Monday, Feb 14

Assumptions

Mathematical (statistical) models make assumptions, and results (statistical inferences) based on the models are derived using those assumptions.

Example: Assume that an object is a *cone*. It can be shown (with a little calculus) that

$$V = \pi r^2 h/3$$
 and $A = \pi r \left(r + \sqrt{r^2 + h^2}\right)$,

based on the assumption that $the\ object\ is\ a\ cone.$

"All models are wrong but some are useful." — George E. P. Box

Implicit Assumptions of Linear and Nonlinear Regression

Discussions of assumptions are based on an alternative representation of a regression model. A linear model can be written as

$$Y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_k x_{ik} + \epsilon_i$$

and a nonlinear model can be written as

$$Y_i = f(x_{i1}, x_{i2}, \dots, x_{ik}) + \epsilon_i,$$

where the linear model with

$$f(x_{i1}, x_{i2}, \dots, x_{ik}) = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \dots + \beta_k x_{ik}$$

is a special case.

There are four implicit assumptions about ϵ_i that go into the derivation of routine/default methods for making inferences concerning the model.

- 1. $E(\epsilon_i) = 0$ for all i.
- 2. $Var(\epsilon_i) = \sigma^2$ for all i.
- 3. $Cov(\epsilon_i, \epsilon_{i'}) = 0$ for all $i \neq i'$.
- 4. Each ϵ_i has a normal distribution.

How should we approach each assumption?

- 1. How do we define each assumption?
- 2. What are the *consequences* if the assumption is (very) wrong?
- 3. How do we *detect* if the assumption is (very) wrong?
- 4. What is/are the solution(s) if the assumption is (very) wrong?

Assumption 1: Zero Expectations of Errors

Definition: The assumption $E(\epsilon_i) = 0$ implies that $E(Y_i)$ depends on the explanatory variables in the way assumed by the model. That is, if we have the *assumed* model

$$E(Y_i) = f(x_{i1}, x_{i2}, \dots, x_{ik})$$

then

$$E(\epsilon_i) = 0 \Rightarrow E(Y_i) = f(x_{i1}, x_{i2}, \dots, x_{ik}).$$

For the model to be a correct representation of the relationship between $E(Y_i)$ and $x_{i1}, x_{i2}, \ldots, x_{ik}$ we require that $E(\epsilon_i) = 0$ for all $i = 1, 2, \ldots, n$.

Consequences: Estimates of parameters or some functions thereof (e.g., linear combinations) may be biased.

Detection: Residuals are statistics that are frequently used to empirically investigate assumption violations. There are several types of residuals.

1. Raw residuals. These are simply estimates of ϵ_i . In a linear model, for example, the error is

$$\epsilon_i = Y_i - (\beta_0 + \beta_1 x_{i1} + \dots + \beta_k x_{ik}),$$

by definition, so a simple estimator of ϵ_i is the residual

$$e_i = Y_i - \hat{Y}_i$$

where

$$\hat{Y}_i = \hat{\beta}_0 + \hat{\beta}_1 x_{i1} + \dots + \hat{\beta}_k x_{ik}.$$

We can define the raw residual in a similar way for a nonlinear model.

2. Standardized residuals. Defined as

$$z_i = \frac{e_i}{SE(e_i)}.$$

If the model assumptions are *correct* then z_i is approximately standard normal in distribution so we expect that about 95% of such residuals to satisfy $|z_i| < 2$.

3. Studentized residuals. Defined as

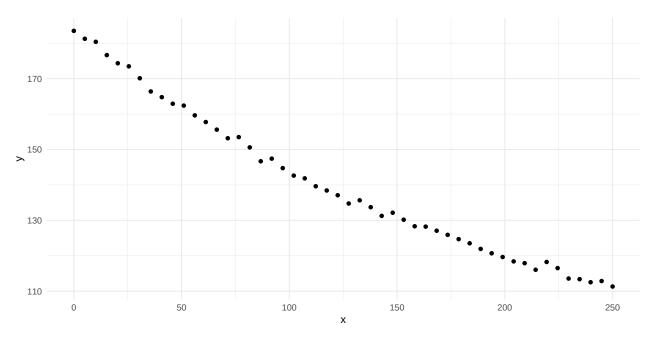
$$t_i = \frac{e_i}{SE_{(i)}(e_i)},$$

where $\mathrm{SE}_{(i)}(e_i)$ is the standard error of e_i estimated by leaving out that observation. This avoids bias in the standard error in cases where $E(\epsilon_i) \neq 0$. If the model assumptions are met then each t_i has a t distribution with one less degree of freedom than the residual degrees of freedom (i.e., n-p-1 where p is the number of parameters in the model). Unless n-p-1 is very small, we expect that about 95% of studentized residuals satisfy $|t_i| < 2$.

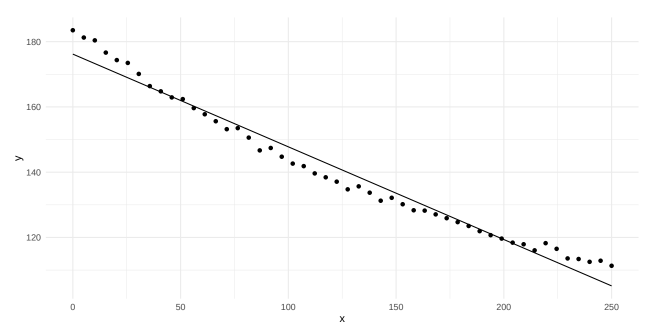
What to look for in residuals:

- 1. Individual observations with exceptional residuals.
- 2. More residuals than expected overall that are exceptional.
- 3. Changes in the distribution of residuals when plotting against \hat{y}_i .

Example: Consider the following artificial data.



Suppose we tried a linear model.



What are the observations with "exceptionally large" residuals?

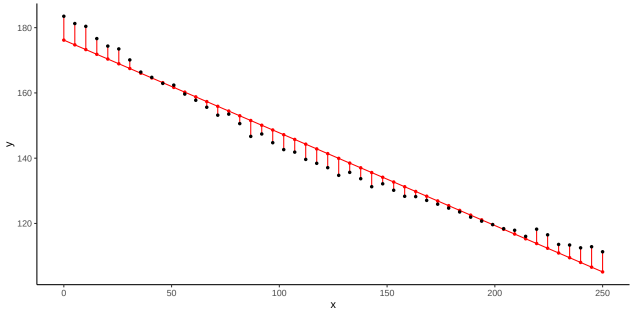
subset(fakedata, abs(rest) > 2)

x y yhat rese rest 1 0.0 183.5 176.2 7.311 2.198 3 10.2 180.4 173.3 7.113 2.121

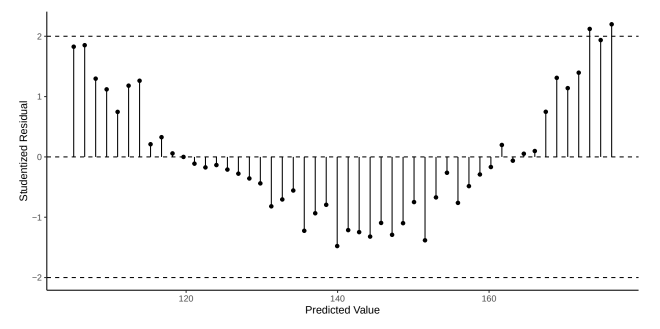
library(dplyr)
fakedata %>% filter(abs(rest) > 2)

x y yhat rese rest 1 0.0 183.5 176.2 7.311 2.198 2 10.2 180.4 173.3 7.113 2.121

Are there any patterns?



```
p <- ggplot(fakedata, aes(x = yhat, y = rest))
p <- p + geom_segment(aes(x = yhat, xend = yhat, y = 0, yend = rest))
p <- p + geom_point() + theme_classic()
p <- p + labs(x = "Predicted Value", y = "Studentized Residual")
p <- p + geom_hline(yintercept = c(-2, 0, 2), linetype = 2)
plot(p)</pre>
```

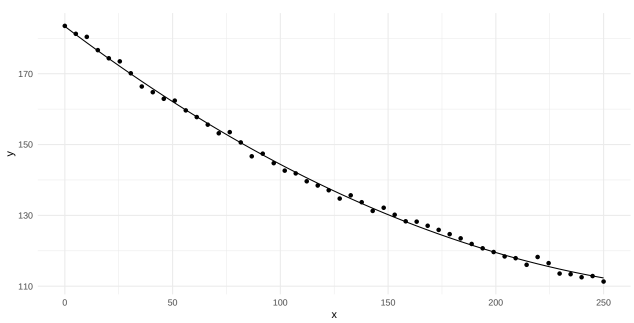


Let's try the polynomial model $E(Y_i) = \beta_0 + \beta_1 x_i + \beta_2 x_i^2$.

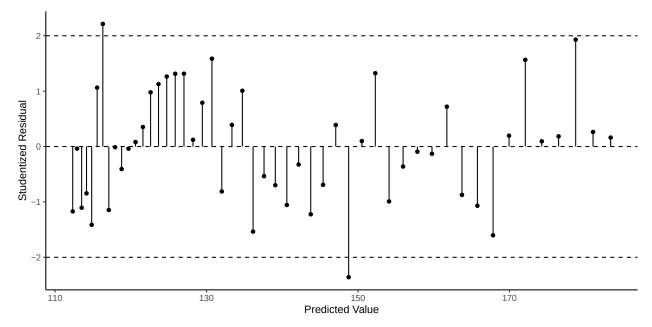
```
mpoly <- lm(y ~ poly(x, degree = 2), data = fakedata)
fakedata$yhat <- predict(mpoly)
fakedata$rest <- rstudent(mpoly)

p <- ggplot(fakedata, aes(x = x, y = y)) + theme_minimal()</pre>
```

```
p <- p + geom_point() + geom_line(aes(y = yhat))
plot(p)</pre>
```



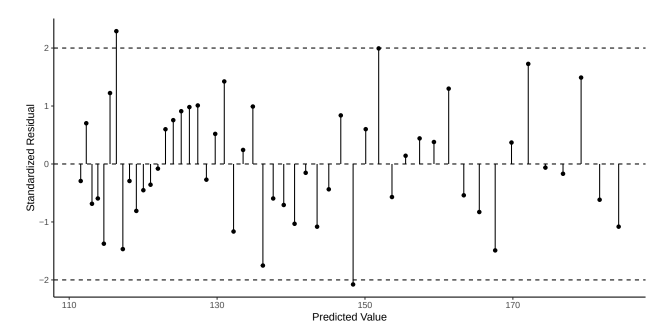
```
p <- ggplot(fakedata, aes(x = yhat, y = rest)) +
  geom_segment(aes(x = yhat, xend = yhat, y = 0, yend = rest)) +
  geom_point() + theme_classic() +
  labs(x = "Predicted Value", y = "Studentized Residual") +
  geom_hline(yintercept = c(-2, 0, 2), linetype = 2)
plot(p)</pre>
```



The "correct" model is the nonlinear model

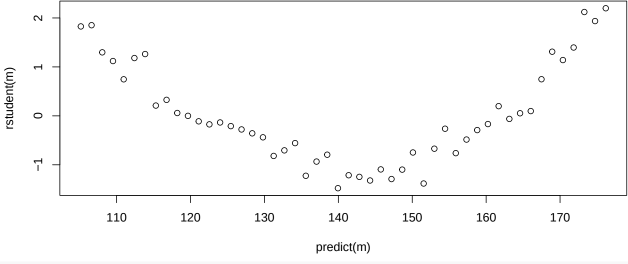
$$E(Y_i) = \alpha + (\delta - \alpha)2^{-x_i/\gamma}.$$

```
mnlin <- nls(y ~ alpha + (delta - alpha) * 2^(-x / gamma),</pre>
  data = fakedata, start = list(alpha = 80, delta = 180, gamma = 140))
d <- nlsint(mnlin, residuals = TRUE) # nlsint is from trtools package</pre>
head(d)
    fit
            se
                 lwr
                        upr
                                hat
1 184.3 0.3894 183.5 185.1 0.21231 -1.08147
2 181.7 0.3481 181.0 182.4 0.16966 -0.61684
3 179.2 0.3112 178.6 179.9 0.13564 1.49073
4 176.8 0.2788 176.2 177.3 0.10888 -0.16877
5 174.4 0.2510 173.9 174.9 0.08820 -0.06274
6 172.1 0.2276 171.6 172.5 0.07256 1.72687
p \leftarrow ggplot(d, aes(x = fit, y = res)) +
  geom_segment(aes(x = fit, xend = fit, y = 0, yend = res)) +
  geom_point() + theme_classic() +
  labs(x = "Predicted Value", y = "Standardized Residual") +
  geom_hline(yintercept = c(-2, 0, 2), linetype = 2)
plot(p)
```

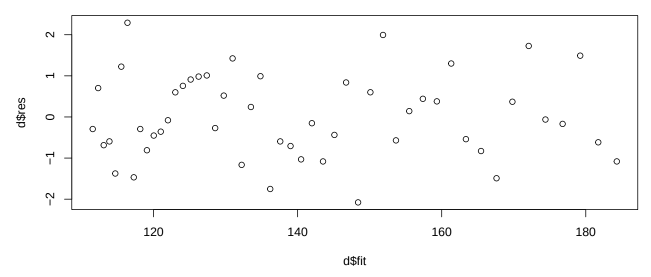


Note that since we are usually not interested in created publication quality residual plots, we can do some quick-and-dirty plots with simpler (but uglier) graphics.

```
# when using lm
m <- lm(y ~ x, data = fakedata)
plot(predict(m), rstudent(m))</pre>
```



```
# when using nls
mnlin <- nls(y ~ alpha + (delta - alpha) * 2^(-x / gamma),
   data = fakedata, start = list(alpha = 80, delta = 180, gamma = 140))
d <- nlsint(mnlin, residuals = TRUE) # nlsint is from trtools package
plot(d$fit, d$res)</pre>
```



Example: Consider a model where the expected MPG of cars is assumed to be a linear function of weight and rear axle ratio.

```
cars <- read.csv("http://webpages.uidaho.edu/~trjohns/cars.csv")
cars <- cars[,c(2,3,4,5)] # select columns 2, 3, 4, and 5
head(cars)</pre>
```

```
car mpg weight ratio
         Buick Estate Wagon 16.9
                                  4.360
                                         2.73
                                         2.26
2 Ford Country Squire Wagon 15.5
                                  4.054
         Chevy Malibu Wagon 19.2
                                  3.605
                                         2.56
4
     Chrysler LeBaron Wagon 18.5
                                  3.940
                                         2.45
5
                   Chevette 30.0
                                  2.155
                                         3.70
6
              Toyota Corona 27.5 2.560
                                         3.05
```

```
m <- lm(mpg ~ weight + ratio, data = cars)</pre>
cars$yhat <- predict(m)</pre>
cars$rest <- rstudent(m)</pre>
subset(cars, abs(rest) > 2)
                        car mpg weight ratio yhat rest
1 Buick Estate Wagon 16.9
                                      4.36 2.73 10.32 3.944
34
             Fiat Strada 37.3
                                      2.13 3.10 32.67 2.348
p \leftarrow ggplot(cars, aes(x = rest, y = car))
p <- p + geom_point() + theme_classic()</pre>
p \leftarrow p + geom_segment(aes(x = 0, xend = rest, y = car, yend = car))
p \leftarrow p + labs(x = "Studentized Residual", y = NULL)
plot(p)
           VW Scirocco
             VW Rabbit
            VW Dasher
           Volvo 240 GL
          Toyota Corona
          Saab 99 GLE
        Pontiac Phoenix
        Plymouth Horizon
        Peugeot 694 SL
           Ölds Omega
         Mercury Zephyr
   Mercury Grand Marquis
            Mazda GLC
       Honda Accord LX
      Ford Mustang Ghia
         Ford Mustang 4
              Ford LTD
Ford Country Squire Wagon
            Fiat Strada
        Dodge St Regis
           Dodge Omni
            Dodge Colt
          Dodge Aspen
            Datsun 810
            Datsun 510
            Datsun 210
  Chrysler LeBaron Wagon
     Chevy Malibu Wagon
          Chevy Citation
    Chevy Caprice Classic
              Chevette
          Buick Skylark
     Buick Estate Wagon
    Buick Century Special
             BMW 320i
             Audi 5000
            AMC Spirit
```

What is up with the Buick Estate Wagon?

AMC Concord D/L

Solutions: Modify the model. Drop offending observations(s) if and only if you can justify restricting the scope of the model.

Studentized Residual

Assumption 2: Equality of Error Variances

Definition: In the regression model

$$Y_i = f(x_{i1}, x_{i2}, \dots, x_{ik}) + \epsilon_i,$$

we assume $Var(\epsilon_i) = \sigma^2$ which implies $Var(Y_i) = \sigma^2$. This is called "homoscedasticty" or sometimes "homogeneity of variance" in the context of linear models for designed experiments. A more complete

description of the assumed model is

$$E(Y_i) = f(x_{i1}, x_{i2}, \dots, x_{ik}), \tag{1}$$

$$Var(Y_i) = \sigma^2. (2)$$

Note that the estimator $\hat{\sigma}^2$, the square of the "residual standard error" reported by summary, is computed as

$$\hat{\sigma}^2 = \frac{\sum_{i=1}^n (y_i - \hat{y}_i)^2}{n - p},$$

where p is the number of parameters in the part of the model for $E(Y_i)$ (which is p = k + 1 in a linear model with a β_0 term and k explanatory variables).

Consequences: Severe violations of homoscedaticity can result in two problems.

- 1. Biased standard errors, incorrect p-values, and incorrect confidence/prediction intervals.
- 2. Inefficient estimation of model parameters (and functions thereof).

Detection: Many common patterns of heteroscedasticity can be found by plotting standardized or studentized residuals against \hat{y}_i .

Example: Consider the following data on survival time of terminal cancer patients given a supplement of ascorbate (i.e., vitamin C).

```
library(Stat2Data)
data(CancerSurvival) # this package requires that we "load" the data
head(CancerSurvival)
```

```
Survival Organ
1 124 Stomach
2 42 Stomach
3 25 Stomach
4 45 Stomach
5 412 Stomach
6 51 Stomach
```

For plotting purposes we can order the levels of Organ according to mean survival using reorder.

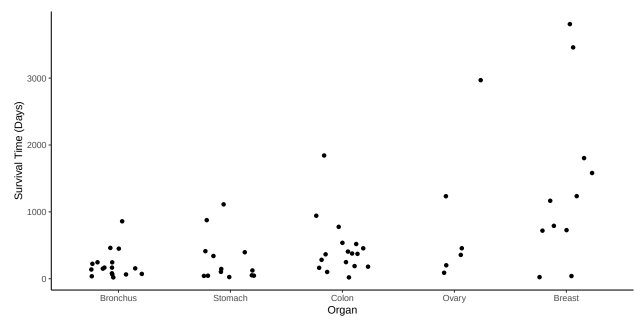
```
CancerSurvival$Organ <- with(CancerSurvival, reorder(Organ, Survival, mean))
```

The with function implies that each variable is "with" the data frame CancerSurvival. This is sometimes nicer than having to identify the data frame explicitly as in the following.

```
CancerSurvival$Organ <- reorder(CancerSurvival$Organ, CancerSurvival$Survival, mean)
```

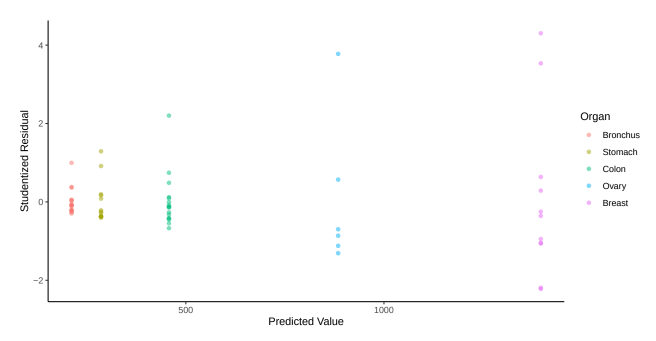
Here is a plot of the data using geom_jitter to space out the points.

```
p <- ggplot(CancerSurvival, aes(x = Organ, y = Survival)) +
  geom_jitter(height = 0, width = 0.25) +
  labs(y = "Survival Time (Days)") + theme_classic()
plot(p)</pre>
```



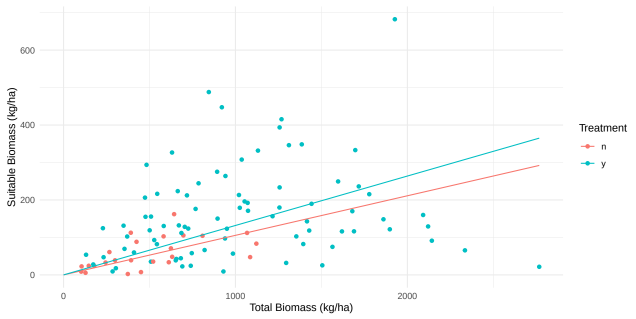
Here we can see some descriptive statistics.

```
library(dplyr)
CancerSurvival %>% group_by(Organ) %>%
  summarize(mean = mean(Survival), stdev = sd(Survival), obs = n())
# A tibble: 5 x 4
  Organ
            mean stdev
                          obs
  <fct>
            <dbl> <dbl> <int>
1 Bronchus
            212.
                   210.
                           17
2 Stomach
            286
                   346.
                           13
3 Colon
            457.
                   427.
                           17
4 Ovary
            884. 1099.
                            6
5 Breast
           1396. 1239.
                           11
Now consider a linear model that assumes homoscedasticity.
m <- lm(Survival ~ Organ, data = CancerSurvival)</pre>
CancerSurvival$yhat <- predict(m)</pre>
CancerSurvival$rest <- rstudent(m)</pre>
head(CancerSurvival)
  Survival
             Organ yhat
1
       124 Stomach 286 -0.2498
2
        42 Stomach
                     286 -0.3765
3
        25 Stomach
                     286 -0.4029
4
                     286 -0.3719
        45 Stomach
5
       412 Stomach
                     286 0.1943
        51 Stomach 286 -0.3626
p <- ggplot(CancerSurvival, aes(x = yhat, y = rest, color = Organ)) +
  geom_point(alpha = 0.5) + theme_classic() +
  labs(x = "Predicted Value", y = "Studentized Residual")
plot(p)
```



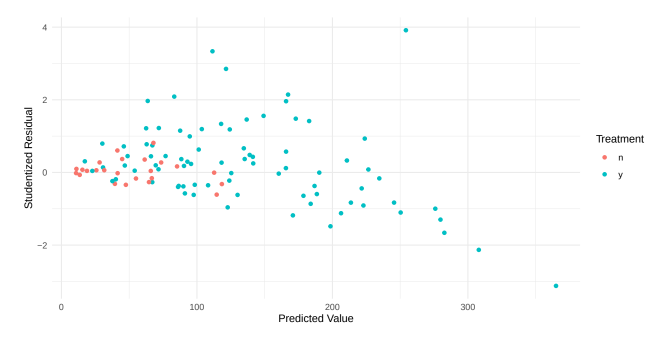
Example: Consider the following data from a study on the effects of fuel reduction on biomass.

```
library(trtools) # for biomass data
m <- lm(suitable ~ -1 + treatment:total, data = biomass)</pre>
summary(m)$coefficients
                  Estimate Std. Error t value Pr(>|t|)
treatmentn:total
                    0.1056
                              0.04183
                                         2.524 1.31e-02
treatmenty:total
                    0.1319
                              0.01121 11.773 7.61e-21
d \leftarrow expand.grid(treatment = c("n","y"), total = seq(0, 2767, length = 10))
d$yhat <- predict(m, newdata = d)</pre>
p \leftarrow ggplot(biomass, aes(x = total, y = suitable, color = treatment)) +
  geom_point() + geom_line(aes(y = yhat), data = d) + theme_minimal() +
  labs(x = "Total Biomass (kg/ha)", y = "Suitable Biomass (kg/ha)",
    color = "Treatment")
plot(p)
```



```
biomass$yhat <- predict(m)
biomass$rest <- rstudent(m)

p <- ggplot(biomass, aes(x = yhat, y = rest, color = treatment)) +
    geom_point() + theme_minimal() +
    labs(x = "Predicted Value", y = "Studentized Residual",
        color = "Treatment")
plot(p)</pre>
```



Solutions: There are several possible solutions.

- 1. Response variable transformation.
- 2. Weighted least squares.

- 3. Robust standard error estimators.
- 4. Models that do not assume constant variance.

We will discuss each of these in turn soon.