Lesson 19: ACFT Problem (solution)

Is there an association between the Standing Power Throw and Maximum Deadlift raw scores?

We are going to determine if an association exists between spr raw and mdl raw scores.

The relationship we are modeling looks like this:

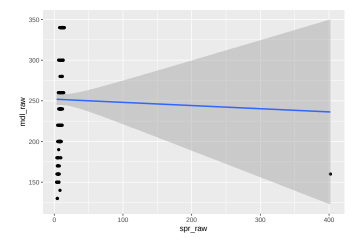
$$\widehat{mdl} = \beta_0 + \beta_1 \cdot spr$$

1. Provide the hypotheses in symbols and words:

 $H_0: \beta_1 = 0$. My null hypothesis is that there is no association between 'spr_raw' and 'mdl_raw' $H_A: \beta_1 \neq 0$. My null hypothesis is that there is an association between 'spr_raw' and 'mdl_raw'.

2. Plot the association, including the regression line.

```
df %>%
    ggplot(aes(x = spr_raw, y = mdl_raw)) +
    geom_point() +
    geom_smooth(method = "lm")
```



3. Is there anything unusual about the data?

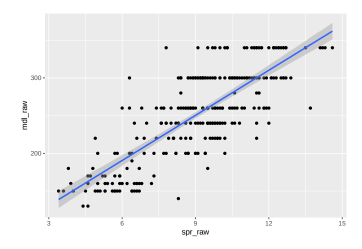
Yes. Outlier.

4. Remove the outlier using the filter() command. How many observations were dropped?

```
df <- df %>% filter(spr_raw < 100)</pre>
```

5. Create a new scatter plot with updated dataframe.

```
df %>%
   ggplot(aes(x = spr_raw, y = mdl_raw)) +
   geom_point() +
   geom_smooth(method = "lm")
```



6. Create a linear model to test your hypotheses.

```
lrmodel <- df %>%
  lm(mdl_raw ~ spr_raw, data = .)
summary(lrmodel)
```

```
##
## Call:
## lm(formula = mdl_raw ~ spr_raw, data = .)
##
## Residuals:
##
      Min
                1Q
                   Median
                                3Q
                                       Max
  -96.394 -24.950
                   -2.174
                           23.672 113.572
##
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 70.9488
                            8.7017
                                     8.153 1.08e-14 ***
                19.9332
                            0.9299 21.436 < 2e-16 ***
## spr_raw
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 36.03 on 290 degrees of freedom
## Multiple R-squared: 0.6131, Adjusted R-squared: 0.6117
## F-statistic: 459.5 on 1 and 290 DF, p-value: < 2.2e-16
```

7. Write out the linear model with the coefficients determined by the linear regression.

```
\widehat{mdl} = 70.95 + 19.93 \cdot spr
```

8. Interpret the intercept term.

With a 'spr_raw' of 0, we can expect an 'mdl_raw' of 70.95lbs. This does not make sense because we are extrapolating outside of our data.

9. Interpret the slope term.

On average, every one unit (meter) increase in Standing Power Throw is associated with an increase of Maximum Deadlift by 19.93lbs, holding all else equal.

10. Conclusion of your hypothesis test at a 5% significance level.

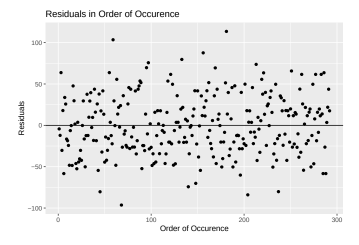
I reject my null at the 5% significance level that there is no association between 'spr_raw' and 'mdl_raw'.

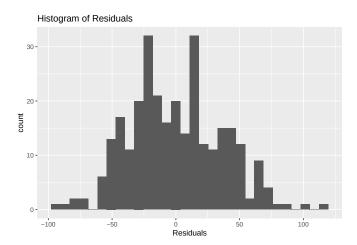
11. Interpret the R-squared value.

About 61% of the variation in 'mdl_raw' is explained by 'spr_raw'.

12. Check the 4 Validity Conditions (L.I.N.E.). Are any not met?

Residuals vs. Predicted Values 100 - 50 - 150 - 200 250 300 350 Predicted Values





While there are "bands" in this data, that is due to the limited number of 'mdl_raw' values. Overall, the plot appears to be somewhat linear but variance is not constant! There does not appear to be independence issues and the histogram of the residuals seems fairly normal/bell-shaped.

13. Can we determine causation? Why or why not.

No, because this was an observational study and there was no random assignment.

Does adding the variable sex change the association between the Standing Power Throw (spr_raw) and Leg Tuck (ltk_raw) raw scores?

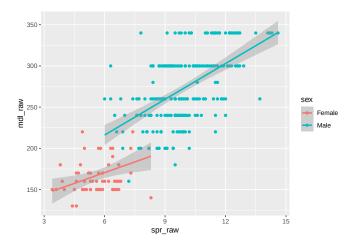
Now, we add sex as another variable to the model but also include it as an interaction with spr_raw.

Our updated model looks like this:

$$\widehat{mdl} = \beta_0 + \beta_1 \cdot spr + \beta_2 \cdot sexmale + \beta_3 (spr \cdot sexmale)$$

14. Plot the data (including the interaction term), including the regression line.

```
df %>%
  ggplot(aes(x = spr_raw, y = mdl_raw, color = sex)) +
  geom_point() +
  geom_smooth(method = "lm")
```



15. Does there appear to be a different relationship between mdl_raw and spr_raw based on sex?

Maybe. The lines don't have a very different slope.

16. Conduct a linear regression with the updated model.

```
lrmodel <- df %>%
  lm(mdl_raw ~ spr_raw * sex, data = .)
summary(lrmodel)
##
## Call:
## lm(formula = mdl_raw ~ spr_raw * sex, data = .)
##
##
  Residuals:
##
       Min
                 1Q
                     Median
                                  3Q
                                         Max
##
   -86.729 -24.267
                     -1.634
                             27.365
                                     97.850
##
```

Coefficients: ## Estimate Std. Error t value Pr(>|t|) 118.529 24.401 4.858 1.95e-06 *** ## (Intercept) ## spr_raw 8.662 4.207 2.059 0.0404 * 10.844 28.003 0.387 0.6989 ## sexMale ## spr_raw:sexMale 5.797 4.423 1.310 0.1911 ## ---## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1

##

Residual standard error: 33.84 on 288 degrees of freedom ## Multiple R-squared: 0.6611, Adjusted R-squared: 0.6576 ## F-statistic: 187.3 on 3 and 288 DF, p-value: < 2.2e-16

17. Interpret the coefficient on the interaction term.

The coefficient indicates that, for Males, a one-unit increase in 'spr_raw' results in an 8.66+5.797 increase in 'mdl_raw', on average, holding all else constant.

18. Is this significant at the 5% level?

No, because our p-value (.1911) exceeds our significance level.

19. Interpret the intercept term.

With a zero 'spr_raw', Females will exhibit a 'mdl_raw' of 119lbs.

20. Interpret the base slope term for spr_raw.

For Females, a one-unit increase in 'spr_raw' results in an 8.67 increase in 'mdl_raw', on average, holding all else equal.

- 21. Using this model (even with no significance), calculate the following:
- The mdl_raw for a male cadet with an spr_raw of 6m:

```
(118.529 + 10.84) + 6*(8.662 + 5.797)
```

```
## [1] 216.123
```

• The mdl_raw for a female cadet with an spr_raw of 6m:

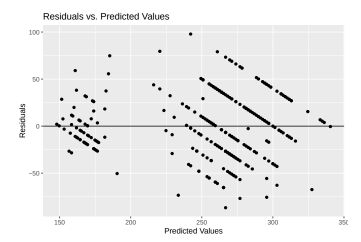
```
(118.529) + 6*(8.662)
```

```
## [1] 170.501
```

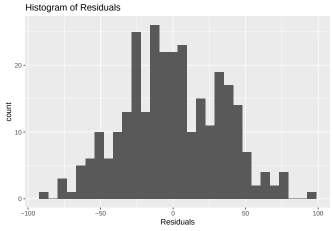
22. Interpret the R-squared value.

About 66% of the variation in 'mdl_raw' is explained by 'spr_raw' and sex and the interaction between the two.

23. Check the 4 Validity Conditions (L.I.N.E.). Are any not met?



Residuals in Order of Occurence



The plots look very similar to before. However, there are

two-groups of residuals now...