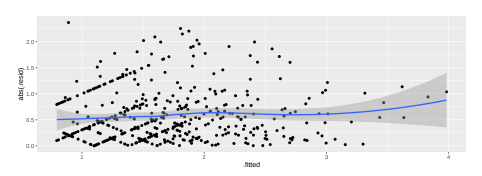
# Normal quantile plot of residuals

```
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:
0 '***' 0.001 '**' 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".
- ullet Taking out those x's is a mistake. Or putting them in is a good idea.

# 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.

# Reading in

• Separated by multiple spaces with columns lined up:

```
my_url <- "http://ritsokiguess.site/datafiles/punting.txt"
punting <- read_table(my_url)</pre>
```

#### The data

#### punting

```
A tibble: 13 x 4
    left right punt
                        fred
   <dbl> <dbl> <dbl> <dbl>
     170
            170
                  162.
                          171
1
2
     130
            140
                  144
                          136
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)
summary(punting.1)</pre>
```

#### Call:

```
lm(formula = punt ~ left + right + fred, data = punting)
```

#### Residuals:

```
Min 1Q Median 3Q Max -14.9325 -11.5618 -0.0315 9.0415 20.0886
```

#### Coefficients:

|             | Estimate | Std. Error | t value | Pr(> t ) |
|-------------|----------|------------|---------|----------|
| (Intercept) | -4.6855  | 29.1172    | -0.161  | 0.876    |
| left        | 0.2679   | 2.1111     | 0.127   | 0.902    |
| right       | 1.0524   | 2.1477     | 0.490   | 0.636    |
| fred        | -0.2672  | 4.2266     | -0.063  | 0.951    |

Residual standard error: 14.68 on 9 degrees of freedom Multiple R-squared: 0.7781, Adjusted R-squared: 0.7042 F-statistic: 10.52 on 3 and 9 DF, p-value: 0.00267

### 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>
```

Analysis of Variance Table

```
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
```

No significant loss by dropping other two variables.

# Comparing R-squareds

• All three *x*-variables:

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

[1] 0.7781401

• Only right:

```
summary(punting.2)$r.squared
```

[1] 0.7753629

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

### Regression results

summary(punting.2)

```
Call:
lm(formula = punt ~ right, data = punting)
Residuals:
    Min 1Q Median 3Q
                                     Max
-15.7576 -11.0611 0.3656 7.8890 19.0423
Coefficients:
          Estimate Std. Error t value Pr(>|t|)
(Intercept) -3.6930 25.2649 -0.146
                                       0.886
right 1.0427 0.1692 6.162 7.09e-05 ***
Signif. codes:
0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 13.36 on 11 degrees of freedom
Multiple R-squared: 0.7754, Adjusted R-squared: 0.7549
F-statistic: 37.97 on 1 and 11 DF, p-value: 7.088e-05
```

Regression revisited 77 / 83

### 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.

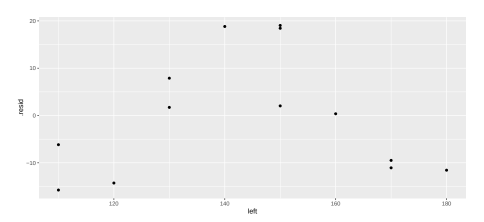
## Augmenting punting.2

```
punting.2 %>% augment(punting) -> punting.2.aug
punting.2.aug
```

```
# A tibble: 13 \times 10
   left right punt
                   fred .fitted .resid .hat .sigma
  <dbl> <dbl> <dbl> <dbl> <
                          <dbl>
                                 <dbl>
                                       <dbl>
                                              <dbl>
    170
          170
1
              162.
                    171 174. -11.1
                                       0.157
                                               13.5
2
    130
         140
              144
                    136
                           142.
                                 1.72
                                       0.0864
                                               14.0
3
    170
         180
              174.
                    174
                           184. -9.49
                                       0.244
                                               13.6
4
    160
         160
              164.
                    161
                           163. 0.366 0.101
                                               14.0
5
    150
         170
              192
                    159
                           174. 18.4
                                       0.157
                                               12.5
6
    150
         150
              172.
                    151
                           153. 19.0
                                       0.0778
                                               12.5
7
    180
         170
              162
                    174
                           174. -11.6
                                       0.157
                                               13.4
8
    110
         110
              105.
                    111
                           111. -6.17
                                       0.305
                                               13.8
9
    110
         120
              106.
                    114
                           121. -15.8
                                       0.2
                                               12.9
10
    120
         130
              118.
                    126
                           132. -14.3
                                       0.127
                                               13.1
11
    140
         120
              140.
                    129
                           121. 18.8
                                       0.2
                                               12.3
12
    130
          140
              150.
                    136
                           142. 7.89
                                       0.0864
                                               13.8
13
    150
          160
              165.
                    154
                           163.
                                 2.04
                                       0.101
                                               14.0
   2 more variables: .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.