Introduction to linear regression

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The Human Freedom Index is a report that attempts to summarize the idea of "freedom" through a bunch of different variables for many countries around the globe. It serves as a rough objective measure for the relationships between the different types of freedom - whether it's political, religious, economical or personal freedom - and other social and economic circumstances. The Human Freedom Index is an annually copublished report by the Cato Institute, the Fraser Institute, and the Liberales Institut at the Friedrich Naumann Foundation for Freedom.

In this lab, you'll be analyzing data from Human Freedom Index reports from 2008-2016. Your aim will be to summarize a few of the relationships within the data both graphically and numerically in order to find which variables can help tell a story about freedom.

Getting Started

Load packages

In this lab, you will explore and visualize the data using the **tidyverse** suite of packages. The data can be found in the companion package for OpenIntro resources, **openintro**.

Let's load the packages.

```
library(tidyverse)
library(openintro)
data('hfi', package='openintro')
```

The data

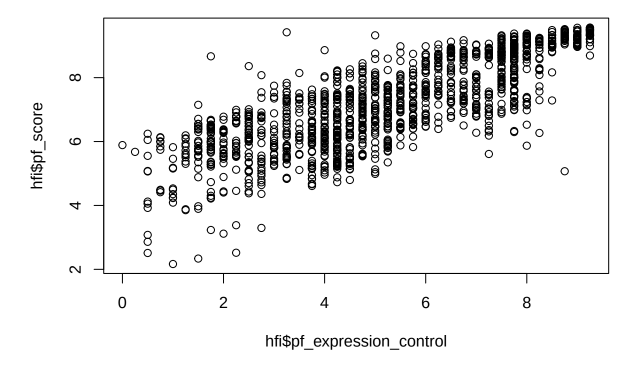
The data we're working with is in the openintro package and it's called hfi, short for Human Freedom Index.

1. What are the dimensions of the dataset? 1458rows x 123 columns

```
dim(hfi)
```

```
## [1] 1458 123
```

2. What type of plot would you use to display the relationship between the personal freedom score, pf_score, and one of the other numerical variables? Plot this relationship using the variable pf_expression_control as the predictor. Does the relationship look linear? If you knew a country's pf_expression_control, or its score out of 10, with 0 being the most, of political pressures and controls on media content, would you be comfortable using a linear model to predict the personal freedom score? A standard plot of two variables would be sufficient as this factor is numerical and could be pit against another numerical variable to recognize trends. No I would not be comfortable, we need to see the data through more lens to determine the progression forward.



If the relationship looks linear, we can quantify the strength of the relationship with the correlation coefficient.

Here, we set the use argument to "complete.obs" since there are some observations of NA.

Sum of squared residuals

In this section, you will use an interactive function to investigate what we mean by "sum of squared residuals". You will need to run this function in your console, not in your markdown document. Running the function also requires that the hfi dataset is loaded in your environment.

Think back to the way that we described the distribution of a single variable. Recall that we discussed characteristics such as center, spread, and shape. It's also useful to be able to describe the relationship of two numerical variables, such as pf_expression_control and pf_score above.

3. Looking at your plot from the previous exercise, describe the relationship between these two variables. Make sure to discuss the form, direction, and strength of the relationship as well as any unusual observations. The points have a strong correlation coefficient of 0.796 which suggests they strongly influence each other. Just as you've used the mean and standard deviation to summarize a single variable, you can summarize the relationship between these two variables by finding the line that best follows their association. Use the following interactive function to select the line that you think does the best job of going through the cloud of points.

```
# This will only work interactively (i.e. will not show in the knitted document)
hfi <- hfi %>% filter(complete.cases(pf_expression_control, pf_score))
DATA606::plot_ss(x = hfi$pf_expression_control, y = hfi$pf_score)
```

After running this command, you'll be prompted to click two points on the plot to define a line. Once you've done that, the line you specified will be shown in black and the residuals in blue. Note that there are 30 residuals, one for each of the 30 observations. Recall that the residuals are the difference between the observed values and the values predicted by the line:

$$e_i = y_i - \hat{y}_i$$

The most common way to do linear regression is to select the line that minimizes the sum of squared residuals. To visualize the squared residuals, you can rerun the plot command and add the argument **showSquares** = TRUE.

```
DATA606::plot_ss(x = hfi$pf_expression_control, y = hfi$pf_score, showSquares = TRUE)
```

Note that the output from the plot_ss function provides you with the slope and intercept of your line as well as the sum of squares.

4. Using plot_ss, choose a line that does a good job of minimizing the sum of squares. Run the function several times. What was the smallest sum of squares that you got? How does it compare to your neighbors? 997 was the lowest. This was similar to the neighbors i built as i aimed to reach the bottom left to top right each time ## The linear model

It is rather cumbersome to try to get the correct least squares line, i.e. the line that minimizes the sum of squared residuals, through trial and error. Instead, you can use the lm function in R to fit the linear model (a.k.a. regression line).

```
m1 <- lm(pf_score ~ pf_expression_control, data = hfi)</pre>
```

The first argument in the function lm is a formula that takes the form y ~ x. Here it can be read that we want to make a linear model of pf_score as a function of pf_expression_control. The second argument specifies that R should look in the hfi data frame to find the two variables.

The output of 1m is an object that contains all of the information we need about the linear model that was just fit. We can access this information using the summary function.

```
summary(m1)
```

```
##
## Call:
## lm(formula = pf_score ~ pf_expression_control, data = hfi)
```

```
##
## Residuals:
##
       Min
                1Q
                   Median
                                       Max
   -3.8467 -0.5704
                    0.1452
                            0.6066
                                    3.2060
##
##
##
  Coefficients:
##
                         Estimate Std. Error t value Pr(>|t|)
                          4.61707
                                                        <2e-16 ***
## (Intercept)
                                     0.05745
                                                80.36
  pf_expression_control 0.49143
                                     0.01006
                                                48.85
                                                        <2e-16 ***
##
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 0.8318 on 1376 degrees of freedom
##
     (80 observations deleted due to missingness)
## Multiple R-squared: 0.6342, Adjusted R-squared: 0.634
## F-statistic: 2386 on 1 and 1376 DF, p-value: < 2.2e-16
```

Let's consider this output piece by piece. First, the formula used to describe the model is shown at the top. After the formula you find the five-number summary of the residuals. The "Coefficients" table shown next is key; its first column displays the linear model's y-intercept and the coefficient of pf_expression_control. With this table, we can write down the least squares regression line for the linear model:

```
\hat{y} = 4.61707 + 0.49143 \times pf\_expression\_control
```

One last piece of information we will discuss from the summary output is the Multiple R-squared, or more simply, R^2 . The R^2 value represents the proportion of variability in the response variable that is explained by the explanatory variable. For this model, 63.42% of the variability in runs is explained by at-bats.

5. Fit a new model that uses pf_expression_control to predict hf_score, or the total human freedom score. Using the estimates from the R output, write the equation of the regression line. What does the slope tell us in the context of the relationship between human freedom and the amount of political pressure on media content? 5.153687 + 0.349862 x pf_expression_control The slope tells us the steepness of the relationship, ie how direct their relationship is

```
m2<-lm(hf_score ~ pf_expression_control, data = hfi)
summary(m2)</pre>
```

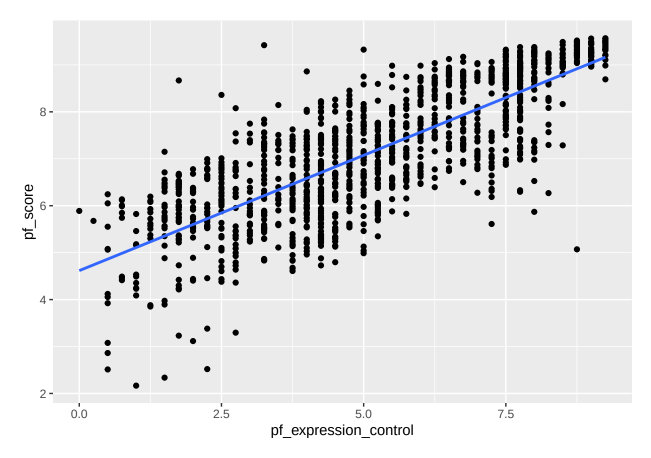
```
##
## Call:
##
  lm(formula = hf_score ~ pf_expression_control, data = hfi)
##
## Residuals:
##
       Min
                1Q
                    Median
                                3Q
                                       Max
##
  -2.6198 -0.4908
                   0.1031
                           0.4703
                                    2.2933
##
##
  Coefficients:
##
                         Estimate Std. Error t value Pr(>|t|)
                         5.153687
                                    0.046070
                                              111.87
                                                        <2e-16 ***
## (Intercept)
## pf expression control 0.349862
                                    0.008067
                                                43.37
                                                        <2e-16 ***
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

```
## Residual standard error: 0.667 on 1376 degrees of freedom
## (80 observations deleted due to missingness)
## Multiple R-squared: 0.5775, Adjusted R-squared: 0.5772
## F-statistic: 1881 on 1 and 1376 DF, p-value: < 2.2e-16</pre>
```

Prediction and prediction errors

Let's create a scatterplot with the least squares line for ${\tt m1}$ laid on top.

```
ggplot(data = hfi, aes(x = pf_expression_control, y = pf_score)) +
geom_point() +
stat_smooth(method = "lm", se = FALSE)
```



Here, we are literally adding a layer on top of our plot. **geom_smooth** creates the line by fitting a linear model. It can also show us the standard error **se** associated with our line, but we'll suppress that for now.

This line can be used to predict y at any value of x. When predictions are made for values of x that are beyond the range of the observed data, it is referred to as extrapolation and is not usually recommended. However, predictions made within the range of the data are more reliable. They're also used to compute the residuals.

6. If someone saw the least squares regression line and not the actual data, how would they predict a country's personal freedom school for one with a 6.7 rating for pf_expression_control? Is this an overestimate or an underestimate, and by how much? In other words, what is the residual for this prediction? I would predict it by finding the yhat for the regression where 6.7 meets the point. I don't

think this is an overestimate as it falls within the range of the points, it would be accurate dependent on the parameters of the regression

```
hfi6.7 <- hfi %>%
    filter(pf_expression_control > 6.7 & pf_expression_control <6.8)
mean(hfi6.7$pf_score)

## [1] 8.006315

y = 4.61707 + 0.49143*6.7

y

## [1] 7.909651

residual = mean(hfi6.7$pf_score) - y
residual

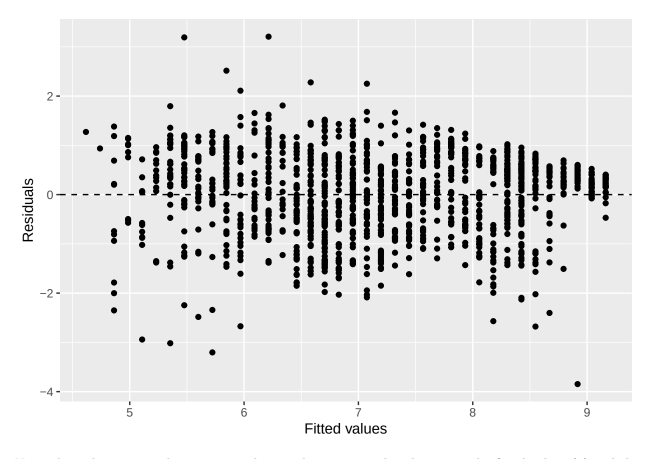
## [1] 0.09666408
```

Model diagnostics

To assess whether the linear model is reliable, we need to check for (1) linearity, (2) nearly normal residuals, and (3) constant variability.

Linearity: You already checked if the relationship between pf_score and 'pf_expression_control' is linear using a scatterplot. We should also verify this condition with a plot of the residuals vs. fitted (predicted) values.

```
ggplot(data = m1, aes(x = .fitted, y = .resid)) +
  geom_point() +
  geom_hline(yintercept = 0, linetype = "dashed") +
  xlab("Fitted values") +
  ylab("Residuals")
```



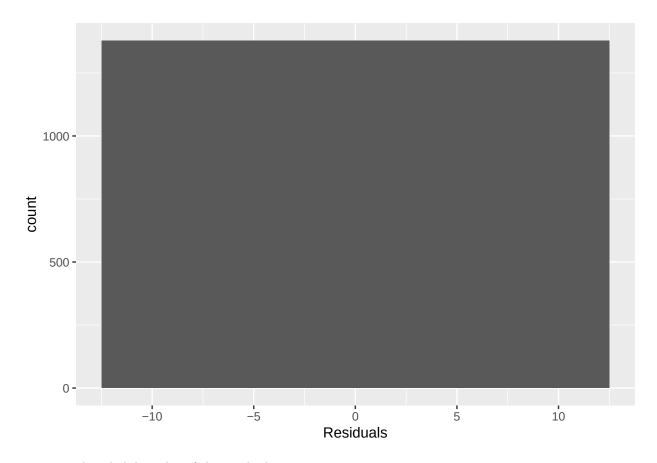
Notice here that m1 can also serve as a data set because stored within it are the fitted values (\hat{y}) and the residuals. Also note that we're getting fancy with the code here. After creating the scatterplot on the first layer (first line of code), we overlay a horizontal dashed line at y = 0 (to help us check whether residuals are distributed around 0), and we also reanme the axis labels to be more informative.

7. Is there any apparent pattern in the residuals plot? What does this indicate about the linearity of the relationship between the two variables?

There is no apparent pattern. Thus illustrating the likelihood of a linearity

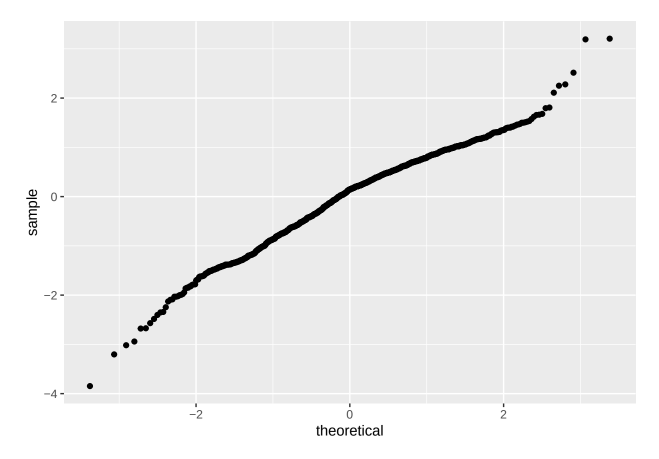
Nearly normal residuals: To check this condition, we can look at a histogram

```
ggplot(data = m1, aes(x = .resid)) +
  geom_histogram(binwidth = 25) +
  xlab("Residuals")
```



or a normal probability plot of the residuals.

```
ggplot(data = m1, aes(sample = .resid)) +
  stat_qq()
```



Note that the syntax for making a normal probability plot is a bit different than what you're used to seeing: we set sample equal to the residuals instead of x, and we set a statistical method qq, which stands for "quantile-quantile", another name commonly used for normal probability plots.

8. Based on the histogram and the normal probability plot, does the nearly normal residuals condition appear to be met? Yes it does, it seems a tiny bit skewed though

Constant variability:

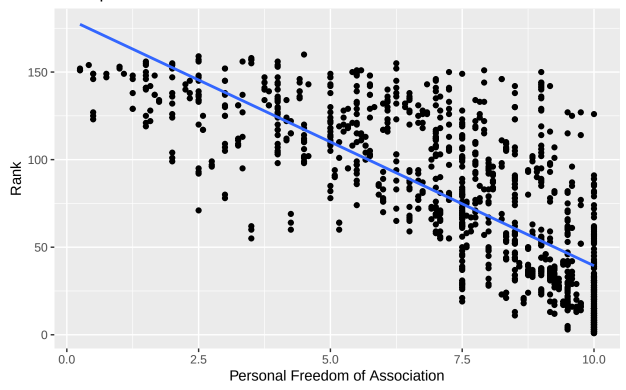
9. Based on the residuals vs. fitted plot, does the constant variability condition appear to be met? Yes it appears to be met

More Practice

• Choose another freedom variable and a variable you think would strongly correlate with it.. Produce a scatterplot of the two variables and fit a linear model. At a glance, does there seem to be a linear relationship? pf_association and pf_rank It appears linear from a glance but no strong concentration of points along the regression

```
ggplot(hfi, aes(x=pf_association, y=pf_rank)) + geom_point() +
geom_smooth(method = "lm", se = FALSE) +
ggtitle("Human Freedom Index-Personal Freedom \nRank per Association") +
xlab("Personal Freedom of Association") + ylab("Rank")
```

Human Freedom Index-Personal Freedom Rank per Association



```
m3 <- lm(pf_rank ~ pf_association, data = hfi)
summary(m3)</pre>
```

```
##
## Call:
## lm(formula = pf_rank ~ pf_association, data = hfi)
##
## Residuals:
##
      Min
               1Q Median
                                3Q
                                      Max
## -76.308 -21.902 -6.476 20.781
                                   96.455
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 180.7927
                              2.8036
                                       64.49
                                               <2e-16 ***
                                      -39.48
                                                <2e-16 ***
## pf_association -14.1386
                              0.3581
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 29.91 on 1127 degrees of freedom
     (329 observations deleted due to missingness)
## Multiple R-squared: 0.5804, Adjusted R-squared:
## F-statistic: 1559 on 1 and 1127 DF, p-value: < 2.2e-16
```

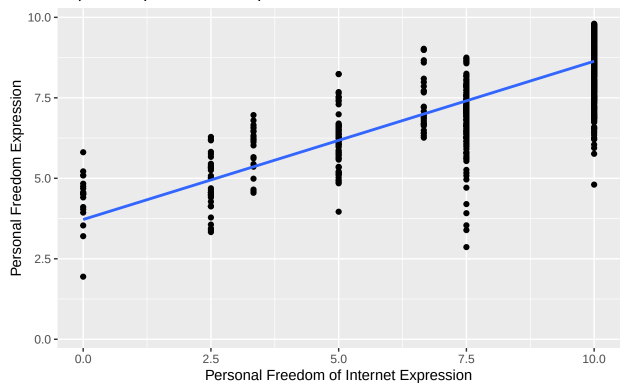
• How does this relationship compare to the relationship between $pf_{expression}_{ontrol}$ and $pf_{expression}_{ontrol}$. Use the R^2 values from the two model summaries to compare. Does your independent

variable seem to predict your dependent one better? Why or why not? This prediction is less accurate as it has a R-squared of 0.5804 while pf_expression_control and pf_score had a R-squared of 0.6342

• What's one freedom relationship you were most surprised about and why? Display the model diagnostics for the regression model analyzing this relationship. I was surprised by pf_expression_internet, pf_expression, you'd expect them to walk hand in hand as the appeared to internet gave freedom of expression. Initially i though it had no correlation but it had a 0.6277 R-squared and a successful q plot but a weird looking fitted vs residual

```
ggplot(hfi, aes(x=pf_expression_internet, y=pf_expression)) + geom_point() +
geom_smooth(method = "lm", se = FALSE) +
ggtitle("Human Freedom Index-Personal Freedom \nExpressionper Internet Expression ") +
xlab("Personal Freedom of Internet Expression") + ylab("Personal Freedom Expression")
```

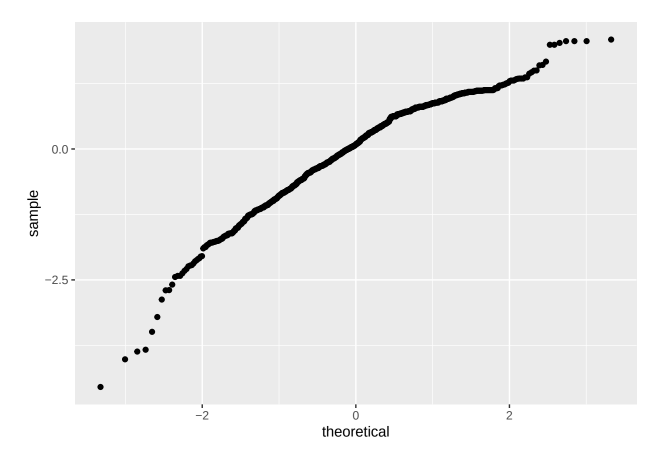
Human Freedom Index–Personal Freedom Expressionper Internet Expression



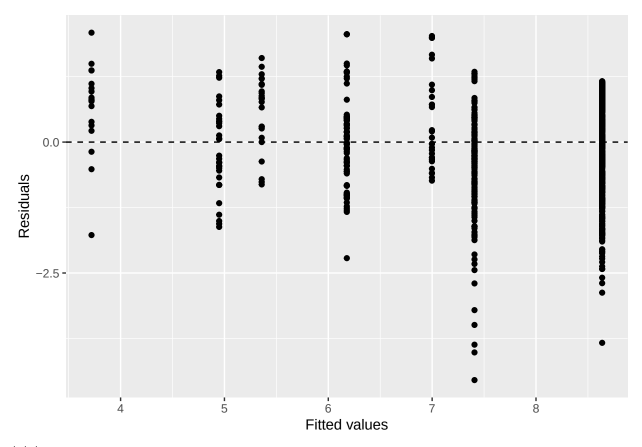
```
m5 <- lm(pf_expression ~ pf_expression_internet, data = hfi)
summary(m5)</pre>
```

```
##
## Call:
## lm(formula = pf_expression ~ pf_expression_internet, data = hfi)
##
## Residuals:
## Min    1Q Median    3Q Max
## -4.5446 -0.5575    0.0862    0.7155    2.0895
```

```
##
## Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                          3.72006
                                     0.10012
                                               37.15
                                                       <2e-16 ***
## pf_expression_internet 0.49170
                                     0.01128
                                               43.59
                                                       <2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 0.8949 on 1127 degrees of freedom
     (329 observations deleted due to missingness)
## Multiple R-squared: 0.6277, Adjusted R-squared: 0.6274
## F-statistic: 1900 on 1 and 1127 DF, p-value: < 2.2e-16
ggplot(data = m5, aes(sample = .resid)) +
 stat_qq()
```



```
ggplot(data = m5, aes(x = .fitted, y = .resid)) +
  geom_point() +
  geom_hline(yintercept = 0, linetype = "dashed") +
  xlab("Fitted values") +
  ylab("Residuals")
```



* * *