ENV 710 – Applied Data Analysis Fall 2014

Lab 9: Correlation and Linear Regression

In this lab, we will acquaint ourselves with correlation and linear regression. Recall that correlation explores the association between two variables, whereas linear regression describes the functional relationship between the explanatory variable and the response variable. Regression analysis is the statistical method used when both the response variable and the explanatory variable are continues variables (i.e. real numbers with decimal places).

The learning goals of the lab are to:

- Recognize when to use correlation or regression
- Practice implementing correlation and regression in R
- Understand how to interpret the output from a correlation and linear regression
- Learn to read the diagnostic tests to evaluate the fit of the model to the data.

At the end of the lab, there are two analyses to conduct. Please type your answers to each problem, including any requested graphs, in a Word document. Submit your answers and your R-code to the class Sakai site under the folder Assignments before 5 pm on either Mon., Nov. 3 (Section 07) or Wed., Nov., 5 (Sections 01 and 02).

More functions in R

In this lab, we introduce a few new R commands.

```
pairs () - Produces a matrix of scatter plots.
cor.test() -Test for association between paired samples, using Pearson's correlation coefficient, Kendall's tau or Spearman's rho.
lm() - Fits a linear model to the data; used for regression and ANOVA (when the independent variable is a factor).
```

Linear Correlation

Download the dataset "TreePlots.csv"), which consists of data from 73 tree plots. The dataset includes a number of variables related to tree inventory, including: Plot (plot ID), mDBH.cm (mean DBH of trees in cm), mBA.cm2 (mean Basal area in cm²), mWD.g.m3 (mean wood density g/m³), mH.m (mean height in meters), AGBH.Mg.ha (above ground biomass in Mg/ha), and Trees (number of trees).

```
tdat <-read.csv("TreePlots.csv", header=T)
attach(tdat)</pre>
```

To get a look at the data, we can dispense with all the histograms and scatter plots

that we have been graphing one-by-one, and use the pairs () command to get a matrix of scatterplots of all the different variables.

```
pairs(tdat)
```

We can spruce up this graph and get even more information on the data, by loading a few functions stored in a script called "pair.fun.R". (Download the script from Sakai.)

Study the plots and you should be able to get a good sense of the relationship between the different variables. In fact, the plot even provides the correlation coefficients for the relationship between each of the variables. We can verify this, by running a correlation on two of the variables.

The data point with the tallest mean tree height seems like it might be having a lot of influence on this correlation. Let's take it out and try again.

```
cor2 <- with(tdat[mH.m<22,], cor.test(mH.m, mWD.g.m3))
with(tdat[mH.m<22,], plot(mH.m, mWD.g.m3))</pre>
```

What is your conclusion? Here we used correlation to check out how it works. But could we have been justified in using linear regression? Why or why not?

What does the output from <code>cor.test()</code> tell you? Can you interpret all the statistics? What is the null hypothesis for the test? Also, look up <code>?cor.test</code> and see what other options exist for correlations.

While cor.test() runs a correlation on two variables and provides tests of significance, you can find the correlation for an entire matrix using cor().

```
cor(tdat)
```

Linear Regression

Now let's do a linear regression on tree height and aboveground biomass. Our hypothesis is that plots with greater mean height of trees have greater biomass. (You don't have to be a genius to figure that out, but let's proceed anyway.)

```
Analysis of Variance Table

Response: AGBH.Mg.ha

Df Sum Sq Mean Sq F value Pr(>F)

mH.m 1 448405 448405 46.339 2.645e-09 ***

Residuals 71 687046 9677

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

First, let's look at the ANOVA printout on the model since this should be familiar. The small p-value tells us that our regression model explains a significant part of the variation in the data. It tests our alternative hypothesis (tree height is related to biomass) against the null hypothesis (tree height has no effect on biomass), and demonstrates that by incorporating tree height, $\beta_1 X_i$, we can explain a significant part of the variation in above ground biomass, Y_i .

$$H_0: Y_i = \beta_0 + \varepsilon_i$$

$$H_a: Y_i = \beta_0 + \beta_1 X_i + \varepsilon_i$$

If we had additional independent variables (multiple regression), then it would test: $H_0: \beta_1 = \beta_2 = 0$ vs. $H_a: at least one of <math>\beta_1 and \beta_2 \neq 0$.

In a simple model like this, the ANOVA table does not provide much information different from 1m () output. However, ANOVA tables can be useful for comparing more sophisticated models. Let's move onto the rest of the output.

```
summary(lm1)
```

```
Call:
lm(formula = AGBH.Mg.ha ~ mH.m, data = tdat)
Residuals:
                 Median
    Min
             10
                               30
                                       Max
-242.031 -41.889
                           56.276 283.406
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -499.926 114.246 -4.376 4.07e-05 ***
                               6.807 2.64e-09 ***
mH.m
             45.704
                        6.714
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 98.37 on 71 degrees of freedom
Multiple R-squared: 0.3949, Adjusted R-squared:
F-statistic: 46.34 on 1 and 71 DF, p-value: 2.645e-09
```

Let's work through this piece by piece.

Call repeats the function call to the lm() function.

Residuals displays a five-number summary of the model residuals. This is the output obtained from applying the quantile () function to the residuals. The residuals are defined as $e_i = y_i - \hat{y}_i$ where y_i is the observed value and \hat{y}_i is the predicted value (from the model) of the response variable for that observation. For example, take the case of Plot 1, where the observed biomass (AGBH.Mg.ha) is 157.45 and the observed mean height (mH.m) is 14.90. Plug the height value into our regression formula:

$$\hat{y}_1 = -499.926 + 45.704(14.90)$$

results in $\hat{y}_1 = 181.09$. And, the residual would be $e_i = 181.09 - 157.45 = -23.64$. To get all the residuals for our model, type: lm1\$residuals.

Coefficients contains the parameter estimates from the regression model. So the estimated model (rounded to 2 decimal places) is:

$$\hat{y}_i = -499.93 + 45.70 \times mH. m_i$$

We can use the equation to calculate the fitted values – the values for an output variable that have been predicted by a model fitted to a set of data.

```
y.fitted <- lm1$coefficient[1] + lm1$coefficient[2]*mH.m</pre>
```

This should produce the same numbers as lm1\$fitted. To check that they are the same, let's run a correlation between them:

```
cor.test(y.fitted, lm1$fitted)
```

What is your conclusion?

Because the coefficient, *mH.m*, is positive the model predicts that aboveground biomass will increase as tree height increases. Every meter in additional height will lead to a 45.70 Mg/ha increase in biomass. The intercept in this model is not interpretable. Technically it represents the biomass of a plot that is predicted when the average height of trees is 0. The model predicts a negative biomass when trees have a height of 0, which is nonsensical. In discussing the intercept we are extrapolating beyond the range of data that are possible, one of the no-no's in regression analysis.

The rest of the display includes the standard deviation of the estimates, and the t statistic and corresponding p-value for the null hypothesis that the true value of each coefficient is 0.

The individual *t* tests are variable-added-last tests. They test whether the coefficient of the given variable is significantly different from 0, in the context of a model that already contains the rest of the independent variables. These tests are not tests of the importance of a variable by itself, but only in the context of the rest of the model. More about this when we discuss whether to include a variable in a model or not.

Residual Standard Error, $\hat{\sigma}$, is the square root of the sum of the squared residuals divided by their degrees of freedom, where n is the number of observations and p is the number of estimated regression parameters (73-2 = 71, in this example).

$$\hat{\sigma} = \sqrt{\frac{\sum_{i=1}^{n} e_i^2}{n-p}}$$

When the residual standard error is 0, then the model fits the data perfectly (likely due to overfitting). If the residual standard error cannot be shown to be significantly different from the variability in the null model, then there is little evidence that the linear model has any predictive ability.

Multiple R-squared is the coefficient of determination: the proportion of the variation in the response that is explained by the regression. Here 39.5% of the variation in aboveground biomass across the plots is explained by its linear relationship to mean tree height. R² compares the amount of unexplained variation before and after a regression model is fit. The before variation is the sum of squares total (SST).

$$SST = \sum_{i=1}^{n} (y_i - \bar{y})^2$$

Having fit the regression, the model we obtain replaces the sample mean as the best predictor of an individual observation.

Adjusted R-square introduces a penalty term for each regressor (independent variable) in the R² calculation. Without this adjustment, R² will usually increase (at least a little) with each additional regressor; and thus we will always end up choosing the most complicated model as the best model. The penalty in the adjusted R² is chosen to yield an unbiased estimate of the population R², and works along the same principal as the AIC, which we will talk about later.

The rest of the output (*F* statistic, *p*-value) summarizes the ANOVA results from above.

Reporting results from linear regression

For the above example, we might say: Average tree height in forest plots significantly increase plot-level biomass, with a 1 m increase in average tree height resulting in a 45.70 Mg/ha increase in plot biomass ($R^2 = 0.386$, $F_{1.71} = 46.34$, p < 0.001).

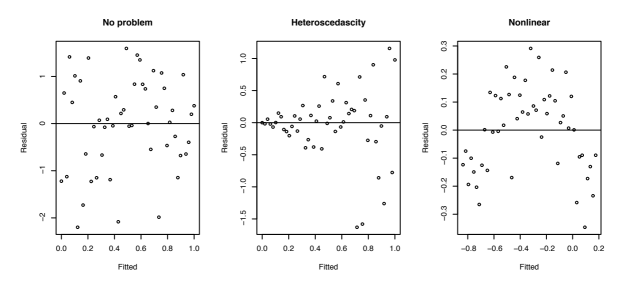
In some cases, you may want to present the entire equation: Average forest biomass is significantly related to average tree height (biomass = -499.9 + 45.70 x height, R^2 = 0.386, $F_{1,71}$ = 46.34, p < 0.001).

Checking the fit of the model

When conducting any statistical analysis it is important to evaluate (i) how well the model fits the data and (ii) that the data meet the assumptions of the model. There are numerous ways to do this and a variety of statistical tests to evaluate deviations from model assumptions. Generally statisticians examine various diagnostic plots after running their regression models. See the below description (which is exactly the same as the description from last week's lab) of the diagnostic tests.

```
par (mfrow=c(2,2))
plot(lm1)
```

Plotting of the model provides four diagnostic plots. The first is plot of the residuals (distance of the data points from the fitted regression line) versus the fitted data. Points should be randomly scattered around the centerline. Any pattern indicates either violation of linearity or homoscedasticity.



The second plot is a q-q plot, which we have already used to evaluate the normality of a variable. Significant departures from the line suggest violations of normality. If the pattern were S-shaped or banana shaped, we would need a different model. You can also perform a Shapiro-Wilk test of normality with the ${\tt shapiro.test}()$

function, but be careful...

Against my better judgment, in the past we have used the shapiro.test() to assess normality. Remember that no test will show that your data has a normal distribution. Normality statistics show when your data is sufficiently inconsistent with a normal distribution that you would reject the null hypothesis of "no difference from a normal distribution". However, when the sample size is small, even big departures from normality are not detected, and when the sample size is large, even the smallest deviation from normality will lead to a rejected null. In other words, if we have enough data to fail a normality test, we always will because real-world data won't be clean enough. See (http://www.r-bloggers.com/normality-and-testing-for-normality/) for an example with simulated data. So, where does that leave us? Explore your data for large deviations from normality and make sure to assess heteroscedasticity and outliers. But, don't get hung up on whether your data are normally distributed or not. As the author of the above link suggests: "When evaluating and summarizing data, rely mainly on your brain and use statistics to catch really big errors in judgment."

The third plot is a plot of standardized residuals versus the fitted values. It repeats the first plot, but on a different scale. It shows the square root of the standardized residuals (where all the residuals are positive). If there was a problem, the points would be distributed inside a triangular shape, with the scatter of the residuals increasing as the fitted values increase.

The fourth plot is a residuals-leverage plots that shows the Cook's distance for each of the observed values. Cook's distance measures relative change in the coefficients as each replicate is deleted. So the point is to highlight those y_i (response) values that have the biggest effect on parameter estimates. The idea is to verify that no single data point is so influential that leaving it out changes the structure of the model. The potential trouble points are labeled.

In the diagnostic plots for our model, both the Scale-Location and Residuals vs. Leverage plots show points scattered away from the center, suggesting that some points have excessive leverage. Another pattern is that points 25, 43, and 63 stick out in nearly every plot. This warns us that something could be odd with those observations. We might want to redo the analysis without those plots and see if our inference changes.

```
lm2 < - with(tdat[-c(25,43,63), ], lm(AGBH.Mg.ha ~ mH.m))
```

Taking out these observations actually improves our R² and doesn't change the estimate for the effect of mean tree height, mH.m, much at all.

We can plot the regression line, this time getting fancier with ggplot2.

```
require(ggplot2)
tdat1 <- tdat[-c(25,43,63), ]
ggplot(tdat1, aes(x = mH.m, y = AGBH.Mg.ha)) +
        geom_point(shape = 1) +
        geom_smooth(method = lm) +</pre>
```

```
xlab("Mean height, m") +
ylab(expression(paste("Biomass Mg ", ha^-1)))
```

<code>ggplot2</code> offers a lot of options and makes attractive graphs, but feel free to stick with the standard plotting procedures that we have used up to now. Note that using $geom_smooth$ produces a 95% confidence region. This can be suppressed by revising the above to read: $geom_smooth$ (method = lm, se = F).

Your assignment

Your assignment is to conduct two different linear regressions on the "TreePlots.csv" data: (1) mean tree diameter (mDBH.cm) versus plot biomass (AGBH.Mg.ha), and (2) mean height (mH.m) versus mean wood density (mWD.g.m3). In the first regression, plot biomass should be your dependent variable; in the second regression, mean height should be your dependent variable.

See my above comment about normality, and don't sweat departure from normality too much.

For each regression, write a 1-page description of your analysis, results, and inference. Each write-up should include the following information:

- Null and alternative hypotheses of your tests
- Results of your statistical test, interpreting your test in 2-3 sentences that include the appropriate reporting of the statistics
- An interpretation of the regression model (equation) from each analysis (e.g. how does plot biomass and mean height vary with different levels of tree diameter and wood density)
- A description of how you checked the assumptions of your statistical test
- An interpretation of diagnostic figures