

6: Part 1 - Generalized Linear Models

Environmental Data Analytics | John Fay and Luana Lima | Developed by Kateri Salk

Spring 2023

Objectives

1. Describe the components of the generalized linear model (GLM)
2. Apply special cases of the GLM (linear regression) to real datasets
3. Interpret and report the results of linear regressions in publication-style formats
4. Apply model selection methods to choose model formulations

Generalized Linear Models (GLMs)

The analysis of variance (ANOVA), and linear regression are special cases of the **generalized linear model** (GLM). The GLM also includes analyses not covered in this class, including logistic regression, multinomial regression, chi square, and log-linear models. The common characteristic of general linear models is the expression of a continuous response variable as a linear combination of the effects of categorical or continuous explanatory variables, plus an error term that expresses the random error associated with the coefficients of all explanatory variables. The explanatory variables comprise the deterministic component of the model, and the error term comprises the stochastic component of the model. Historically, artificial distinctions were made between linear models that contained categorical and continuous explanatory variables, but this distinction is no longer made. The inclusion of these models within the umbrella of the GLM allows models to fit the main effects of both categorical and continuous explanatory variables as well as their interactions.

Choosing a model from your data: A “cheat sheet”

One-way ANOVA (Analysis of Variance): Continuous response, one categorical explanatory variable with more than two categories. When the categorical explanatory variable just have two categories use the t-test

Two-way ANOVA (Analysis of Variance) Continuous response, two categorical explanatory variables.

Single Linear Regression Continuous response, one continuous explanatory variable.

Multiple Linear Regression Continuous response, two or more continuous explanatory variables.

ANCOVA (Analysis of Covariance) Continuous response, categorical explanatory variable(s) and continuous explanatory variable(s).

If multiple explanatory variables are chosen, they may be analyzed with respect to their **main effects** on the model (i.e., their separate impacts on the variance explained) or with respect to their **interaction effects**, the effect of interacting explanatory variables on the model.

Assumptions of the GLM

The GLM is based on the assumption that the data residuals approximate a normal distribution (or a linearly transformed normal distribution). We will discuss the non-parametric analogues to several of these tests if the assumptions of normality are violated. For tests that analyze categorical explanatory variables, the assumption is that the variance in the response variable is equal among groups. Note: environmental data often violate the assumptions of normality and equal variance, and we will often proceed with a GLM even if these assumptions are violated. In this situation, justifying the decision to proceed with a linear model must be made.

Set up

```
library(tidyverse)
options(scipen = 4)
library(here)
here()

## [1] "/Users/min/Desktop/EDA-Spring2023/R/EDA"

PeterPaul.chem.nutrients <- read.csv(here("Data/Processed_KEY/NTL-LTER_Lake_Chemistry_Nutrients_PeterPaul"))

# Set theme
mytheme <- theme_classic(base_size = 14) +
  theme(axis.text = element_text(color = "black"),
        legend.position = "top")
theme_set(mytheme)
```

Linear Regression

A linear regression is comprised of a continuous response variable, plus a combination of 1+ continuous response variables (plus the error term). The deterministic portion of the equation describes the response variable as lying on a straight line, with an intercept and a slope term. The equation is thus a typical algebraic expression:

$$y = \alpha + \beta * x + \epsilon$$

The goal for the linear regression is to find a **line of best fit**, which is the line drawn through the bivariate space that minimizes the total distance of points from the line. This is also called a “least squares” regression. The remainder of the variance not explained by the model is called the **residual error**.

The linear regression will test the null hypotheses that

1. The intercept (alpha) is equal to zero.
2. The slope (beta) is equal to zero

Whether or not we care about the result of each of these tested hypotheses will depend on our research question. Sometimes, the test for the intercept will be of interest, and sometimes it will not.

Important components of the linear regression are the correlation and the R-squared value. The **correlation** is a number between -1 and 1, describing the relationship between the variables. Correlations close to -1 represent strong negative correlations, correlations close to zero represent weak correlations, and correlations close to 1 represent strong positive correlations. The **R-squared value** is the correlation squared, becoming a number between 0 and 1. The R-squared value describes the percent of variance accounted for by the explanatory variables.

Simple Linear Regression

For the NTL-LTER dataset, can we predict irradiance (light level) from depth?

```
irradiance.regression <- lm(PeterPaul.chem.nutrients$irradianceWater ~ PeterPaul.chem.nutrients$depth)
summary(irradiance.regression)
```

```
## 
## Call:
## lm(formula = PeterPaul.chem.nutrients$irradianceWater ~ PeterPaul.chem.nutrients$depth)
## 
## Residuals:
##     Min      1Q  Median      3Q     Max 
## -458.9  -144.1   -41.2    90.3 23813.0 
## 
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 486.818    4.063   119.82 <2e-16 ***
## PeterPaul.chem.nutrients$depth -95.890    1.153   -83.14 <2e-16 ***
## ---      
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1 
## 
## Residual standard error: 303.4 on 15449 degrees of freedom
## (7557 observations deleted due to missingness)
## Multiple R-squared:  0.3091, Adjusted R-squared:  0.3091 
## F-statistic:  6912 on 1 and 15449 DF,  p-value: < 2.2e-16
```

```
# another way to format the lm() function
irradiance.regression <- lm(data = PeterPaul.chem.nutrients, irradianceWater ~ depth)
summary(irradiance.regression)
```

```
## 
## Call:
## lm(formula = irradianceWater ~ depth, data = PeterPaul.chem.nutrients)
## 
## Residuals:
##     Min      1Q  Median      3Q     Max 
## -458.9  -144.1   -41.2    90.3 23813.0 
## 
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 486.818    4.063   119.82 <2e-16 ***
## depth       -95.890    1.153   -83.14 <2e-16 *** 
## ---      
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1 
## 
## Residual standard error: 303.4 on 15449 degrees of freedom
## (7557 observations deleted due to missingness)
## Multiple R-squared:  0.3091, Adjusted R-squared:  0.3091 
## F-statistic:  6912 on 1 and 15449 DF,  p-value: < 2.2e-16
```

```

# Correlation
cor.test(PeterPaul.chem.nutrients$irradianceWater, PeterPaul.chem.nutrients$depth)

##
## Pearson's product-moment correlation
##
## data: PeterPaul.chem.nutrients$irradianceWater and PeterPaul.chem.nutrients$depth
## t = -83.137, df = 15449, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.5667674 -0.5449776
## sample estimates:
## cor
## -0.555968

```

Question: How would you report the results of this test (overall findings and report of statistical output)?

So, we see there is a significant negative correlation between irradiance and depth (lower light levels at greater depths), and that this model explains about 31 % of the total variance in irradiance. Let's visualize this relationship and the model itself.

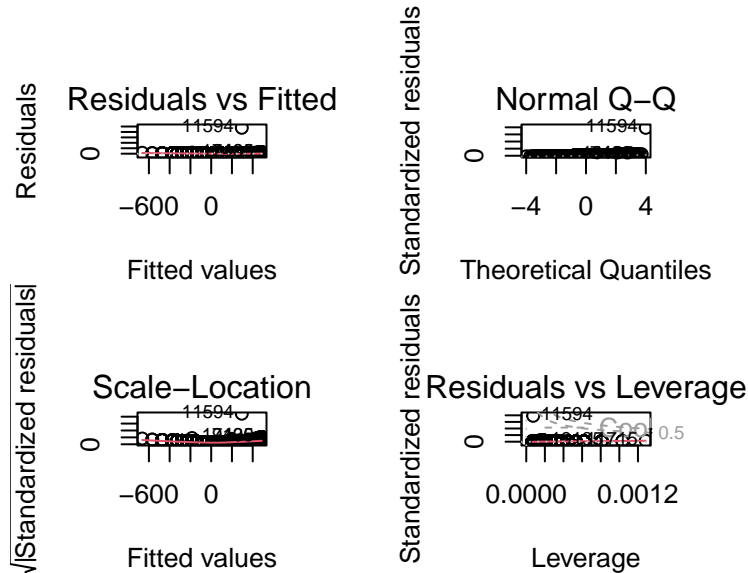
An exploratory option to visualize the model fit is to use the function `plot`. This function will return four graphs, which are intended only for checking the fit of the model and not for communicating results. The plots that are returned are:

1. **Residuals vs. Fitted.** The value predicted by the line of best fit is the fitted value, and the residual is the distance of that actual value from the predicted value. By definition, there will be a balance of positive and negative residuals. Watch for drastic asymmetry from side to side or a marked departure from zero for the red line - these are signs of a poor model fit.
2. **Normal Q-Q.** The points should fall close to the 1:1 line. We often see departures from 1:1 at the high and low ends of the dataset, which could be outliers.
3. **Scale-Location.** Similar to the residuals vs. fitted graph, this will graph the squared standardized residuals by the fitted values.
4. **Residuals vs. Leverage.** This graph will display potential outliers. The values that fall outside the dashed red lines (Cook's distance) are outliers for the model. Watch for drastic departures of the solid red line from horizontal - this is a sign of a poor model fit.

```

par(mfrow = c(2,2), mar=c(4,4,4,4))
plot(irradiance.regression)

```



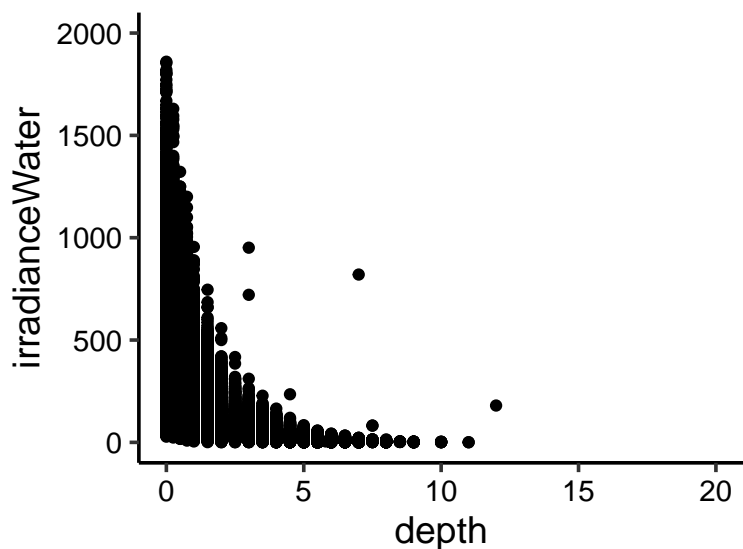
```
#note, plot 1, right side not fit, maybe remove these.plot4 dash line --outlayer
```

```
par(mfrow = c(1,1))
```

The option best suited for communicating findings is to plot the explanatory and response variables as a scatterplot.

```
# Plot the regression
irradiancebydepth <-
  ggplot(PeterPaul.chem.nutrients, aes(x = depth, y = irradianceWater)) +
  ylim(0, 2000) +
  geom_point()
print(irradiancebydepth)
```

Warning: Removed 7558 rows containing missing values ('geom_point()').

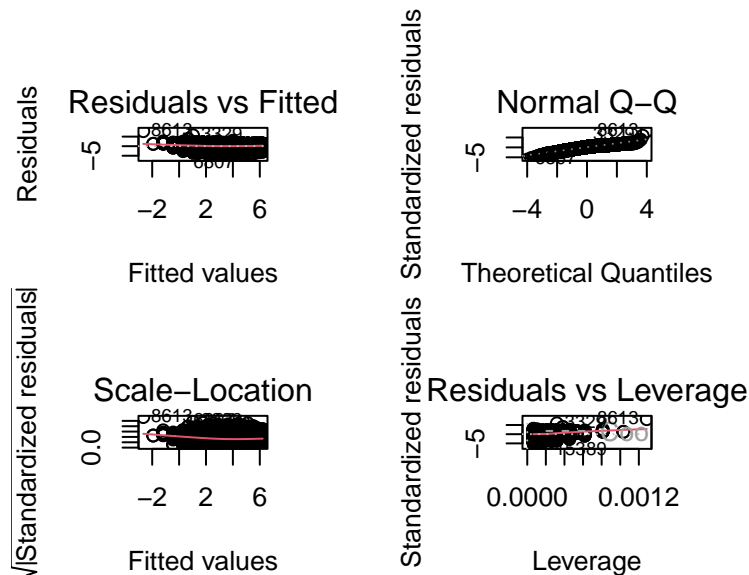


Given the distribution of irradiance values, we don't have a linear relationship between x and y in this case. Let's try log-transforming the irradiance values. Note we also removing the observations that seems to be an outlier.

```
PeterPaul.chem.nutrients <- filter(PeterPaul.chem.nutrients,
                                     irradianceWater != 0 & irradianceWater < 5000)
irradiance.regression2 <- lm(data = PeterPaul.chem.nutrients, log(irradianceWater) ~ depth)
summary(irradiance.regression2)

##
## Call:
## lm(formula = log(irradianceWater) ~ depth, data = PeterPaul.chem.nutrients)
##
## Residuals:
##      Min        1Q    Median        3Q       Max
## -5.9421 -0.5745  0.1930  0.7215  7.8568
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)
## (Intercept) 6.218503  0.012903 481.9   <2e-16 ***
## depth      -0.740198  0.003664 -202.0   <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.9634 on 15445 degrees of freedom
## Multiple R-squared:  0.7255, Adjusted R-squared:  0.7254
## F-statistic: 4.081e+04 on 1 and 15445 DF, p-value: < 2.2e-16

par(mfrow = c(2,2), mar=c(4,4,4,4))
plot(irradiance.regression2)
```

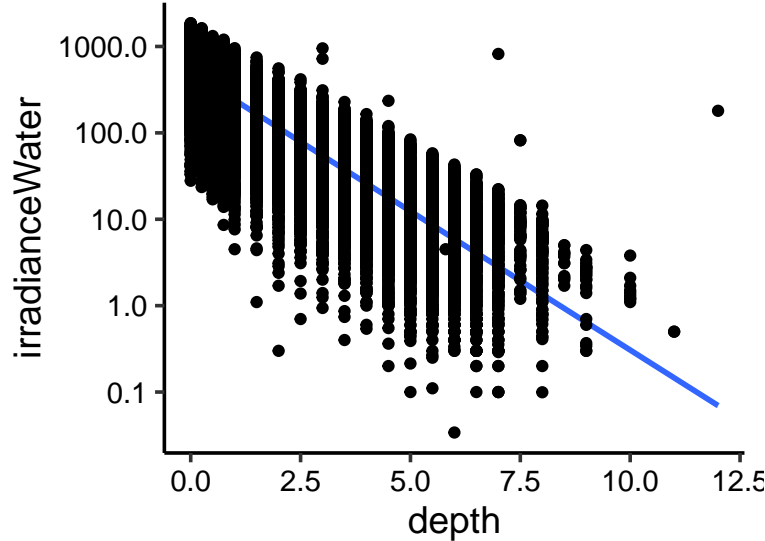


```
par(mfrow = c(1,1))

# Add a line and standard error for the linear regression
```

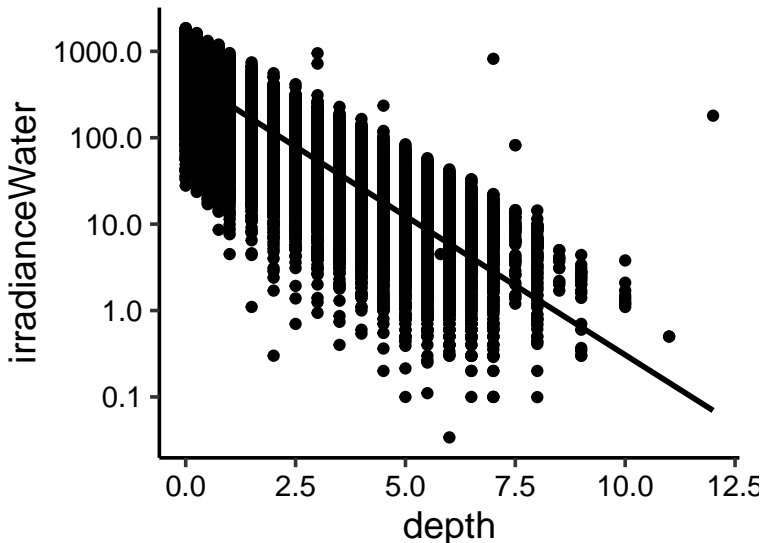
```
#note y-axis transfer to log
irradiancebydepth2 <-
  ggplot(PeterPaul.chem.nutrients, aes(x = depth, y = irradianceWater)) +
  geom_smooth(method = "lm") +
  scale_y_log10() +
  geom_point()
print(irradiancebydepth2)
```

‘geom_smooth()’ using formula = ’y ~ x’



```
# SE - confidence interval around smooth can also be removed
irradiancebydepth2 <-
  ggplot(PeterPaul.chem.nutrients, aes(x = depth, y = irradianceWater)) +
  geom_point() +
  scale_y_log10() + #log transformed values for irradianceWater
  geom_smooth(method = 'lm', se = FALSE, color = "black")
print(irradiancebydepth2)
```

‘geom_smooth()’ using formula = ’y ~ x’



```
# Make the graph attractive
```

Multiple Linear Regression

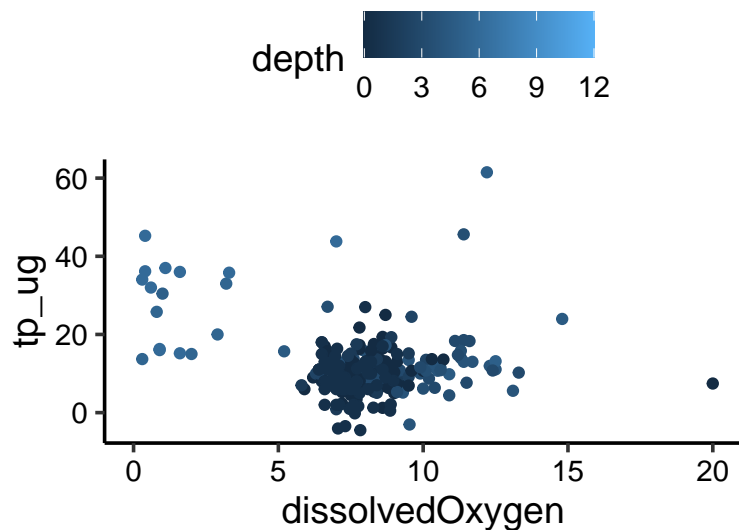
It is possible, and often useful, to consider multiple continuous explanatory variables at a time in a linear regression. For example, total phosphorus concentration in Paul Lake (the unfertilized lake) could be dependent on depth and dissolved oxygen concentration:

```
TPregression <- lm(data = subset(PeterPaul.chem.nutrients, lakename == "Paul Lake"),
                     tp_ug ~ depth + dissolvedOxygen)
summary(TPregression)
```

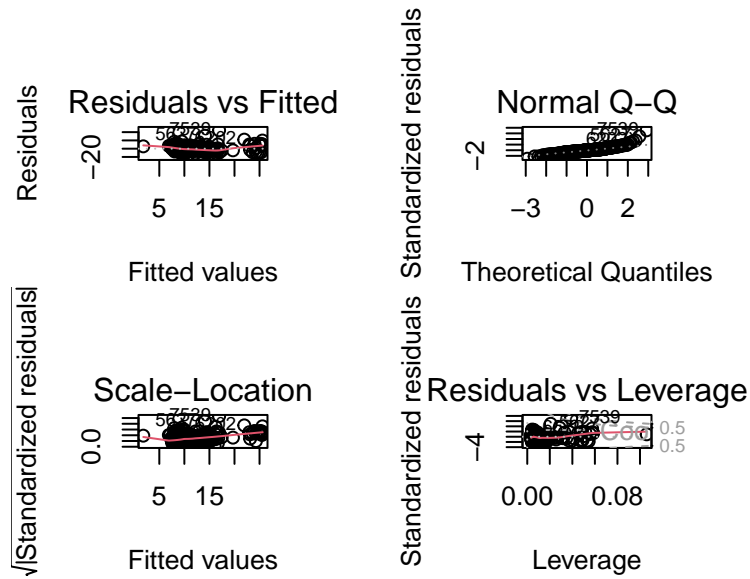
```
##
## Call:
## lm(formula = tp_ug ~ depth + dissolvedOxygen, data = subset(PeterPaul.chem.nutrients,
##   lakename == "Paul Lake"))
##
## Residuals:
##    Min      1Q  Median      3Q     Max
## -19.266 -3.436 -0.534  2.425 44.559
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)
## (Intercept) 12.7869    1.4985  8.533 8.27e-16 ***
## depth       2.1691    0.2222  9.762 < 2e-16 ***
## dissolvedOxygen -0.5494    0.1726 -3.184  0.00161 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 6.405 on 288 degrees of freedom
##   (7283 observations deleted due to missingness)
## Multiple R-squared:  0.2985, Adjusted R-squared:  0.2936
## F-statistic: 61.28 on 2 and 288 DF,  p-value: < 2.2e-16
```

```
TPplot <- ggplot(subset(PeterPaul.chem.nutrients, lakename == "Paul Lake"),
  aes(x = dissolvedOxygen, y = tp_ug, color = depth)) +
  geom_point() +
  xlim(0, 20)
print(TPplot)
```

Warning: Removed 7283 rows containing missing values ('geom_point()').



```
par(mfrow = c(2,2), mar=c(4,4,4,4))
plot(TPregression)
```



```
par(mfrow = c(1,1))
#note, plot1 , what the red line flat.
```

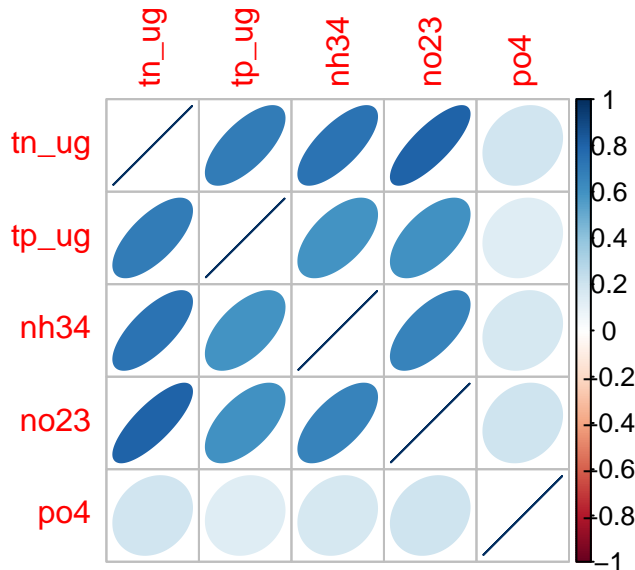
Correlation Plots

We can also make exploratory plots of several continuous data points to determine possible relationships, as well as covariance among explanatory variables.

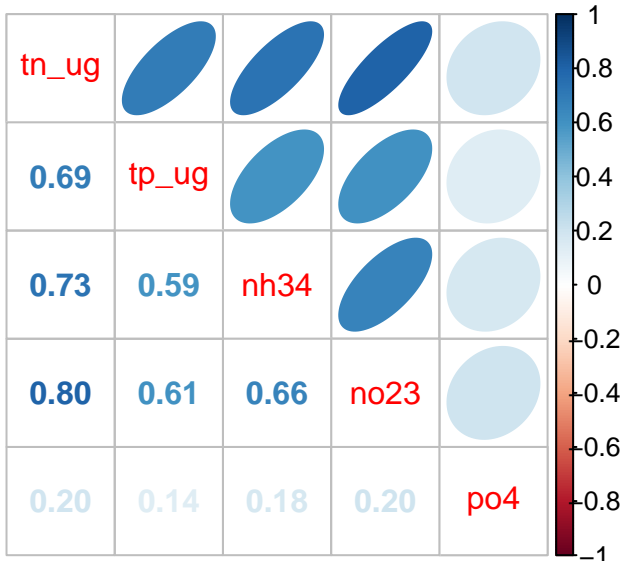
```
#install.packages("corrplot")
#note correlation
library(corrplot)

## corrplot 0.92 loaded

PeterPaulnutrients <-
  PeterPaul.chem.nutrients %>%
  select(tn_ug:po4) %>%
  na.omit()
PeterPaulCorr <- cor(PeterPaulnutrients)
corrplot(PeterPaulCorr, method = "ellipse")
```



```
#note--thin----straight estimation line. color---positive to negative.
corrplot.mixed(PeterPaulCorr, upper = "ellipse")
```



AIC to select variables

However, it is possible to over-parameterize a linear model. Adding additional explanatory variables takes away degrees of freedom, and if explanatory variables co-vary the interpretation can become overly complicated. Remember, an ideal statistical model balances simplicity and explanatory power! To help with this tradeoff, we can use the **Akaike's Information Criterion (AIC)** to compute a stepwise regression that either adds explanatory variables from the bottom up or removes explanatory variables from a full set of suggested options. The smaller the AIC value, the better.

Let's say we want to know which explanatory variables will allow us to best predict total phosphorus concentrations. Potential explanatory variables from the dataset could include depth, dissolved oxygen, temperature, PAR, total N concentration, and phosphate concentration.

```
Paul.naomit <- PeterPaul.chem.nutrients %>%
  filter(lakename == "Paul Lake") %>%
  na.omit()

TPAIC <- lm(data = Paul.naomit, tp_ug ~ depth + dissolvedOxygen +
  temperature_C + tn_ug + po4)

#Choose a model by AIC in a Stepwise Algorithm
step(TPAIC)
```

```
## Start:  AIC=353.95
## tp_ug ~ depth + dissolvedOxygen + temperature_C + tn_ug + po4
##
##          Df Sum of Sq    RSS    AIC
## - po4      1     1.111 2330.8 352.00
## - depth    1    16.739 2346.4 352.76
## - tn_ug    1    30.674 2360.3 353.43
## <none>           2329.7 353.95
## - temperature_C  1   178.398 2508.1 360.29
## - dissolvedOxygen 1   232.194 2561.9 362.68
##
```

```

## Step: AIC=352
## tp_ug ~ depth + dissolvedOxygen + temperature_C + tn_ug
##
##          Df Sum of Sq    RSS    AIC
## - depth      1   15.745 2346.5 350.76
## - tn_ug      1   38.270 2369.1 351.84
## <none>           2330.8 352.00
## - temperature_C  1   194.143 2524.9 359.04
## - dissolvedOxygen 1   250.981 2581.8 361.56
##
## Step: AIC=350.76
## tp_ug ~ dissolvedOxygen + temperature_C + tn_ug
##
##          Df Sum of Sq    RSS    AIC
## <none>           2346.5 350.76
## - tn_ug      1   46.09  2392.6 350.96
## - dissolvedOxygen 1   305.67 2652.2 362.60
## - temperature_C  1   562.60 2909.1 373.05

##
## Call:
## lm(formula = tp_ug ~ dissolvedOxygen + temperature_C + tn_ug,
##      data = Paul.naomit)
##
## Coefficients:
## (Intercept)  dissolvedOxygen  temperature_C          tn_ug
## 23.905781     -0.834405       -0.487603        0.007002

TPmodel <- lm(data = Paul.naomit, tp_ug ~ dissolvedOxygen + temperature_C + tn_ug)
summary(TPmodel)

```

```

##
## Call:
## lm(formula = tp_ug ~ dissolvedOxygen + temperature_C + tn_ug,
##      data = Paul.naomit)
##
## Residuals:
##      Min      1Q      Median      3Q      Max
## -12.9443 -2.8674 -0.4834   2.1666  14.4713
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)
## (Intercept) 23.905781  3.586389  6.666 1.12e-09 ***
## dissolvedOxygen -0.834405  0.221438 -3.768 0.000267 ***
## temperature_C  -0.487603  0.095382 -5.112 1.37e-06 ***
## tn_ug          0.007002  0.004786   1.463 0.146304
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 4.64 on 109 degrees of freedom
## Multiple R-squared:  0.2774, Adjusted R-squared:  0.2575
## F-statistic: 13.95 on 3 and 109 DF,  p-value: 0.00000009208

```

```
##note when check AIC, value, when the value increase with removing, it means the condition become worse  
## note, meanwhile the end of the result aslo shows the Coef
```