BIFX 553 - Discussion 4

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Modeling and testing complex interactions

As a reminder, let's write out our model from a few weeks back:

$$nodes_i = \beta_0 + \beta_1 age_i + \beta_2 size_i + \beta_3 grade_i + \varepsilon_i$$

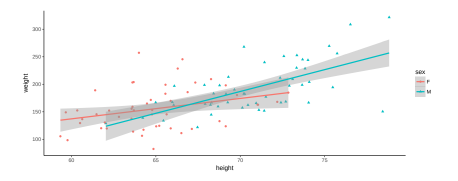
- nodes; is our outcome variable
- β_0, β_1, \ldots are our regression coefficients
- ▶ age_i, size_i, . . . are our predictors
- \triangleright ε_i are the error terms for the model (i.e. how far off is the model prediction from what we observed)

For variables that are categorical (e.g. sex, race), we normally include dummy variables for them in the model.

$$weight_i = \beta_0 + \beta_1 height_i + \beta_2 (sex_i = M) + \varepsilon_i$$

```
set.seed(293847)
# data for our categorical predictor example
bmi <- data frame(</pre>
         sex = c(rep(c('M', 'F'), each = 50)),
         # average US height
         height = rnorm(100, mean = ifelse(sex == 'M', 70, 65),
                         sd = ifelse(sex == 'M', 4, 3.5)),
                                   # ave BMT - men
         bmi = ifelse(sex == 'M', 2^rnorm(100, 4.7004, 0.2305),
                                   # ave BMI - women
                                   2<sup>rnorm(100, 4.6439, 0.3086))</sup>
         # calculate weight
         weight = ((height / 12 / 3.28)^2 * bmi) * 2.2)
```

```
# graphical view of relationsihp btween height and weight
ggplot(bmi, aes(x = height, y = weight, group = sex)) +
geom_point(aes(color = sex, shape = sex)) +
geom_smooth(method = 'lm', aes(color = sex))
```



```
# and now for a statistical look
lm(weight ~ height + sex, data = bmi) %>%
tidy()
```

```
## term estimate std.error statistic p.value

## 1 (Intercept) -236.278051 69.799873 -3.3850785 1.028381e-03

## 2 height 6.032593 1.071821 5.6283572 1.764689e-07

## 3 sexM 2.437032 9.625050 0.2531969 8.006519e-01
```

In the background R is doing the following:

```
# create a dummy variable for "sex == 'M'"
bmi$male <- as.integer(bmi$sex == 'M')

# this model is exactly the same as the model above
lm(weight ~ height + male, data = bmi) %>%
tidy()
```

```
## term estimate std.error statistic p.value

## 1 (Intercept) -236.278051 69.799873 -3.3850785 1.028381e-03

## 2 height 6.032593 1.071821 5.6283572 1.764689e-07

## 3 male 2.437032 9.625050 0.2531969 8.006519e-01
```

When we write this out in a formula, we get

weight_i =
$$\beta_0 + \beta_1 height_i + \beta_2 (sex_i = M) + \varepsilon_i$$

= $\beta_0 + \beta_1 height_i + \beta_2 (1) + \varepsilon_i$
= $\beta_0 + \beta_1 height_i + \beta_2 + \varepsilon_i$

when sex_i is 'M'

weight_i =
$$\beta_0 + \beta_1 height_i + \beta_2 (0) + \varepsilon_i$$

= $\beta_0 + \beta_1 height_i + \varepsilon_i$

when sex_i is 'F'

So, exactly what does β_2 represent? It is interpreted as the difference in height between a male and a female who are otherwise similar (i.e. in our case, they are the same height).

$$\begin{split} \mathsf{E}(\textit{weight}_i - \textit{weight}_j | \textit{height}_i &= \textit{height}_j) = \beta_0 + \beta_1 \textit{height}_i + \beta_2 (\textit{sex}_i = \textit{M}) \\ &- (\beta_0 + \beta_1 \textit{height}_j + \beta_2 (\textit{sex}_j = \textit{M})) \\ &= \beta_0 + \beta_1 \textit{height}_i + \beta_2 (1) \\ &- (\beta_0 + \beta_1 \textit{height}_i + \beta_2 (0)) \\ &= \beta_2 \\ &= 2.4 \text{ in} \end{split}$$

An asside on the interpretation of the intercept

What does the intercept (β_0) really mean? In the model above, β_0 is what we get when $sex_i = {}^{\iota}F'$ and $height_i = 0$. So, the predicted weight of a Female who is 0 inches tall would be

$$E(weight_i) = \beta_0 + \beta_1 height_i + \beta_2 (sex_i = M)$$

$$= \beta_0 + \beta_1 (0) + \beta_2 (0)$$

$$= \beta_0$$

$$= -236 \text{ lbs}$$

This, of course, makes no sense, neither the height nor the weight. For this reason, the intercept is often ignored.

An asside on the interpretation of the intercept

If we wanted the intercept to make sense, we could recenter $height_i$ like so:

```
\# lets center height at the sample median (or there abouts) median(bmi\$height)
```

```
## [1] 67.89892
```

```
## term estimate std.error statistic p.value
## 1 (Intercept) 167.905685 5.608466 29.9378991 8.506954e-51
## 2 ht_med 6.032593 1.071821 5.6283572 1.764689e-07
## 3 sexM 2.437032 9.625050 0.2531969 8.006519e-01
```

An asside on the interpretation of the intercept

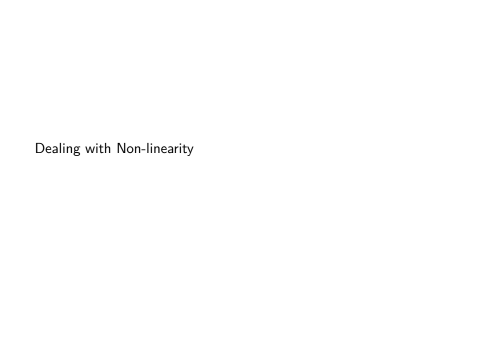
Now, the intercept has a more useful meeing. It represents the expected weight of a 5'7" female:

$$E(weight_{i}) = \beta_{0} + \beta_{1}(height_{i} - 67) + \beta_{2}(sex_{i} = M)$$

$$= \beta_{0} + \beta_{1}(67 - 67) + \beta_{2}(0)$$

$$= \beta_{0}$$

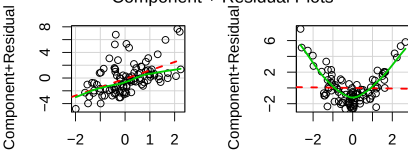
$$= 168 \text{ lbs}$$



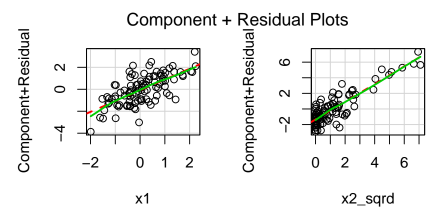
Simple non-linear terms

We saw in Discussion 3 that the number of nodes in the gbsg data set was more log-normally than normally distributed. Also, we looked at an example where the relationship between a predictor was a quadratic relationship, rather than a linear relationship. What do we do in these cases?



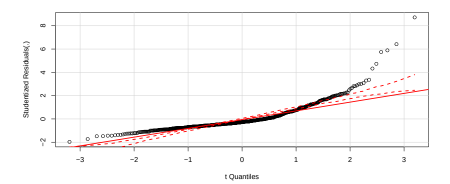


```
# create a new variable and try this
mutate(tmp, x2_sqrd = x2^2) %>%
lm(formula = y ~ x1 + x2_sqrd) %>%
crPlots() # looks much better!
```

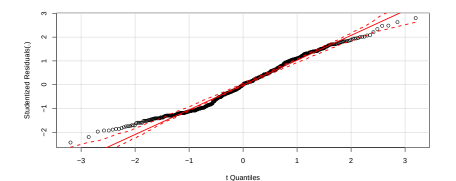


```
# log-normally distributed outcome variable
load('../1-26/gbsg.RData')

# wrong model... nodes is log-normally distributed
lm(nodes ~ size + grade, data = gbsg) %>%
    qqPlot()
```

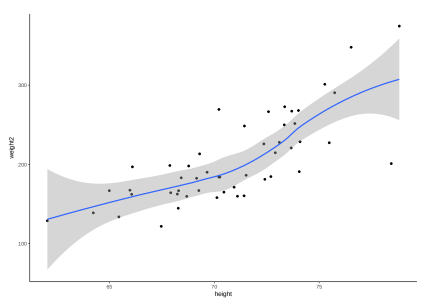


```
# lets try with log(nodes) ... still not perfect
mutate(gbsg, lnodes = log(nodes)) %>%
lm(formula = lnodes ~ size + grade) %>%
# a lot better -- we will revisit this in a future discussion
qqPlot()
```

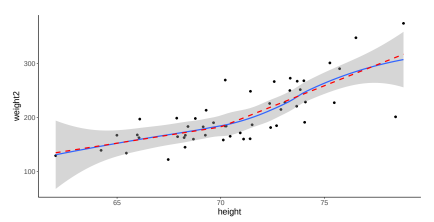


Sometimes, the relationship between a predictor and the outcome is neither linear, nor does it conform to a nice function that we can use for a simple transformation. In such a case, splines can help.

```
## `geom_smooth()` using method = 'loess'
```



```
## `geom_smooth()` using method = 'loess'
```



Splines: Interpretation/Prediction

Interpreting the regression coefficients directly is beyond the scope of this class, but there is an easy way to use this model to predict. This should work with any of the models we will be generating in this class. All you need is the model output from 1m and a data frame with the values needed to predict.

```
# for the male-only model
data_frame(height = 70) %>%
  predict(object = mdl1)
```

```
## 1
## 182.2475
```

Splines: Interpretation/Prediction

In mdl1, we only needed to include height, but we also need to include sex for mdl2.

```
## 1 2
## 179.8799 163.0864
```

Splines: Higher order splines

You can also use the bs function to fit higher order polynomial splines (the default is a cubic spline).

Non-parametric regression

You may have noticed that the default output of <code>geom_smooth()</code> is a smooth function that looks sort of like a moving average. The default smoother used for data sets with less than 1000 observations is called loess, and a generalized additive model is used with larger data sets. You won't get p-values from this, but it can be a valuable tool to graphically see what is going on.

Non-parametric regression

```
## `geom_smooth()` using method = 'loess'
```

