Linear Models with R

Emphasis on 2-IV models: Basics of Multiple Regression

Bruce Dudek

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Contents

Pı	eface	2
1	Introduction and Goals 1.1 A note on the R Programming environment	2 3
2	Import the Data Set and Do Numerical and Graphical EDA 2.1 Read the data	4 4 5 12 21
3	Bivariate Correlation and Linear Regression 3.1 Bivariate computations: Covariances and Pearson product-moment correlations 3.2 Test correlations with the corr.test function 3.3 Simple regressions of Salary on each of the two IVs	21 22 23 24
4	Implementation of Standard Multiple Regression Analyses: Linear modeling with multiple IVs 4.1 The core linear modeling approach with R 4.2 Diagnostic and Added-variable plots for the full two-IV model	30 31 39
5	Review Unique and common proportions of variance and Type I vs III Sums of Squares5.1SS components, semi-partial correlations, unique and common proportions/SS5.2F tests on the unique components?5.3Compare and Contrast information from fit3 and fit4	3 43 43 45 46
6	Formally Comparing Models	48
7	Further work with residuals and yhats	49
8	Examine the fit of the plane in a 3D wireframe plot:	52
9	Inferential tests for assumptions 9.1 Evaluation of the Residual Normality Assumption 9.2 Inferential test of Homoscedasticity 9.3 Global Evaluation of Linear Model Assumptions using the 'gvlma' package.	54 54 57 59
10	Further capabilities in R: Assumptions, Diagnostics, Model Criticism	60

11 Regression diagnostics and Influence Analysis	61
11.1 A set of standard influence and diagnostic statistics	61
12 Resampling and Robust methods for Linear Models	68
12.1 Ordinary nonparametric bootstrapping with the boot function	68
12.2 An alternative bootstrap approach from car package functions	75
12.3 Robust Regression for heteroscedastic data sets	79
12.4 Robust regression methods when outliers or highly influential cases are present	82
13 Extensions to larger models and to categorical IVs	82
13.1 Evaluate a 3-IV model	82
13.2 Categorical IVs	87
14 Efficiency in OLS analysis using the olsrr package	90
14.1 A basic analysis	90
14.2 Collinearity diagnostics	90
14.3 Added-variable plots	91
14.4 Residual Assumptions	92
14.5 Plots for examination of Influence	97
15 Bayes Factor approach to multiple regression	102
15.1 Basic approaches with Bayes Factors	103
15.2 An admonition regarding Bayesian Inference	106
16 Documentation for Reproducibility	106
16.1 Revision History	108

Preface

Linear Modeling, in it's most rudimentary form is also termed multiple regression. This document provides a template for execution of most of the basic analyses associated with this methodology. It is intended for students of the APSY510/511 introductory statistics sequence at the University at Albany, but can be a standalone document for others learning to use R for data analysis. The level of the document is targeted to an audience of researchers in training who are simultaneously learning linear modeling/regression theory and R programming. Some introduction to regression modeling with the 1m function previously had been covered for simple regression and can be found in an accompanying document. The current document extends that work to a model with two IVs as an extensive illustration and then briefly covers models with more than two IVs. The document emphasizes models where all variables are numeric/quantitative. Categorical IVs are covered in later documents, although one brief illustration is included in this document. Some extension to models with more than two IVs is also included in the "extensions" chapter. Using the **bookdown** suite of tools permits organization of the document into chapters.

This book/monograph uses the **bookdown** package (Xie, 2020a) for R (R Core Team, 2020), which was built on top of **rmarkdown** (Allaire et al., 2020) and **knitr** (Xie, 2015). RStudio (RStudio Team, 2015) was used for all writing and programming.

1 Introduction and Goals

This document lays out basic strategies for linear modeling in R. It is structured as a reflection of what, in Social, Behavioral, and Life Sciences is called Multiple Regression. It approaches linear modeling where one outcome variable (dependent variable) is predicted from multiple independent variables. Each variable is a quantitative measurement. The document presumes a background in the basics of multiple regression from both equational and conceptual perspectives, including the roles of partial and semi-partial correlation. The document is intended for students in the APSY510/511 course sequence at the University at Albany, but can be more generally applicable.

For the most part, the approach here will use a multiple regression problem with only two independent variables. This permits comparability for a companion presentation on implementation of multiple regression with SPSS that used the same data set. The data set is the Cohen, Cohen, West and Aiken textbook chapter 3 example on faculty salaries, publications and citations (Cohen et al., 2003).

The flow of the document moves from univariate description, to bivariate description, to all aspects of linear modeling. It extends to topics of evaluation of assumptions for inferential tests, influence analysis, and model criticism.

The emphasis is narrowly placed on implementation of the standard methods in R. Conceptual rationales for the approach, alternative methods (e.g., bayesian), variable selection strategies, model building, and the role of regression in causality assessment are treated elsewhere although a brief introduction to Bayes Factors is included here. The document does not address nonparametric or semi-parametric regression such as quantile regression nor does it address curvilinear regression.

1.1 A note on the R Programming environment

All of the analyses and graphical displays found in this document were produced in R. Usually, the document shows the relevant R code for each topic. The purpose of the document is has a primary focus on the how-to in R, but also emphasizes the conceptual progression related to understanding linear modeling in the simplest of multiple regression applications, the two-IV model. The document can be an extensive template for R usage in these types of analyses. Some extension to models with larger numbers of IVs is also included. To that end, all the code is available both in this document and one other source. In the spirit of reproducible and open source research, this document was created in **rmarkdown** and **bookdown**. The Rmd file contains ALL of the R code required to reproduce the analyses and figures contained in the document.

Graphs are drawn with **ggplot2**, base system graphics, and a convenient 3D surface plotting capability from the **plot3D** package. Analyses are extensively reliant on the base system 1m function. Additional analyses use other packages and BCD-created functions introduced as the document progresses.

Several packages are required for the work in this document.

```
library(BayesFactor)
library(bcdstats)
library(boot)
library(broom)
library(car)
library(GGally)
library(ggExtra)
library(ggplot2)
library(ggthemes)
library(grid)
library(gvlma)
library(gt)
library(HH)
library(knitr)
library(lattice)
library(lmtest)
library(MASS)
library(moments)
library(nortest)
library(olsrr)
library(psych)
library(plot3D)
```

library(plot3Drgl)
library(plyr)
library(rcompanion)
library(rmarkdown)
library(sandwich)
library(tseries)
library(UsingR)
library(yhat)

Package citations for packages loaded here (in the above order): BayesFactor (Morey and Rouder, 2018), bcdstats (Dudek, 2020), boot (Canty and Ripley, 2019), broom (Robinson and Hayes, 2020), car (Fox et al., 2020), GGally (Schloerke et al., 2020), ggExtra (Attali and Baker, 2019), ggplot2 (Wickham et al., 2020), ggthemes (Arnold, 2019), grid (Auguie, 2017), gvlma (Pena and Slate, 2019), gt (Iannone et al., 2019), HH (Heiberger, 2020), knitr (Xie, 2020b), lattice (Sarkar, 2020), lmtest (Hothorn et al., 2019), MASS (Ripley, 2019), moments (Komsta and Novomestky, 2015), nortest (Gross and Ligges, 2015), olsrr (Hebbali, 2020), psych (Revelle, 2020), plot3D (Soetaert, 2019), plot3Drgl, (Soetaert, 2016), plyr (Wickham, 2020), rcompanion (Mangiafico, 2020), rmarkdown (Allaire et al., 2020), sandwich (Zeileis and Lumley, 2019), tseries (Trapletti and Hornik, 2019), UsingR (Trapletti and Hornik, 2019), yhat (Nimon et al., 2013)

Package citations for packages loaded elsewhere in this document: **bookdown** (Xie, 2020a)

The **bcdstats** package can be installed with instructions found at its github repository:

https://github.com/bcdudek/bcdstats

2 Import the Data Set and Do Numerical and Graphical EDA

Cohen, Cohen, West and Aiken (2003) have presented a data set where, ostensibly, faculty salaries are predicted from several IVs. For the detailed example in this document two IVS are used. Both are quantitative: number of publications (pubs) and number of citations (cits). In this illustration all three variables are quantitative and can be thought of as measured on ratio scales (although pubs and cits are integer). The data are in a .csv file. It contains the three variables of interest plus three more. A subject/case number variable, time since PhD degree awarded, and sex. The file is "cohen.csv". The elaboration of this two-IV example is intended to parallel and extend the implementation previously done with SPSS for the same variables.

2.1 Read the data

The data file is read and a data frame is produced in the typical manner for .csv files that contain a header row with variable names. We also need to establish that our quantitative variables are read, by R, as numeric variables. We can examine the data frame with the 'View' function but I just show the first chunk of the data frame with a screen capture here. Later in this document the additional IVs, degree_yrs, and gender will be considered, but most analyses just use pubs and cits.

Finally, the data frame is "attached" so that easier naming of variables can be accomplished in the many of the initial R functions to be employed. Although this is often not a recommended approach in R, we will not encounter the downside in this document. The data frame will be "detached" prior to use in lm functions.

```
cohen1 <- read.csv("data/cohen.csv", stringsAsFactors=T)
cohen1$degree_yrs <- as.numeric(cohen1$degree_yrs)
cohen1$pubs <- as.numeric(cohen1$pubs)
cohen1$cits <- as.numeric(cohen1$cits)
cohen1$salary <- as.numeric(cohen1$salary)
#View(cohen1)
attach(cohen1)</pre>
```

	case 🔅	$degree_yr\hat{s}$	pubs 🍦	cits 🗦 🗘	salary $\hat{\circ}$	$\mathbf{gender}^{\updownarrow}$
1	1	3	18	50	51876	female
2	2	6	3	26	54511	female
3	3	3	2	50	53425	female
4	4	8	17	34	61863	male
5	5	9	11	41	52926	female
6	6	6	6	37	47034	male
7	7	16	38	48	66432	male
8	8	10	48	56	61100	male
9	9	2	9	19	41934	male
10	10	5	22	29	47454	male

Here is a screen capture of what the first few lines of the data frame look like:

2.2 Graphical and Numerical Description of the Variables

The best practices strategy we have put in place prioritizes exploratory data analysis at the outset. "Get to know your Data." This involves graphical and numerical summaries. The exact approaches taken here are somewhat arbitrary. There are many ways to draw useful graphs in R and an equally large number of ways to obtain numerical descriptive information. The functions used here are similar or identical to those used for previous topics in the course, beginning with the location test problems and progressing through simple regression and bivariate correlation.

2.2.1 Univariate Distributional Displays for each of the three Variables, salary, publications, and citations

Each of our three variables is examined here in several ways. First, I have used base system graphics to plot a few standard exploratory graphs. The basic set of four figures includes the frequency histogram with a kernel density function overlaid, a qq normal probability plot, box plot, and a violinplot.

Salary, the DV is first.

```
#win.graph(8,6) # use quartz() on MAC OS
layout(matrix(c(1,2,3,4),2,2)) #optional 4 graphs/page
hist(salary,probability=TRUE,breaks=6,
    ylim= c(0, 1.2*max(density(salary)$y)),
    main="Histogram/Density Salary",
    cex.main=.7)
lines(density(salary),col= "red",lwd = 3)
qqnorm(salary,main="Normal Probability Plot of Salary",
    cex.main=.7);qqline(salary)
boxplot(salary,xlab="Salary",ylab="")
violinplot(salary,col="cornsilk",names="Salary")
```

Histogram/Density Salary







Normal Probability Plot of Salary



Theoretical Quantiles



Then, publications and citations, the IV's.

```
#win.graph(8,6) # use quartz() on MAC OS
layout(matrix(c(1,2,3,4),2,2)) #optional 4 graphs/page
hist(pubs,probability=TRUE,
    ylim= c(0, 1.5*max(density(pubs)$y)),
    main="Histogram/Density of Publications",
    cex.main=.7)
lines(density(pubs),col = "red",lwd = 3)
qqnorm(pubs,main="Normal Probability Plot of Publications",
    cex.main=.7);qqline(pubs)
boxplot(pubs,xlab="Publications",ylab="")
violinplot(pubs,col="cornsilk",names="Publications")
```

Histogram/Density of Publications





```
Publications
```

Normal Probability Plot of Publications



Theoretical Quantiles



```
#win.graph(8,6) # use quartz() on MAC OS
layout(matrix(c(1,2,3,4),2,2)) #optional 4 graphs/page
hist(cits,probability=TRUE,
    ylim= c(0, 1.5*max(density(cits)$y)),
    main="Histogram/Density of Citations",
    cex.main=.7)
lines(density(cits),col = "red",lwd = 3)
qqnorm(cits,main="Normal Probability Plot of Citations",
    cex.main=.7);qqline(cits)
boxplot(cits,xlab="Citations",ylab="")
violinplot(cits,col="cornsilk",names="Citations")
```







Citations





Theoretical Quantiles



An alternative that is a bit more efficient is to use the 'explore' function in the 'bcdstats' package. This produces not only graphs, but also numeric summaries of the variables.

explore(salary)



vars n mean sd median trimmed mad min max range skew
X1 1 62 54815.76 9706.02 53681 54226.1 9119.47 37939 83503 45564 0.64
kurtosis se
X1 0.5 1232.67

Univariate Plots: pubs

Histogram with Kernel Densi Histogram with Normal Curv Boxplot with Jittered Rug





Kernel Bandwidth based on Wand, 19



Univariate Plots: cits

Histogram with Kernel Densi Histogram with Normal Curv Boxplot with Jittered Rug





##		vars	n	mean	sd	median	trimmed	mad	min	\max	range	skew	kurtosis	se
##	X1	1	62	40.23	17.17	35	39.08	14.08	1	90	89	0.68	0.57	2.18

If you don't want to use the 'explore' function from 'bcdstats', then you can still easily obtain numerical summaries with the 'describe' function from the 'psych' package as we have done in prior work, and perhaps just go with the earlier graphs instead of the six panel figure from explore.

```
# our three variables are the third through fifth in the data frame
psych::describe(cohen1[,c("salary","pubs","cits")], skew=T, type=2)
```

mad## sd median trimmed max range skew vars n mean min 53681 54226.10 9119.47 37939 83503 45564 0.64 ## salary 1 62 54815.76 9706.02 ## pubs 2 62 18.18 14.00 13 16.28 10.38 1 69 68 1.38 3 62 35 39.08 ## cits 40.23 17.17 14.08 1 90 89 0.68 ## kurtosis se 0.50 1232.67 ## salary ## pubs 2.01 1.78 ## cits 0.57 2.18

2.2.2 Summary of Univariate EDA

All three variables have some degree of positive skewness, pubs more than cits and salary. This may mean that the residual normality assumption for the linear models is not satisfied. We will look carefully at that assumption following those analyses.

2.3 Bivariate Relationships among the three variables

The most important starting point for evaluation of a linear models regression system is visualization of the zero-order bivariate relationships. We do this with scatter plots. The two sections here do this either with base system graphics or ggplot2 in a more sophisticated and perhaps more publication ready format.

Readers are reminded that we earlier examined numerous ways of drawing bivariate scatter plots with R. This was done in the R implementation section when we considered simple regression. It included a base system graphics approach to what I thought might be a publication quality figure. In the present document, I present simple approaches to obtaining maximum information in an exploratory data analysis (EDA) type of approach where speed is important and the investigator is at initial stages of analysis.

2.3.1 Bivariate Scatterplots and SPLOM with base system functions

Initially, we can draw bivariate scatter plots for all three pairs of variables. Using base system graphics, I have done this in a more complicated fashion by adding boxplots to the margins for each variable

```
par(fig=c(0,0.8,0,0.8))
plot(cits, salary, xlab="Citations", ylab="Salary")
abline(lm(salary~cits), col="red") # regression line (y~x)
par(fig=c(0,0.8,0.55,1), new=TRUE)
boxplot(cits, horizontal=TRUE, axes=FALSE)
par(fig=c(0.65,1,0,0.8),new=TRUE)
boxplot(salary,axes=FALSE)
mtext("Scatterplot & Linear Regression\n Plus Univariate Boxplots", side=3, outer=TRUE, line=-3)
```



```
par(fig=c(0,0.8,0,0.8))
plot(pubs, salary, xlab="Publications",
    ylab="Salary")
abline(lm(salary~pubs), col="red") # regression line (y~x)
par(fig=c(0,0.8,0.55,1), new=TRUE)
boxplot(pubs, horizontal=TRUE, axes=FALSE)
par(fig=c(0.65,1,0,0.8), new=TRUE)
boxplot(salary,axes=FALSE)
mtext("Scatterplot & Linear Regression Plus Univariate Boxplots", side=3, outer=TRUE, line=-3)
```

Scatterplot & Linear Regression Plus Univariate Boxplots



```
par(fig=c(0,0.8,0,0.8))
plot(pubs, cits, xlab="Publications",
    ylab="Citations")
abline(lm(cits~pubs), col="red") # regression line (y~x)
par(fig=c(0,0.8,0.55,1), new=TRUE)
boxplot(pubs, horizontal=TRUE, axes=FALSE)
par(fig=c(0.65,1,0,0.8),new=TRUE)
boxplot(cits,axes=FALSE)
mtext("Scatterplot & Linear Regression Plus Univariate Boxplots", side=3, outer=TRUE, line=-3)
```

Scatterplot & Linear Regression Plus Univariate Boxplots



A preferred approach for the generation of bivariate scatter plots is creation of a scatter plot matrix (SPLOM). There are many ways of doing this in R. Here I use the **pairs.panels** function. I also permitted the 'pairs.panels' function from the **psych** package to include the time variable so that a sense of how it handles larger numbers of variables. Note the "subsetting" syntax for asking to leave out the first and sixth variables from the data frame (case number and gender)

pairs.panels(cohen1[c(-1,-6)])



The pairs.panels function creates a SPLOM with quite a few attributes beyond the simple bivariate scatter plots. The frequency histograms and kernel density functions as well as the display of the Pearson product-moment zero-order correlation is obvious. However, the added elements on the scatter plots require explanation. Rather than plotting a linear regression function, by default, pairs.panels draws a loess fit line. The ellipse that is drawn is based on the 'ellipse' function from John Fox's 'car' package and is implemented by default in pairs.panels. It is called a correlation ellipse and gives a sense of the correlation at limits of one std deviation above and below the mean for each variable. Each of these latter two graph elements can be controlled with arguments passed to 'pairs.panels'. For example, instead of a loess fit, a linear regression line can be chosen. See the help for pairs.panels for more information (?pairs.panels)

2.3.2 Bivariate Scatterplots and SPLOM with 'ggplot2' and 'lattice'

As a reminder of basic work we did with the introduction to using R for simple regression, bivariate scatter plots are drawn here with the ggplot function fromg the **gplot2* package. This set of plots is intended to give a sense of different kinds of information display that is possible. The plots are not refined to a publication quality level, and I only take space for the salary-pubs pair of variables. The reader can extend beyond this basic set with only salary and pubs by changing the variables

```
# the base plot:
#create base graph entity wthout displaying anything
pbase <- ggplot(cohen1, aes(x=pubs,y=salary))
#display the scatterplot with the data points
pbase + geom_point()
```







or map the height of the density curve to a color progression using the "level" specification
pbase + stat_density2d(aes(fill=..density..), geom="raster",contour=FALSE)



or, do it with points and map density a different way
pbase + geom_point() +



Two more ways to obtain a SPLOM are illustrated below. First, we can use the **splom** function from the **lattice** package. Note the exact specification required for requesting the subset of variables from the cohen data frame, and the tilda model symbol used to specify that data frame, and the subsetting to request the same four variables as above.

splom(~cohen1[2:5])



I have never found that **splom** function to yield a useful or appealing figure. An alternative, and easily implemented, function in the **GGally** package is called **ggpairs**. It is based on ggplot2 and is both efficient and helpful (but I probably still prefer 'pairs.panels')

ggpairs(cohen1[2:5], progress=F)



Each of these SPLOMS can be modified/extended to provide more useful information and to tailor the appearance.

2.4 Summary of EDA

The EDA approaches outlined above are views as an essential starting point to any more advanced modeling and inference. "Get to Know Your Data!" It is particularly important to be aware of univariate or bivariate outliers and non-normal distributions of variables. These issues can have major impacts on the ability of the linear modeling results to be accurate, generalizable, and useful. Our three primary variables (salary, pubs and cits) all appeared to have some degree of positive skewness, with pubs having the most. Pubs has an outlier that contributes to this skewness. It will be interesting to examine the residuals-based assumptions for our linear modeling when we know that non-normality in a univariate sense is present. The bivariate EDA approaches did not give any strong sense of multi-dimensional outliers, so this data set set may not have any particularly influential individual cases.

3 Bivariate Correlation and Linear Regression

These bivariate analyses are done largely to reiterate the approaches we established earlier for two variable systems (i.e., simple regression and bivariate correlation). The R implementations are identical to what was covered earlier, except that with more variables, there are more pairs and more than one simple regression possible.

This review established some important information to compare to the approaches we took in SPSS, and allows us to remember the meaning of a simple regression regression coefficient and compare its value to coefficients found for the same variables in the multiple regression section of this document that follows this bivariate section. The sequence of our approach here is to:

- a. obtain the variance-covariance matrix for our three variables.
- b. obtain the zero-order bivarate correlation matrix (Pearson product-moment correlations).
- c. and to test the null hypotheses that the three population correlations are zero.
- d. obtain the two different simple regressions of salary on each of the two IVs.
- e. graphically evaluate the assumptions of inferences in simple regression.

3.1 Bivariate computations: Covariances and Pearson product-moment correlations

Analyses requested here repeat/extend the implementation approach we learned with R for simple regression.

First, obtain variance-covariance and correlation matrices for the three variables of interest. Recall that we can submit a whole data frame or matrix of data to cov and cor. That is done here along with subsetting the data frame to use only the three variables for our defined analyses.

```
cat(paste("cov produces a matrix with variances on the leading diagonal
and covariances in the upper and lower triangles.
```

"))

```
## cov produces a matrix with variances on the leading diagonal
## and covariances in the upper and lower triangles.
```

cov(cohen1[,3:5])

```
##
                pubs
                            cits
                                      salary
## pubs
            196.1155
                         80.1724
                                    68797.68
## cits
             80.1724
                        294.8662
                                    91628.81
## salary 68797.6830 91628.8096 94206882.35
corr1 <- cor(cohen1[,3:5], use="complete.obs", method="pearson")</pre>
cat(paste("
cor produces a symmetrical matrix of Pearson correlation coefficients
Here, they are rounded to two places.
```

```
"))
```

##
cor produces a symmetrical matrix of Pearson correlation coefficients
Here, they are rounded to two places.
round(corr1,2)

pubs cits salary
pubs 1.00 0.33 0.51
cits 0.33 1.00 0.55
salary 0.51 0.55 1.00

Now we will test each of the three zero-order correlations with the standard t-test and n-2 df:

```
## Pearson's product-moment correlation
##
## data: pubs and salary
## t = 4.5459, df = 60, p-value = 2.706e-05
```

```
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
##
  0.2934802 0.6710778
## sample estimates:
##
         cor
## 0.5061468
cor.test(pubs,cits, method = "pearson", alternative = "two.sided",
         exact = FALSE)
##
##
   Pearson's product-moment correlation
##
## data: pubs and cits
## t = 2.7392, df = 60, p-value = 0.008098
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
  0.09122073 0.53833356
##
## sample estimates:
##
         cor
## 0.3333929
cor.test(cits,salary, method = "pearson", alternative = "two.sided",
         exact = FALSE)
##
   Pearson's product-moment correlation
##
##
## data: cits and salary
## t = 5.098, df = 60, p-value = 3.69e-06
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
  0.3477491 0.7030024
##
## sample estimates:
##
         cor
## 0.5497664
```

3.2 Test correlations with the corr.test function

A more efficient way of testing correlations from a whole correlation matrix is to use the corr.test and corr.p functions from the **psych** package. This method also provides the ability to adjust p values for multiple testing (using the Holm method), and to produce confidence intervals.

```
ct1 <- psych::corr.test(cohen1[,3:5])</pre>
print(psych::corr.p(ct1$r, n=62), short=FALSE)
## Call:psych::corr.p(r = ct1$r, n = 62)
## Correlation matrix
##
          pubs cits salary
## pubs
          1.00 0.33
                      0.51
          0.33 1.00
                      0.55
## cits
## salary 0.51 0.55
                      1.00
## Sample Size
## [1] 62
## Probability values (Entries above the diagonal are adjusted for multiple tests.)
##
          pubs cits salary
          0.00 0.01
## pubs
                          0
```

```
## cits 0.01 0.00
                        0
## salary 0.00 0.00
                        0
##
   Confidence intervals based upon normal theory. To get bootstrapped values, try cor.ci
##
##
             lower
                      r upper
                                 р
              0.09 0.33 0.54 0.01
## pubs-cits
## pubs-salry 0.29 0.51 0.67 0.00
## cits-salry 0.35 0.55 0.70 0.00
#bcdstats::adjust.corr(cohen1[,3:5])
```

3.3 Simple regressions of Salary on each of the two IVs

Next, use the 'lm' function from R's base installation to do linear modeling. Initially, lets do the two simple regressions of salary (the DV) on each of the IV's separately. We do this for comparability to how we approached the multiple regression problem before, but mostly to have the regression coefficients in hand in order to compare them to the regression coefficients for the IVs when included in the two-IV model.

First, we detach the cohen1 data frame so that we can use the "data" argument in the 1m function, a better practice.

```
detach(cohen1)
#produce the two single-IV models
fit1 <- lm(salary~pubs, data=cohen1)</pre>
fit2 <- lm(salary~cits, data=cohen1)</pre>
# obtain relevant inferences and CI's for fit1 with pubs as IV
summary(fit1)
##
## Call:
## lm(formula = salary ~ pubs, data = cohen1)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     ЗQ
                                             Max
                         333.7
##
  -20660.0 -7397.5
                                 5313.9
                                        19238.7
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 48439.09
                            1765.42 27.438 < 2e-16 ***
## pubs
                 350.80
                             77.17
                                      4.546 2.71e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 8440 on 60 degrees of freedom
## Multiple R-squared: 0.2562, Adjusted R-squared: 0.2438
## F-statistic: 20.67 on 1 and 60 DF, p-value: 2.706e-05
confint(fit1, level=0.95) # CIs for model parameters
##
                   2.5 %
                              97.5 %
## (Intercept) 44907.729 51970.4450
## pubs
                 196.441
                            505.1625
anova(fit1)
## Analysis of Variance Table
##
```

```
## Response: salary
##
                             Mean Sq F value
            Df
                   Sum Sq
                                                 Pr(>F)
## pubs
              1 1472195326 1472195326 20.665 2.706e-05 ***
## Residuals 60 4274424497
                            71240408
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Anova(fit1,type="III") # note this is Anova, not anova
## Anova Table (Type III tests)
##
## Response: salary
                   Sum Sq Df F value
                                        Pr(>F)
##
## (Intercept) 5.3632e+10 1 752.831 < 2.2e-16 ***
              1.4722e+09 1 20.665 2.706e-05 ***
## pubs
## Residuals 4.2744e+09 60
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
# obtain relevant inferences and CI's for fit2 with cits as IV
summary(fit2)
##
## Call:
## lm(formula = salary ~ cits, data = cohen1)
##
## Residuals:
##
       Min
                  1Q
                       Median
                                    3Q
                                            Max
                     -884.1 5461.7 24096.2
## -16462.0 -5822.4
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 42315.71
                          2662.69 15.892 < 2e-16 ***
## cits
                310.75
                             60.95
                                   5.098 3.69e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 8175 on 60 degrees of freedom
## Multiple R-squared: 0.3022, Adjusted R-squared: 0.2906
## F-statistic: 25.99 on 1 and 60 DF, p-value: 3.69e-06
confint(fit2, level=0.95) # CIs for model parameters
##
                    2.5 %
                              97.5 %
## (Intercept) 36989.5395 47641.8738
## cits
                 188.8201
                            432.6741
anova(fit2)
## Analysis of Variance Table
##
## Response: salary
##
             Df
                   Sum Sq
                             Mean Sq F value
                                                Pr(>F)
              1 1736876419 1736876419
## cits
                                        25.99 3.69e-06 ***
## Residuals 60 4009743405
                             66829057
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Anova(fit2,type="III") # note this is Anova, not anova
## Anova Table (Type III tests)
##
## Response: salary
##
                   Sum Sq Df F value
                                        Pr(>F)
## (Intercept) 1.6878e+10 1
                              252.56 < 2.2e-16 ***
               1.7369e+09 1
                               25.99 3.69e-06 ***
## cits
## Residuals
               4.0097e+09 60
##
  ___
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Summary of findings from bivariate analyses/inferences

- 1. Notice in the above analyses that each regression coefficient has a t value equivalent to what was found for the analogous bivariate correlation test. Not a coincidence - make sure you still recall why this is the case.
- 2. In addition, the R squareds reported by the 'summary' function are the square of the zero-order correlations and the F tests are the squares of the t values for testing the regression coefficients.
- 3. The ANOVA tables produced by anova and Anova do not give SS total. This is simply SS for Salary, the DV, and can be found other ways (e.g. var(salary)*(n-1)). This approach is used below in the multiple regression section.
- 4. The SS regression for each of the fits is labeled as a SS for that IV and not called SS regression as we have done previously. Make sure that understand this equivalence and the labeling choice made in R.
- 5. Compare the SS for each IV in the anova table with the comparable value in the Anova table. Ignoring the fact that 'Anova' uses exponential notation, these SS regression values are the same for the two different ANVOA functions. This is the expected outcome in simple regression. But they will not always be equivalent for a particular IV in multiple regression. We explore this below where the differences between anova and Anova become important in multiple regression.
- 6. The SS residual value for each IV is the same in the **anova** and **Anova** output for each of the two fits. Again, this is expected in simple regression because the SS regression and SS residual must sum to SS total. This point is emphasized here because it is one area where numerical and conceptual differences occur in multiple regression (see below), compared to these simple regressions.
- 7. Finally, notice that the F test values obtained by either the **anova** function or the **Anova** function (upper/lower case A) are identical. This is established now for comparison to what happens when we have two IVs below.

3.3.1 Diagnostic plots for simple regressions

Next, we want to obtain the base set of diagnostic plots that R makes available for 'lm' objects. This is the same set that we reviewed in the simple regression section of the course plus an additional normal probability plot of the residuals.

First for fit1:

layout(matrix(c(1,2,3,4),2,2)) # optional 4 graphs/page
plot(fit1)



I prefer to draw the qqplot with the 'qqPlot' function from the 'car' package since it provides confidence bounds. See Fox and Weinberg (2011).

qqPlot(fit1, distribution="norm", id=F)



norm Quantiles

Next, the same plots for fit2:

layout(matrix(c(1,2,3,4),2,2)) # optional 4 graphs/page
plot(fit2)



And the preferred qqplot for fit2: qqPlot(fit2, distribution="norm", id=FALSE)



Since the tests of these two simple regression fits would not typically be used, the diagnostic plots would also not typically be of interest. The analyses seen above are largely here to put in place approaches and numeric values to compare the more interesting multiple regression results found below.

4 Implementation of Standard Multiple Regression Analyses: Linear modeling with multiple IVs

When we covered implementation of simple regression procedures in R, we covered most of the R syntax required for multiple regression. Only a few additional coding strategies are necessary to employ models with more than one IV. We also reviewed those simple regression techniques in the previous section of this document. The illustrations outlined below emphasize a two-IV model for comparability to the example we worked out in SPSS. Some sections extend this capability to more than two IVs, but we will emphasize that to a greater degree at a later point in the course and in other documents.

The reader should also note that the following sections have two interrelated goals. One is simply the illustration of coding strategies in R for basic linear modeling. Some sections go beyond that narrow goal and continue to reinforce and expand the conceptual framework we have put in place for multiple regression. Some more advanced R strategies are also employed to obtain useful graphics and output that is formatted well for the markdown approach taken in this document.

The general sequence of the implementation is the following:

- a. perform the two-IV multiple regression
- b. obtain all the components and tests the we implemented in SPSS for the multiple regression
- c. obtain the basic evaluations of residual normality and homoscedasticity
- d. obtain additional information such as semi partial correlations, beta weights and CI's
- e. reinforce the concepts of unique and common proportions of variance, R squared, SS partitioning, etc
- f. perform some diagnostics (topics not yet covered in lecture)

4.1 The core linear modeling approach with R

Requesting a multiple regression using lm is an easy extension of the basic expressions used for simple regression. The "model" specification still uses the "tilda" symbol to indicate the DV to the left and the IV's to the right. All IV's are named, and the plus symbol is used to essentially list the full set of IV's. Here, I fit two models, each with the same two IVs, but I named them in opposite order. We will explore whether this makes any difference in outcome.

Typically we would NOT fit both models of the different IV orders. It is done here for instructional purposes that play out below and later in the semester.

First, establish the two alternative model fits:

```
fit3 <- lm(salary~pubs+cits, data=cohen1)
fit4 <- lm(salary~cits+pubs, data=cohen1)</pre>
```

4.1.1 Explore fit3 (salary~pubs+cits)

Obtain the same set of additional summaries/analyses/plots we did for simple regression. First for model fit3:

summary(fit3)

##

```
## Call:
## lm(formula = salary ~ pubs + cits, data = cohen1)
##
## Residuals:
##
       Min
                  10
                       Median
                                     ЗQ
                                             Max
##
  -17133.1 -5218.3
                       -341.3
                                5324.1
                                         17670.3
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 40492.97
                           2505.39
                                    16.162
                                            < 2e-16 ***
                 251.75
                                      3.452
                                            0.00103 **
## pubs
                             72.92
                                            0.00014 ***
## cits
                 242.30
                             59.47
                                      4.074
##
  ___
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 7519 on 59 degrees of freedom
## Multiple R-squared: 0.4195, Adjusted R-squared: 0.3998
## F-statistic: 21.32 on 2 and 59 DF, p-value: 1.076e-07
confint(fit3, level=0.95) # CIs for model parameters
```

##		2.5 %	97.5 %
##	(Intercept)	35479.6889	45506.2540
##	pubs	105.8395	397.6605
##	cits	123.3023	361.2931

At this point, let's recall the interpretation of the values for the regression coefficients. For comparison's sake, analyses in the previous section found that when, in simple regression, salary was regressed on pubs, the regression coefficient was \$350.80. It was interpreted to mean that each publication was related to an increase in salary of \$350.80. Now, in the two-IV regression, the regression coefficient is seen to be \$251.75. The change is recognized to have come about because the two IV's are correlated and the multiple regression coefficient is said to be a partial regression coefficient. That is, it is the change in the DV for a one unit change in the IV, controlling for the presence of the other IV. So, in fit3, each pub is uniquely associated with about a \$251.75 increase in salary. The interpretation of the regression coefficient for cits is similar.

The conclusion is that the impact of each IV in a regression model is only interpretable in the context of that model. The presence or absence of other IV's will influence the capability of any IV to predict the DV, when the IVs are correlate.

Next, we produce ANOVA summary tables two ways, using both the **anova** and **Anova** functions. Are they different? Note that I have formatted the ANOVA tables in a manner that might be unfamiliar, using the 'kable' and 'tidy' functions. This approach permits direct comparison of the output from the two functions since the default listing for the 'Anova' table uses scientific notation for SS values. See the section below on R Markdown formatting capabilities for more detail on this.

The first thing to notice about the ANOVA tables found below is that they do not simply present SS Regression, SS Residual and SS total as you might have expected from how we saw that SPSS produced ANOVA tables. SS total is not shown at all, and SS Regression appears to be broken apart and assigned to the two IVs. Exactly where these SS for pubs and cits come from is a longer discussion. We covered this point earlier in our multiple regression expositions. Reiteration of the importance of this question is begun just below here and in an later section of this document (where we conceptually connect these SS to squared semi-partial correlations).

Make sure you see that the "statistic" column in the kable/tidy version of the ANOVA tables is actually just the F value. Make sure you know how these F's were computed (what divided by what?)

kable(tidy(anova(fit3)))

term	df	sumsq	meansq	statistic	p.value
pubs	1	1472195326	1472195326	26.03841	0.0000037
cits	1	938602110	938602110	16.60086	0.0001396
Residuals	59	3335822387	56539362	NA	NA

"unique" ss for each IV produced by this Anova function. # Can also be found by multiplying # the squared semi-partial for each IV by SStotal kable(tidy(Anova(fit3,type="III")))# note this is Anova, not anova

term	sumsq	df	statistic	p.value
(Intercept)	14769235188	1	261.22041	0.0000000
pubs	673921018	1	11.91950	0.0010337
cits	938602110	1	16.60086	0.0001396
Residuals	3335822387	59	NA	NA

At this point, in comparing the results from the two anova functions, it is useful to recall that **anova** produces Type I SS and **Anova** was set up to request Type III SS. For the moment, lets just note that the SS and F value for cits is the same in both anova tables. Cits was entered second in the model specification for fit3. This point is expanded below.

Next, we obtain the standard set of diagnostic plots are obtained as they were for simple regression. Recall that I ask for these plots to be put on one figure by using the 'layout' function. At this point in the course, we have not actually covered two of the four plots, but two are our standard assessments of the normality and homoscedasticity assumptions. There do not appear to be strong violations of either assumption (seen in the two plots on the left)

```
#influence(fit3)
# regression diagnostics
layout(matrix(c(1,2,3,4),2,2)) # optional 4 graphs/page
plot(fit3)
```



And we can also obtain the preferred qqplot: qqPlot(fit3, distribution="norm", id=F)



A histogram of the residuals with a normal curve overlaid also provides a helpful visualization.



4.1.2 Explore fit4 (salary~cits+pubs)

Next, we examine model fit4 (the one with the opposite order of IVs listed in the model specification): summary(fit4)

```
##
## Call:
## lm(formula = salary ~ cits + pubs, data = cohen1)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     ЗQ
                                             Max
## -17133.1 -5218.3
                       -341.3
                                5324.1 17670.3
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 40492.97
                           2505.39
                                    16.162 < 2e-16 ***
                 242.30
                             59.47
                                      4.074 0.00014 ***
## cits
## pubs
                 251.75
                             72.92
                                      3.452 0.00103 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 7519 on 59 degrees of freedom
## Multiple R-squared: 0.4195, Adjusted R-squared: 0.3998
```

F-statistic: 21.32 on 2 and 59 DF, p-value: 1.076e-07 confint(fit4, level=0.95) # CIs for model parameters

##		2.5 %	97.5 %
##	(Intercept)	35479.6889	45506.2540
##	cits	123.3023	361.2931
##	pubs	105.8395	397.6605

The ANOVAS for fit4:

kable(tidy(anova(fit4)))

term	df	sumsq	meansq	statistic	p.value
cits	1	1736876419	1736876419	30.71977	0.0000007
pubs	1	673921018	673921018	11.91950	0.0010337
Residuals	59	3335822387	56539362	NA	NA

"unique" ss for each IV produced by this Anova function.

Can also be found by multiplying

the squared semi-partial for each IV by SStotal
kable(tidy(Anova(fit4,type="III")))# note this is Anova, not anova

term	sumsq	df	statistic	p.value
(Intercept)	14769235188	1	261.22041	0.0000000
cits	938602110	1	16.60086	0.0001396
pubs	673921018	1	11.91950	0.0010337
Residuals	3335822387	59	NA	NA

Once again, the SS and F's are identical for pubs, which is the second IV specified in the model specification - is is the "last entered". The SS and F's differ for the first entered variable, cits, as was also the case for fit3. These differences are explored below.

We can also examine diagnostic plots for fit4. They should look identical to those for fit3 since the residuals are the same for each of the two fits.

```
#influence(fit3)
# regression diagnostics
layout(matrix(c(1,2,3,4),2,2)) # optional 4 graphs/page
plot(fit4)
```



Here is the preferred qqplot for fit4: qqPlot(fit4, distribution="norm", id=F)


4.1.3 Comparing fit3 and fit4 results.

We have emphasized how the SS and F-tests for fits 3 and 4 differ, depending on whether we use **anova** or **Anova**. But it is helpful to ask which components are identical for the two Fits. These items/statistics are identical:

- 1. The Multiple R-squared
- 2. SS Regression (although not printed above). It is found by adding the SS for the two IVs from the anova-produced table.
- 3. SS Residual
- 4. df regression/residual
- 5. The regression coefficients, confidence Intervals, and t-tests of the coefficients are also identical for the two models
- 6. Below, we also see that beta weights, partial r's, semi-partial r's, tolerances and unique/common variance proportions are identical for the two models

The primary difference emerges when examining the SS partitioning. Lets do one more comparison here. These SS computations will be revisited in a later chapter. If we rerun the fit3 model here it will permit comparing the SS, F's and t-tests of the regression coefficients when using either anova or Anova. From this analysis, make note of the t-values. I saved them out of the model fit to work with in a section below.

```
fit3b <- lm(salary~pubs+cits,data=cohen1)
summary(fit3b)</pre>
```

```
##
```

```
## Call:
## lm(formula = salary ~ pubs + cits, data = cohen1)
##
## Residuals:
##
        Min
                   1Q
                        Median
                                      3Q
                                              Max
##
   -17133.1
             -5218.3
                        -341.3
                                 5324.1
                                          17670.3
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 40492.97
                            2505.39
                                     16.162
                                             < 2e-16 ***
## pubs
                 251.75
                              72.92
                                      3.452
                                             0.00103 **
## cits
                 242.30
                              59.47
                                      4.074
                                             0.00014 ***
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 7519 on 59 degrees of freedom
## Multiple R-squared: 0.4195, Adjusted R-squared: 0.3998
## F-statistic: 21.32 on 2 and 59 DF, p-value: 1.076e-07
tvals <- summary(fit3b)$coefficients[2:3,3] # subsetting to extract the t's for the two IVs
tvals
##
       pubs
                cits
## 3.452463 4.074415
Next, re-obtain the SS and F tests from the Anova function, and save the F values out to a vector.
```

Anova(fit3b, type=3)

```
## Anova Table (Type III tests)
##
## Response: salary
## Sum Sq Df F value Pr(>F)
```

```
## (Intercept) 1.4769e+10 1 261.220 < 2.2e-16 ***
## pubs     6.7392e+08 1 11.919 0.0010337 **
## cits     9.3860e+08 1 16.601 0.0001396 ***
## Residuals     3.3358e+09 59
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
fvals <- Anova(fit3b, type=3)[2:3,3] # subsetting to extract F's for the two IVs
fvals</pre>
```

[1] 11.91950 16.60086

Now we can take the square roots of the F values and compare those to the t values. Since each F is based on 1 and n-k-1 df, the square roots will produce a t value with those n-k-1 df.

fvals[^].5

[1] 3.452463 4.074415
tvals
pubs cits

3.452463 4.074415

Since the quantities are identical, we can conclude that the t-tests of the regression coefficients are equivalent to tests of Type III SS for each IV. Recalling comparisons of fit3 and fit4 above (the two opposite IV entry orders), we also recall that the coefficients and their t-tests are the same. The conclusion is that tests of the regression coefficients for any model are actually tests of type III SS for those IVs.

The type I SS seen from the **anova** analyses lead us to conclude that those SS are found by computing the unique SS for each variable at the point it enters the equation. That is why the SS for each variable differs, depending on whether it is entered first or second. The type III SS are found be treating each variable as if it were the second (last) to enter. Further implications of this are found in a later chapter.

4.1.4 Efficiently obtaining important descriptive components of a multiple-IV linear model

It has been argued that several core components of multiple regression should be produced whenever any linear model is fit. Some of those components have not yet been requested/produced with the methods outlined above (e.g. beta weights). I have written my own R function that obtains many of these additional items that we have previously discussed. The function is called mrinfo (multiple regression information) and I just call it Mr. Info. It is available in the bcdstats package.

The syntax is to just pass a lm model fit object to the mrinfo function. We do so here with our fit3 model and we will examine the output. The unique proportion of variance accounted for by each IV are equivalent to the square of the semi-partial correlation of that IV with the DV, after residualizing that IV on all other IVs. The argument minimal permits request of only a set of supplemental information or a longer list of regression information that has been covered above in piecemeal fashion.

mrinfo(fit3, minimal=TRUE)

```
## [[1]]
## NULL
##
## $`supplemental information`
##
        beta wt structure r partial r semipartial r tolerances unique common
## pubs 0.36323
                    0.78145
                               0.40996
                                             0.34245
                                                         0.88885 0.11727 0.13891
##
  cits 0.42867
                    0.84880
                               0.46860
                                             0.40414
                                                         0.88885 0.16333 0.13891
##
          total
## pubs 0.25618
## cits 0.30224
```

```
##
## $`var infl factors from HH:vif`
## pubs cits
## 1.12505 1.12505
##
## [[4]]
## NULL
```

We can obtain these statistics for fit4 as well, but all are identical to those from fit3.

```
mrinfo(fit4, minimal=TRUE)
## [[1]]
## NULL
##
## $`supplemental information`
##
        beta wt structure r partial r semipartial r tolerances
                                                                 unique common
                                              0.40414
                                                         0.88885 0.16333 0.13891
## cits 0.42867
                    0.84880
                               0.46860
## pubs 0.36323
                    0.78145
                               0.40996
                                              0.34245
                                                         0.88885 0.11727 0.13891
##
          total
## cits 0.30224
## pubs 0.25618
##
## $`var infl factors from HH:vif`
##
      cits
              pubs
## 1.12505 1.12505
##
## [[4]]
## NULL
```

The beta weights, partial correlations, semi-partial correlations, and tolerances should be values that the reader understands and expects, given our prior conceptual and SPSS work. But what are the "unique", "common", and "total" values. The reader should examine those, keeping in mind core ways of thinking about the Multiple R squared (which you know from above and from SPSS work). Recall that the multiple R-squared from both fits 3 and 4 was about .42. How might we describe the "unique" quantities, and how do they relate to the semi-partials? (Hint: sum the two unique proportions and add the common proportion)

4.1.4.1 A caveat on using 'mrinfo' The original data file from the cohen text called the "years since degree" variable "time". We will examine a 3 IV model using that variable later in this document. At present, the word time can't be used for a variable in a lm model that is submitted to the 'mrinfo' function. It thinks it is an R function called 'time'. So I changed it to degree_yrs.

4.2 Diagnostic and Added-variable plots for the full two-IV model

Many graphical techniques can be explored for evaluation of assumptions, influence, and partial relationships in multivariable systems. A few are explored here, and others are presented in later chapters (e.g., see the chapter on the **olsrr** package).

4.2.1 Working with the residuals and predicted scores (yhats)

Residuals and yhats from any 1m model can be quickly obtained.

```
fit3.resid <- resid(fit3)
fit3.pred <- predict(fit3)</pre>
```

These vectors can also be standardized.

```
fit3.zresid <- scale(fit3.resid)
fit3.zpred <- scale(fit3.pred)</pre>
```

We might also place these new vectors into a data frame along with the original variables, in order to save them for other future work.

```
# cohen1 was already detached
#detach(cohen1)
cohen1new <- cbind(cohen1, fit3.resid, fit3.pred, fit3.zresid, fit3.zpred)
# and then reattach cohen1 for possible use later
#attach(cohen1new)</pre>
```

Now we can work directly with these new residual and yhat vectors. Some of these plots duplicate what was shown above, but it is useful to have the standardized residuals and yhats available.

```
plotNormalHistogram(fit3.zresid)
```



qqPlot(fit3.zresid)



norm quantiles

[1] 32 28
plot(fit3.zpred,fit3.zresid)



Now detach the cohen1new data frame and reattach the original to be used later.

#detach(cohen1new)
#attach(cohen1)

4.2.2 Added Variable Plots

Added variable plots permit evaluation of the so-called partial regressions. These are actually just plots of the partial correlations of each IV (residualized on the other IVs) with Y (also residualized on the other IV). Note that the "test" of these partial correlations is equivalent to the test of the semi-partial correlations and also equivalent to the tests of the individual regression coefficients found in the summary output.

See the later chapters on Extensions and use of the **olsrr** package for further treatment of this.

#require(car)
avPlots(fit3)



5 Review Unique and common proportions of variance and Type I vs III Sums of Squares

We can think about an ability to partition the SS of the DV, not only into Regression and Residual components, but to further break down the Regression SS into unique and common parts. This builds on the concepts just reviewed above and connects to earlier work.

THIS SECTION AND THE NEXT ARE THE CRITICAL CONCEPTUAL FRAMEWORK THAT HAS BEEN DEVELOPED FOR MULTIPLE REGRESSION IN THE 510/511 COURSE. WE COVERD THIS WHEN WE WORKED THROUGH THE SPSS IMPLEMENTATION. IF YOU DID NOT CONSOLIDATE IT THEN, OR ON ASSIGNMENTS/EXAM PREP, THEN THIS IS THE TIME/PLACE TO BE CERTAIN ABOUT IT.

In this section we will: 1. find unique SS for each IV, and the common SS 2. Utilize semi-partial correlations and the derived unique/common proportions from 'mrinfo' 3. use these quantities to reinforce concepts that we have repeatedly covered in slightly different ways 4. use R as a calculator to accomplish some of these things 5. Understand that since we are going to use some rounded quantities from 'mrinfo' some error is introduced into our computations. So, some things may not match exactly as the should if the concepts are correct.

5.1 SS components, semi-partial correlations, unique and common proportions/SS $\,$

A few components are computed "manually" to establish some basic quantities. SS_{total} is found two different ways (the quantities are returned after this code chunk), and they match. Other quantities are calculated, but not printed here - they are used below.

```
# compute a few SS manually.....
# first obtain a few values to use, SS Total, MSresid, and R squared
SStotal <- var(cohen1$salary)*61 # variance times n-1 gives SS for that variable
# name an object that is the anova summary table for the fit3 model
a <- anova(fit3)
#extract the MS Residual and multiple R squared from that table
MSresid <- a$`Mean Sq`[3]
rsquared <- summary(fit3)$r.squared
# use the above quantities to compute SS Regression and SS Residual
SSregression <- rsquared*SStotal
SSresidual <- MSresid*59 # MS resid times its df
# sum them to doublecheck
SSregression + SSresidual # should equal SStotal
```

[1] 5746619823
SStotal # show SStotal to doublecheck

[1] 5746619823

Focus on the information value of the semi-partial correlations (we found them with mrinfo above). By squaring them, we have the unique proportions of variance accounted for by each IV - and the values match what was reported by mrinfo above.

```
semipartialpub <- .3424509
semipartialcit <- .4041425
# square them to produce "unique" proportions of explained variance
# see that it matches the table from mrinfo
part1sqrd <- semipartialpub<sup>2</sup>
part2sqrd <- semipartialcit<sup>2</sup>
part1sqrd
```

[1] 0.1172726

part2sqrd

[1] 0.1633312

Now, we can use the squared semi-partials to find the common proportion of variance shared between the two IVs. Does it match what was returned by mrinfo?

```
# now that we know the unique fractions we can find the common proportion
# and it should match the mrinfo value
# recall that rsquared was defined at the beginning of this section
commonprop <- rsquared -part1sqrd - part2sqrd
commonprop</pre>
```

[1] 0.138912

Now compute the unique and common SS. It would be useful to compare these SS to the table found above in the basic model section for fit3 and fit4 and the section that compared the ANOVA summary tables from anova and Anova.

```
# now compute the unique and common SS
uniqueSSpub <- part1sqrd*SStotal
uniqueSScit <- part2sqrd*SStotal
uniqueSSpub</pre>
```

[1] 673921157

uniqueSScit

[1] 938602084

These do match the SS for the two IVs found by using the Anova function (with type III SS) from either fit3 or fit4. What is their sum? It should be less than SS regression since each is the unique component and the full $SS_{regression}$ includes the common SS where the two IVs overlap in the salary space.

```
# should these two values add up to SSregression?
# No. should be something less
uniqueSSpub + uniqueSScit
```

[1] 1612523240

SSregression

[1] 2410797436

Lets find a common SS by using the common proportion seen above.

```
# how much less?????? the amount due to the shared SS
# its proportion is the common proportion
commonprop
```

[1] 0.138912

SScommon <- commonprop*SStotal SScommon

[1] 798274196

Now verify that all three components sum to $\mathrm{SS}_{\mathrm{regression}},$ as they should:

```
# do all three sum to SSregression now? compare:
SScommon + uniqueSSpub + uniqueSScit
```

[1] 2410797436 SSregression

[1] 2410797436

5.2 F tests on the unique components?

Now that we have derived unique SS for each IV, can we do an F test? That is, can we test a null hypothesis that an individual IV does not uniquely contribute to the R squared (also phrased as the model fit)? Yes. Just like any F test:

First, find the MS for each IV, based on the unique SS.

```
# can we do tests of these unique SS? Sure, F tests with MSresid as "error" term
# We need to convert the unique SS to MS, so we need df
# each has 1 df, since each represents one variable
# the MS are the SS divided by 1
MSuniquepub <- uniqueSSpub/1
MSuniquepub
## [1] 673921157
MSuniquecit</pre>
```

[1] 938602084

Now find the F values by using MSresid as established above.

```
# and now the two F's
Fpubunique <- MSuniquepub/MSresid #(df are 1,59)
Fcitunique <- MSuniquecit/MSresid #(df are 1,59)
Fpubunique</pre>
```

[1] 11.9195 Fcitunique

[1] 16.60086

```
# can you find these F's anywhere above?
# hint: look in the table of the Anova function (not anova)
```

Yes, they match the F's from the Anova tables above, using Type III SS. And, we can take their square roots to compare to the t values for each regression coefficient. This duplicates the illustration seen in a previous chapter.

```
# Now take the square roots of these F's
Fpubunique<sup>.5</sup>
```

[1] 3.452463
Fcitunique^{.5}

[1] 4.074415

```
# look familiar???
# yes, they are the t's that provided the test stat for the regression coefficients
```

Conclusions?

- 1. tests of semipartial correlations can be done as F tests by converting to SS and MS
- 2. these tests are equivalent to the F tests done on TYPE III SS in the Anova function
- 3. since the F's are the squares of the t's for the two regression coefficients, we conclude that tests of the regression coefficients are tantamount to tests of the semi-partial correlations. This is why we can call them partial (actually semi-partial) regression coefficients.
- 4. The unique SS are seen to be equivalent to the SS computed above by the 'Anova' function (rather than 'anova'), and are thus equivalent to what are called Type III SS. To be contained.....

5.3 Compare and Contrast information from fit3 and fit4

Take some time again to ponder the output from above, comparing comparable values from fit3 and fit4 as we did in the previous chapter. Recall that fit3 and fit4 differed only in the order of entry of the two IVs, with pubs first in fit3. Many/most of the components are the same. The primary differences seem to emerge in the 'anova' vs 'Anova' output. Take the square root of each of the F tests from 'anova' and 'Anova'. Which ones match the t's for tests of the comparable regression coefficients and which differ? This was explored briefly earlier in this document. This section reviews that information again in a more succinct tablular form. I have created a table of SS produced by both 'anova' and 'Anova' for each of fits 3 and 4 to facilitate this comparison. Make certain you can see, in the above output, where these values came from.

anova produced Type I SS, and Anova produced Type III SS				
	Fit3 (order	: pubs, cits)	Fit4 (order: cits, pubs)	
Term	anova SS	Anova SS	anova SS	Anova SS
Pubs	1472195326	673921018	673921018	673921018
Cits	938602110	938602110	1736876419	938602110
Residual	3335822387	3335822387	3335822387	3335822387
Sum of Pubs and Cits	2410797436	1612523128	2410797437	1612523128
SS Regression (Rsqrd*SStot)	2410797436	2410797436	2410797436	2410797436
Sum of Pubs, Cits, Resid	5746619823	4948345515	5746619824	4948345515
SS Total (SS of salary)*	5746619823	5746619823	5746619823	5746619823

Compare Type I and Type III SS for the two Im fits

* Note that SS total was independently calculated directly from the DV values

Also note that when SS values don't match in the last decimal place it is due to rounding error

For the moment, notice a few things from this table, recalling that both fit3 and fit4 have the same intercept, regression coefficients and their t's, multiple R squared, SS total, SS residual, SS regression, and F-tests of the whole equation with 2 and 59 df. The SS for pubs and cits should be a partitioning of SS regression.

- 1. In 'anova' output, the sum of the SS for pubs and cits matches SS regression both for fit3, and fit4 which also match each other.
- 2. In 'Anova' output the sum of the SS for pubs and cits is always less than SS Regression, and adding them to SS residual yields a quantity less that SS total.
- 3. For 'anova' output, summing SS for pubs, cits, and residual DOES yield the SS total, as expected.
- 4. Sometimes SS for pubs (or for cits) from 'anova' and 'Anova' match and sometimes they don't. It depends on which fit one examines. So, we must conclude that the 'order' of model specification has something to do with the difference in Type I and Type III SS. Type III SS are found by treating each IV as if it were the last to enter the model, thus controlling for all other IVs.
- 5. The reader might also compare the SS for each IV under each model to the SS Regression from when that IV was used in simple regression in this document (in the bivariate computations section)

A similar table compares F test values from 'anova' and 'Anova' for the two fits. The pattern here follows what was seen for SS above since the F's are derived from those SS. The more interesting comparison is the comparison of those F's to the square of the t's found from testing the two regression coefficients in the two models. This comparison lets us get to the heart of what is being tested with the F tests vis a vis the way we discussed the null hypotheses for the t tests of the regression coefficients.

The t's test a null hypothesis that the "unique" contribution of that predictor is zero, when controlling for the other predictor(s). And the squares of these t's don't always match the F's from the 'anova' function. They do match for the 'Anova' function. Further discussion of this took place above, connecting the findings to concepts deriving from our understanding of semi-partial correlations.

Compare Type I and Type III F tests produced by				
the two ANOVA functions and the t values for the regression coefficients				
Fit3 (order: pubs, cits)				
anova F	Anova F	t*	t squared	
26.03841	11.91950	3.452	11.916	
16.60086	16.60086	4.074	16.597	
Fit4 (order: cits, pubs)				
anova F	Anova F	t*	t squared	
11.91950	11.91950	3.452	11.916	
30.71977	16.60086	4.074	16.597	
30.71977	16.60086	4.074	16.597	
30.71977 ed to 3 decimals	16.60086 and this introdu	4.074 ces some error v	16.597 vhen	
	anova F 26.03841 16.60086 anova F 11.91950	Type I and Type III F tests produ tions and the t values for the reg Fit3 (order: anova F Anova F 26.03841 11.91950 16.60086 16.60086 Fit4 (order: anova F Anova F 11.91950 11.91950	Type I and Type III F tests produced by tions and the t values for the regression coefficient Fit3 (order: pubs, cits) anova F Anova F t* 26.03841 11.91950 3.452 16.60086 16.60086 4.074 Fit4 (order: cits, pubs) anova F Anova F t* 11.91950 11.91950 3.452	

This general topic is further addressed below entitled "Work with Sums of Squares a bit more, along with unique and common proportions of variance."

Typically we would not evaluate both models fit3 and fit4 which entered the IVs in different orders since to final model is largely the same (with the caveat about unique vs sequential or Type I vs Type III SS). If you examined the coefficients tables for fit3 and fit4, you found the regression coefficients to be the same (and the t-tests of them as well) So much of the above work is duplicative. Perhaps we would only run fit3. In that case, the kind of comparison in the next section on formally comparing models might still be of interest.

The issue with types of SS computation is revealing in this comparison of fit3 and fit 4 which have only two IVs. The SS computation issue will be addressed several times later in the semester and becomes more involved when the number of IVs exceed two. For now, focus on the comparability of the fit3 and fit4 models with regard to overall fit, and with regard to coefficients. Additional illumination on fit3 and fit4 differences in SS will be connected to semi-partial correlations and unique vs common SS issues addressed below and that is the part of this topic that is the important conceptual framework. The primary conclusion from the SS comparison work here, is that order of entry of the IVs into a model can affect the SS computation, depending on whether the analyst chooses Type I (sequential) or Type III (unique) SS.

6 Formally Comparing Models

In one more attempt to explore how to think about tests of each IV, vis a vis the kinds of things implied just above, lets use the anova function in a different way.

We can pass two linear models to 'anova' and ask it to compare them. The second model in each specification has to contain a superset of the IV's in the first (i.e., the IV used in the first plus at least one more). It is somewhat like the "stepping" idea we introduced in SPSS and 'anova' essentially tests the improvement in fit of the second model over the first. Carefully look at the F tests for these model differences in Fit, and compare them to what you just examined above for the 'anova' vs 'Anova' approaches in the two different orders of the two-IV fit models. The F's that compare the two models are the same as the test of the R squared "increment" that we covered in SPSS work earlier.

We consider beginning wth fit1 (only pubs was an IV in that simple regression), and compare fit 3 (which also included cits as an IV) to test the increment in the R-squared produced by the inclusion of cits. The F value matches the Type III SS F test for cits and is the square of the t value that tested the regression coefficient of cits.

```
# compare model 3 to model 1 - stepping approach, evaluating a new variable (cits)
anova(fit1,fit3)# note this is anova, not Anova
```

Analysis of Variance Table

The next illustration compares the full model to the simple regression that only contained cits (fit2). The test is therefore the test of an hypothesis that increment in SS accounted for by pubs is zero. And this F also matches the F test of the Type III SS seen above.

```
# compare model 3 to model 2 - stepping approach, evaluating a new variable (pubs)
anova(fit2,fit3)# note this is anova, not Anova
```

```
## Analysis of Variance Table
##
## Model 1: salary ~ cits
## Model 2: salary ~ pubs + cits
##
     Res.Df
                   RSS Df Sum of Sq
                                          F
                                              Pr(>F)
## 1
         60 4009743405
## 2
         59 3335822387
                        1 673921018 11.919 0.001034 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

7 Further work with residuals and yhats

This section is structured with three goals:

- 1. Learning how to extract the residuals and the yhats from a 'lm' fit object in order to do additional work with them. You may recall that we learned to "save" residuals and yhats from an SPSS regression. That is also done here.
- 2. The diagnostic plots from above did not give us a frequency histogram of the residuals or even a simple boxplot. Once extracted, we can easily produce those types of graphs.
- 3. We can revisit some primary concepts in linear modeling by further examining the meaning of the yhats.

First, extract the residuals and yhats from the model fit (fit3 in this instance). We could add them to the data frame containing the original variables for later potential use (although we can work with the newly created vectors directly here.

```
fit3.resid <- residuals(fit3)
fit3.pred <- predict(fit3)
#create a new data frame with the original variables plus the exracted residuals and yhats
cohen2 <- cbind(cohen1,fit3.resid,fit3.pred) #note that fit3.pred is the yhats</pre>
```

Examine the frequency histogram of the residuals. It looks only slightly positively skewed even though the original DV was somewhat skewed. The normality assumption may not be seriously violated.

```
# simple frequency histogram of the residuals from fit3
hist(fit3.resid)
```

Histogram of fit3.resid



Boxplots of the residuals and yhats are produced next, revealing one potential outlier in the yhats.

```
# boxplots
layout(matrix(c(1,2),1,2)) #optional 2graphs/page
boxplot(fit3.resid,xlab="Residuals",data=cohen2,col="lightgrey")
boxplot(fit3.pred, xlab="Yhat",data=cohen2,col="lightgrey")
```



Residuals

Yhat

Numerical summaries of the residuals and yhats are provided by the describe function.

psych::describe(fit3.resid)

vars n mean sd median trimmed range madmin max ## X1 0 7394.97 -341.27 -160.91 7519.46 -17133.14 17670.34 34803.48 1 62 ## skew kurtosis se -0.4 939.16 ## X1 0.22 psych::describe(fit3.pred)

vars n mean sd median trimmed mad min max range
X1 1 62 54815.76 6286.59 53630.77 54275.57 4982.75 42497.52 79670.52 37173
skew kurtosis se
X1 1.13 2.45 798.4

Now that we have the yhat values available, we can do one more analysis that will be familiar. If we correlate the yhats with salary (the DV), we find an interesting quantity, when we square it.

correlate the original DV (salary) with the yhats from the two-IV linear model
r_y_yhat <- with(cohen2, cor(salary, fit3.pred))
r_y_yhat</pre>

[1] 0.6477003
square that value
r_y_yhat² #does the value look like something you recognize?

[1] 0.4195157

When covering this characteristic at earlier points, it was emphasized that the yhats are the best linear combination of the IVs, and carry all the information about the predictability of the DV from those IVs. It is a core feature of regression that the relationship between the DV and the yhats reveals the strength of the prediction, the multiple R (or R-squared).

8 Examine the fit of the plane in a 3D wireframe plot:

R has useful facilities for creating a three dimensional scatter plot. It is also possible to plot the model fit onto that 3D scatter plot. The two-IV model fits a plane. This section creates the model, generates the scatter plot, adds the plane, and permits rotation of the plot if the code is executed in R or RStudio.

Initially we have to relabel the variables in order to use the 3d scatter plot function below.

```
# to simplify the use of the scatter3D function, we re-label the variables.
# scatter3D expects the "Y" variable from a linear model to be in the "z" dimension of the 3D plot
z <- cohen1$salary
x <- cohen1$pubs
y <- cohen1$cits
# Again fit the two-IV model, but using the x,y,z variables
# x,y, and z are used for purposes related to the 3D surfce plotting below.
# This model is the same as fit3 examined above
fit3d1 <- lm(z-x+y)
#summary(fit.dep1b)
tidyfit.3d1 <- tidy(fit3d1) # nicer table than with summary
kable(tidyfit.3d1, format="markdown")
```

term	estimate	std.error	statistic	p.value
(Intercept)	40492.9715	2505.39437	16.162314	0.0000000
х	251.7500	72.91896	3.452463	0.0010337
у	242.2977	59.46809	4.074415	0.0001396

Drawing the 3d scatter plot is somewhat complicated and the student in 510/511 is not expected to master the code at this point. However, one should be able to take this code and modify it to be used in and two-IV model fit with other data sets.

In order to draw the 3D scatter plot we need to do the following:

- 1. Set up a grid of hypothetical points along a plane (I chose 21 lines for visual style reasons). The grid is established with minimal and maximal values in the x and y dimensions by using the minimal and maximal values for the x and y variables (pubs and cits in our case)
- 2. Create a matrix of predicted scores (in the y scale), using the prediction equation that we called fit3d1 above.
- 3. use the 'scatter3D' function from the 'plot3D' package. This draws the basic scatter plot, permits some color control of points, allows text and label editing, and permits addition of a surface.

```
# fitted points for droplines to surface if we want to draw them
fitpoints <- predict(fit3d1)</pre>
```

This 3D scatter plot capability is useful in one important manner not demonstrated in the html or pdf versions of this document. If the code is run in R (preferable RStudio), the 'scatter3D' function can be followed by a 'plotrgl' function (commented out here). This will pop up a graphics window and the user can use the mouse to manually rotate the 3D image. This provides superb visualization capability. In the scatter3D implementation here the theta and phi values are chosen to produce what I thought was the best rendition of the 3D image for the 2 dimensional rendering required for print documents.

```
# scatter plot with regression plane
scatter3D(x, y, z, pch = 20, cex = 1.5, cex.lab=.8,
          theta = 50, phi = 20, ticktype = "detailed",
           theta = 50, phi = 18, ticktype = "detailed",
#
          zlim=c(35000, 82000),
          #cex.axis=.9,
          xlab = "{Publications}", ylab = "Citations",
          zlab = "Salary",
          surf = list(x = x.pred, y = y.pred, z = z.pred,
                      facets = NA,
                      col="grey75",
                      fit = fitpoints),
          colkey=F, col=terrain.colors(length(z)),
          main = "",
          plot=T)
```



execute the plotrgl function outside of the pdf or html doc to interactively work with the plot
plotrgl()

9 Inferential tests for assumptions

There are three primary assumptions that linear modeling with the standard NHST t and F tests rely on:

- 1. The relationships among variables are best described by linear functions.
- 2. The residuals from a model are normally distributed.
- 3. Homoscedasticity.

Graphical evaluation of these assumptions has been the priority in previous considerations, as well in this document. Now we can turn to inferential tests regarding these assumptions.

9.1 Evaluation of the Residual Normality Assumption

Prior work with SPSS and R for linear models has only employed a graphical assessment to the normality assumption for residuals (and skewness/kurtosis computation). Frequency histograms and normal probability plots are useful, but at times one may wish to do an inferential test of a null hypothesis that residuals are normally distributed. Quite a few such inferential tests have been developed, and many are available in R.

While we can easily accomplish these tests, the analyst should be concerned about a binary outcome decision process in evaluation of the assumption. The real question should be something like "with the degree of non-normality present, how much of an impact on the tests of regression coefficients and the test of the whole model is there?" Strong guidance on this is lacking. However, some discussion can be found in standard regression textbooks and the reader is advised to consult Fox (2016), or any number of other sources (e.g., Cohen et al. (2003), Cook and Weisberg (1999), Darlington (1990), Howell (2013), Weisberg (2014), or Wright and London (2009)).

The options available in R will be presented here without regard to that more important question. The Anderson-Darling test may be the test that is most recommended. Historically, the Shapiro test is probably the most commonly used one, based on availability in other software. But with the 'nortest' package in R, many others are also available.

First, we can implement the historically common Shapiro-Wilk test:

```
shapiro.test(residuals(fit3))
```

```
##
## Shapiro-Wilk normality test
##
## data: residuals(fit3)
## W = 0.98639, p-value = 0.7239
```

This next code chunk shows code executing five different tests from the **nortest** package. But only one (Anderson-Darling) is executed here to keep this document small. The reader may recall from graphical approaches accomplished above that there is a slight degree of non-normality in the residuals from the basic two-IV model (fit3 or fit4). Is it a significant departure from normality, based on the sample size employed in this study?

```
# tests for normality of the residuals
#library(nortest)
ad.test(residuals(fit3)) # Anderson-Darling test (from 'nortest' package )
##
## Anderson-Darling normality test
##
## data: residuals(fit3)
## A = 0.34507, p-value = 0.4742
#cvm.test(residuals(fit3)) #Cramer-von Mises test ((from 'nortest' package )
#lillie.test(residuals(fit3)) #Lilliefors (Kolmogorov-Smirnov) test (from 'nortest' package )
#pearson.test(residuals(fit3)) #get Pearson chi-square test (from 'nortest' package )
#sf.test(residuals(fit3)) #get Shapiro-Francia test (from 'nortest' package )
```

Neither of these tests reach significance thresholds at $\alpha = .05$ and this is not suprising given the small amount of skewness visualized in the graphical assessments of the residuals seen in above sections of this document.

9.1.1 Evaluation of the normality assumption by testing skewness and kurtosis

Another approach to inference regarding the normality assumption is to test whether skewness and/or kurtosis of the residuals deviates from a null hypothesis that they are zero (as would be the case for a normal distribution). Once again, this document is more concerned with showing how such tests can be accomplished in R. Discussions about their interpretation and whether they should be used are left to other parts of the course and to the recommendations of the relevant textbooks.

R has two packages (tseries and moments) that contain functions to test skewness and/or kurtosis:

```
# also..... another way to evaluate normality is to test for skewness and kurtosis
#library(tseries)
jarque.bera.test(residuals(fit3)) # Jarque Bera test for normality of a variable
```

```
Jarque Bera Test
##
##
## data: residuals(fit3)
## X-squared = 0.77247, df = 2, p-value = 0.6796
#library(moments)
agostino.test(residuals(fit3))
##
##
   D'Agostino skewness test
##
## data: residuals(fit3)
\#\# skew = 0.22341, z = 0.78060, p-value = 0.435
## alternative hypothesis: data have a skewness
anscombe.test(residuals(fit3)) # Anscombe-Glynn test of kurtosis
##
##
   Anscombe-Glynn kurtosis test
##
## data: residuals(fit3)
\#\# kurt = 2.68478, z = -0.25858, p-value = 0.796
## alternative hypothesis: kurtosis is not equal to 3
```

Again, none of these tests would allow us to reject a null hypothesis of residual normality.

We might also simply perform a Z test by taking the ratio of skewness and kurtosis coefficients to their std errors (called a large-sample approximation in using the std normal Z here). We can obtain the coefficients and std errors in a couple ways:

We can write our own function to test skewness and kurtosis as the Z approximation called the large sample approximation suggested above. First, create the function and test skewness:

```
# we can build our own tests, since we know the
# std errors of skewness and kurtosis (fall semester work)
# these functions test the G1 and G2 statistics with their
# large sample std errors against nulls that the parameters are zero (as in normal distribs)
skewness.test <- function (lmfit) {</pre>
  require(psych)
  G1 <- psych::describe(residuals(lmfit),type=2)$skew # pull G1 from psych:::describe
  skewness <- G1
  N <- length(residuals(lmfit))</pre>
  stderrskew <- sqrt(6/N)</pre>
  teststat <- skewness/stderrskew</pre>
  pvalue <- 2*(pnorm(abs(teststat), lower.tail=F))</pre>
  outskew <- data.frame(cbind(N,G1,stderrskew,teststat,pvalue))</pre>
  colnames(outskew) <- c("N","G1","Std Error", "Z Test Stat","2-tailed p")</pre>
  rownames(outskew) <- c("")</pre>
  return(outskew)
}
# use this approach (z test) with large N. It returns a two-tailed p value
skewness.test(fit3)
##
               G1 Std Error Z Test Stat 2-tailed p
     Ν
##
    62 0.2289903 0.3110855 0.7361008 0.4616693
```

And now test kurtosis:

```
# now kurtosis
kurtosis.test <- function(lmfit) {</pre>
  require(psych)
  G2 <- psych::describe(residuals(lmfit),type=2)$kurtosis # pull G2 from psych:::describe
  kurtosis <- G2
  N <- length(residuals(lmfit))</pre>
  stderrkurt <- sqrt(24/N)</pre>
  teststat <- kurtosis/stderrkurt</pre>
  pvalue <- 2*(pnorm(abs(teststat), lower.tail=F))</pre>
  outk <- data.frame(cbind(N,G2,stderrkurt,teststat,pvalue))</pre>
  colnames(outk) <- c("N","G2","Std Error", "Z Test Stat","2-tailed p")</pre>
  rownames(outk) <- c("")</pre>
  return(outk)
}
kurtosis.test(fit3) # use this approach (z test) with large N. It returns a two-tailed p value
##
     Ν
                G2 Std Error Z Test Stat 2-tailed p
##
    62 -0.2388151 0.622171 -0.3838415 0.7010959
```

9.2 Inferential test of Homoscedasticity

Evaluation of homoscedasticity shares the same graphical-approach history in our prior work as the normality assumption discussed above. There is also the question of relative impact of varying decrees of violation of the assumption that needs to be considered (we have discussed this previously in the context of the 2-sample location test assumption of homogeneity of variance which is essentially the same assumption). But the narrow goal here is the illustration of approaches in R. First, we redo a plot of residuals against yhats using the **spreadLevelPlot** function from the **car** package. The plot gives a hint of larger residual spread for mid range values of yhats, but the pattern is carried by only a few data points.

```
# also a plot slightly different from the base system plot on the fit
# from the car package
spreadLevelPlot(fit3)
```



##

Suggested power transformation: 0.8964362

The ncvTest function (ncv standing for non-constant variance), also from the **car** package provides chisquare based test statistic that evaluates a null hypothesis of homoscedasticity. The test is more typically called the "Score" test.

```
# tests of homoscedasticity
# the ncvTest function is a test of non-constant error variance, called the Score test from car
ncvTest(fit3)
```

```
## Non-constant Variance Score Test
## Variance formula: ~ fitted.values
## Chisquare = 0.2914146, Df = 1, p = 0.58932
```

Another test of homoscedasticity is one that is frequently used. The Breusch-Pagan test is implemented in the **bptest** function found in the **lmtest** package. Neither it, nor the Score test yielded a result that would challenge the null hypothesis of homoscedasticity.

```
#library(lmtest)
bptest(fit3)
##
## studentized Breusch-Pagan test
##
## data: fit3
## BP = 2.665, df = 2, p-value = 0.2638
```

9.3 Global Evaluation of Linear Model Assumptiions using the 'gvlma' package.

A recent paper in JASA (Pena and Slate, 2006) outlined an approach to model assumption evaluation that is novel (and mathematically challenging). An R implementation is now available in the **gvlma** package. Initially, the approach evaluates a global null hypothesis that all assumptions of the linear model hypothesis tests are satisfied:

- 1. Linearity
- 2. Normality evaluation of skewness)
- 3. Normality (evaluation of kurtosis)
- 4. Link Function (linearity and normality)
- 5. Homoscedasticity

If this global null is rejected, then evaluation of each of four component assumptions is evaluated individually.

Fortunately, the 'gvlma' function is simple to use and the output is largely the standard NHST p value for each test.

A caveat here is that this type of approach is even more likely to be the target of the kinds of discussion we have had before on whether binary decision inferential tests are the way to act on knowledge of whether model assumptions are satisfied. I have not yet found a literature commenting on or assessing the Pena & Slate procedure (2006), so use of this method is only recommended with reservation. It does serve as a good example of how new/novel methods become rapidly available in R and are sometimes quite simple to implement.

```
# both the package and the function are called 'gulma'
#library(gulma)
# the gulma function only requires a 'lm' model fit object as a single argument to yield the full asses
gvlmafit3 <- gvlma(fit3)</pre>
#print out the tests
gvlmafit3
##
## Call:
## lm(formula = salary ~ pubs + cits, data = cohen1)
##
## Coefficients:
##
   (Intercept)
                       pubs
                                     cits
##
       40493.0
                      251.8
                                    242.3
##
##
## ASSESSMENT OF THE LINEAR MODEL ASSUMPTIONS
## USING THE GLOBAL TEST ON 4 DEGREES-OF-FREEDOM:
## Level of Significance = 0.05
##
## Call:
   gvlma(x = fit3)
##
##
##
                       Value p-value
                                                     Decision
## Global Stat
                      1.8686 0.7599 Assumptions acceptable.
## Skewness
                              0.4727 Assumptions acceptable.
                      0.5158
## Kurtosis
                      0.2567
                              0.6124 Assumptions acceptable.
## Link Function
                              0.6509 Assumptions acceptable.
                      0.2047
## Heteroscedasticity 0.8914
                              0.3451 Assumptions acceptable.
#plot relevant information
# note: this plot does not play nice in the R markdown format.
    So I generated it outside of the .Rmd file, saved the image and displayed the image below
```

#plot(gvlmafit3)

It is possible to generate a set of very useful plots by passing the gvlma object just created to the base system plot function (plot(gvlmafit3) as was commented out in the code chunk just above). However, the figure doesn't play nice with rmarkdown, so I created it separately, and saved the figure to a .png file that is displayed here.



10 Further capabilities in R: Assumptions, Diagnostics, Model Criticism

At this point in this document, we have covered essentially all of the methods that we first utilized in SPSS and have added several others into the arsenal of tools. The document simultaneously tried to weave conceptual integration/review into the templates for accomplishing the analyses with R. The remainder of the document extends the set of analyses to things not yet covered in the course. This includes resampling methods, regression diagnostics and influence analysis, model criticism, models with more than two IVs, and models with categorical variables. Much of the following are presented with the dual goals of R instruction and further consolidation of the conceptual framework that comprises linear modeling with multiple IV's and serves as a template for conceptual elaboration of those topics at later points in time.

11 Regression diagnostics and Influence Analysis

In a future section of the course, we will look more carefully at procedures for diagnosing problem issues in linear model fits, including the violation of assumptions. We will also build concepts that address how influential specific cases and variables are for the particular outcome that our linear modeling produces. This section of the current document will simply present code for a long list of these things without much comment, annotation, or explanation. It serves to put these capabilities in place for the point in time when we will need them The illustrations are all based on the fit3 model evaluated above:

11.1 A set of standard influence and diagnostic statistics

Several indices are calculated here that produce values for each case in the data set. I show how, with subsetting, to obtain individual case values for some of these (dfbetas and dffits), but don't print all values for any of them. The last code line in this chunk reminds us how to use headtail to examine the first few and last few elements of an object - dfbetas are shown.

```
#
  Diagnostics
#
  create the whole suite; look at each by doing, e.g., print(sresids)
sresids <- rstandard(fit3) #store the standardized residuals in a variable named "sresids"
standresid <- stdres(fit3) #store the standardized residuals in a variable named "standresids"
stud.del.resids <- rstudent(fit3) #store the studentized deleted residuals in a variable named "stud.de
leverage_hats <- hatvalues(fit3) #get the leverage values (hi)</pre>
cooksD <- cooks.distance(fit3) #get Cook's distance</pre>
dfbetas <- dfbetas(fit3) #calculate all dfbetas
dfbetas_4_0 \leftarrow dfbetas(fit3)[4,1] #dfbeta for case 4, first coefficient (i.e., b_0)
dffits <- dffits(fit3) #All dffits
dffits_4 <- dffits(fit3) [4] #dffits for case 4
influence <- influence(fit3) #various influence statistics, including hat values and dfbeta (not dfbeta
# examine dfbetas, for example
headtail(dfbetas)
```

```
##
        (Intercept)
                            pubs
                                         cits
## 1
      0.0081286885 0.019596957 -0.055510624
## 2
       0.1608435212 -0.105631169 -0.061098985
## 3
      0.0009316857 -0.007992130 0.005670943
##
## 59 -0.0742222603 0.056792584 0.011273927
## 60 0.2251262055
                    0.277341258 -0.484391375
## 61 -0.1449618287 -0.126384177 0.158434883
## 62 0.2185338279 -0.150682252 -0.072804543
```

Variance Inflation factors are useful indices for assessing multicolinearity. In a two-IV model such as this one, these will not be very interesting unless the two IVs are extremely highly correlated. And for a two-IV model the VIFs will be the same for each IV, since there is only one other IV to be correlated with for each of them (and it is the same correlation). So, this illustration is not very informative beyond showing how to obtain the VIFs.

```
#library(car) #load the package car if not still loaded from above
vif <- vif(fit3) #variance inflation factors
vif
## pubs cits
## 1.12505 1.12505
# one rule of thumb is to look for square root of vif values > 2
vif(fit3)^.5
```

pubs cits ## 1.060684 1.060684

vif(fit3)^.5 > 2

pubs cits
FALSE FALSE

Added variable plots are a tool that permits examination of the partial correlations between pairs of variables. Each of the two variables in a plot has been residualized on all other variables.

avPlots(fit3) #added variable plots



This code chunk creates a plot of Cooks D values (an influence statistic) as a function of sequential case number in the data set. It also identifies extreme cases (case numbers labeled) based on a threshold set in the threshold object. As discussed in class, it is best to look for Cook's D values divergent from others rather than those exceeding a numerical criterion, but some criteria have been proposed and one of those defines the threshold here.

```
# Cook's D plot
# identify D values > 4/(n-k-1)
threshold <- 4/((nrow(cohen1)-length(fit3$coefficients)-2))
plot(fit3, which=4, cook.levels=threshold)</pre>
```



This influence plot is a standard scatterplot of residuals against yhats, and is the plot used to visually assess heteroscedasticity. But the added benefit here is that the size (area) of each point is scaled to the Cook's D statistic thus permitting a visualization of the most influential cases in their residual/yhat position.



Influence Plot

Hat–Values Circle area is proportial to Cook's Distance

##		StudRes	Hat	CookD
##	24	-0.6018576	0.28463388	0.04856737
##	28	-2.4663093	0.07292105	0.14683221
##	32	2.4982801	0.03657164	0.07253089
##	48	2.5025997	0.10097428	0.21527380
##	53	0.8821127	0.13170293	0.03949029

Fox' **car** package has a useful function that identifies outliers and evaluates their likelihood of occurence with a p value. But doing this for N cases creates a multiple testing problem, so a Bonferroni adjusted p value is also presented. In this data set no exreme outliers were detected. This **outlierTest** function is evaluating outlier status relative to th studentized residuals.

#library(car) # if not loaded previously
outlierTest(fit3) # Bonferonni p-value for most extreme obs\

```
## No Studentized residuals with Bonferroni p < 0.05
## Largest |rstudent|:
## rstudent unadjusted p-value Bonferroni p
## 48 2.5026 0.015169 0.94046</pre>
```

We have already used the qq normal plot of residuals in many places. We should not forget that it is a

diagnostic tool and thus properly included in this chapter.

qqPlot(fit3, main="QQ Plot", distribution="norm", id=F) #qq normal plot for studentized resid



QQ Plot

The leverage plots shown here appear to be identical to the added variable plots shown above, and they are. The illustration here is done as a placeholder for later analyses where some IVs (such as categorical IVs with more than two levels) would have more than 1 df. In that case, this plot is useful, in addition to added variable plots.

library(car) # the function is from the car package leveragePlots(fit3) # leverage plots



A method of choice for addressing any violations of assumptions is often simple scale transformation of either the DV or the IVs, or both. We covered this idea with the household water use data set. We will revisit the scale transformation topic again at a later point in the course and will introduce Tukey's ladder of scales and Box-Cox transformation. Here a quick implementation of Box-Cox transformation capabilities is introduced. In this instance, the goal is to find an exponent to use to transform the DV to help produce normality and homoscedasticity of the residuals. The peak of the curve in the plot yields λ , the scaling component.

```
# scale transformations of the DV and/or IVs can help alleviate the impact of many issues
# #evaluate possible Box-Cox transformations (MASS package must be installed)
# look for the lambda at the log likelihood peak
#library(MASS)
boxcoxfit3 <- boxcox(fit3,plotit=T)</pre>
```



To find the exact λ value the output from the boxcox function was placed in a data frame and then sorted according to the log-likelihood value. The first case in the resorted dataframe gives the correct λ value. Note that the X and Y values are those used to create the plot above.

```
cox = data.frame(boxcoxfit3$x, boxcoxfit3$y)  # Create a data frame with the results
#str(cox)
cox2 <- cox[with(cox, order(-cox$boxcoxfit3.y)),] # Order the new data frame by decreasing y
#str(cox2)
cox2[1,] # Display the lambda with the greatest log likelihood</pre>
```

boxcoxfit3.x boxcoxfit3.y
59 0.3434343 -3.531357

For this model, λ is was found to be about .343. This value would be used to transform the DV and the analysis rerun. But recall that the normality and homoscedasticity assumptions didn't appear to be violated so this process would be unnecessary for the current data set. I show the code just to provide a template. Notice that an exponent of 1.0 would not be a scale change. The exponent of .343 is approximately equivalent to the cubed root. Also note that with the scale change, the values of the regression coefficients would become less readily interpretable.

```
#tsalary <- salary**(cox2[1,]$boxcoxfit3.x)
#hist(tsalary)
#fit3t <- lm(tsalary~cohen1$pubs+cohen1$cits)
#summary(fit3t)
#qqPlot(fit3$residuals, distribution="norm")</pre>
```

12 Resampling and Robust methods for Linear Models

Two classes of alternative approaches to linear modeling are breifly described in this section. The first is bootstrapping which is an alternative to std error calculation when faced with non-normal residuals. Although there are many "flavors" of bootstrapping, only two are illustrated here. The second class is a broad range of methods falling under the rubric of Robust Regression. The ones emphasized here are those that can help when heteroscedasticity is present in an OLS model. The code is not accompanied by much commentary or explanation. It is presumed that background on bootstrapping methods is obtained elsewhere.

12.1 Ordinary nonparametric bootstrapping with the boot function

The **boot** package has extensive facilities for bootstrapping many kinds of analyses. The essential logic is to tell the **boot** function which statistic from the analysis needs to be bootstrapped, what kind of bootstrapping (e.g., casewise vs residual), and then requests for summaries of the bootstrapping. This first section is based somewhat on the methods outlined in the Quick-R website [https://www.statmethods.net/advstats/bootstr apping.html]. The methods presented here are nonparametric bootstrapping methods.

Each statistic of interest can be bootstrapped. But this requires writing a function to extract that statistic from the lm fit so that it can be repeated with the bootstrap samples. This section will do bootstrap analyses on three statistics:

- 1. The Multiple R-Squared from the two-IV model used throughout this doc.
- 2. The pvalue for the F test of that Multiple R-squared (test of the "whole" equation).
- 3. The regression coefficients from the two-IV equation.

In these illustrations, I ran the boot function (creating the results object) within a call to the system.time function to record how long it took to do the bootstrap resampling. This will vary across platforms.

12.1.1 Bootstrap the multiple R-squared statistic

```
#library(boot)
# Bootstrap 95% CI for R-Squared
\# function to obtain R-Squared from the data
multrsq <- function(formula, data, indices) {</pre>
     d <- data[indices,] # allows boot to select sample</pre>
     fit <- lm(formula, data=d)</pre>
     return(summary(fit)$r.square)
}
# bootstrapping with 2000 replications
# for other kinds of statistics, `boot` may need a "maxit" argument to set the upper limit on iteratio
set.seed(1234) # for reproducibility within this document
system.time(results1 <- boot(data=cohen1, statistic=multrsq,</pre>
   R=2000, formula=salary~pubs+cits))
##
            system elapsed
      user
##
      1.67
              0.00
                       1.67
# view results
results1
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = cohen1, statistic = multrsq, R = 2000, formula = salary ~
```

```
## pubs + cits)
##
##
##
## Bootstrap Statistics :
## original bias std. error
## t1* 0.4195157 0.0118493 0.09665115
plot(results1)
```



```
# show 95% confidence interval
boot.ci(results1, type=c("norm","basic","perc","bca"))
```

```
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 2000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = results1, type = c("norm", "basic", "perc",
       "bca"))
##
##
## Intervals :
## Level
              Normal
                                  Basic
## 95%
         (0.2182, 0.5971)
                               (0.2378, 0.6161)
##
             Percentile
## Level
                                   BCa
                               (0.1806, 0.5768)
## 95%
         (0.2230, 0.6013)
## Calculations and Intervals on Original Scale
```

12.1.2 Bootstrap the omnibus F test p value

```
fpval <- function(formula, data, indices) {</pre>
     d <- data[indices,] # allows boot to select sample</pre>
     fit <- lm(formula, data=d)</pre>
     return(pf(summary(fit)$fstatistic[1],
               df1=summary(fit)$fstatistic[2],
               df2=summary(fit)$fstatistic[3],lower.tail=F))
}
# bootstrapping with 2000 replications
# for other kinds of statistics, `boot` may need a "maxit" argument to set the upper limit on iteratio
set.seed(1234) # for reproducibility within this document
system.time(results2 <- boot(data=cohen1, statistic=fpval,</pre>
   R=2000, formula=salary~pubs+cits))
##
      user system elapsed
##
      2.04
              0.00
                      2.05
# view results
results2
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = cohen1, statistic = fpval, R = 2000, formula = salary ~
##
       pubs + cits)
##
##
## Bootstrap Statistics :
##
           original
                          bias
                                   std. error
## t1* 1.076015e-07 0.0003175022 0.006625147
plot(results2)
```



show 95% confidence interval boot.ci(results2, type=c("bca"))

```
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 2000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = results2, type = c("bca"))
##
## Intervals :
## Level BCa
## 95% ( 0.0000, 0.0041 )
## Calculations and Intervals on Original Scale
```

12.1.3 Bootstrap the regression coefficients

Next, we will bootstrap the regression coefficients

```
# Bootstrap 95% CI for regression coefficients
#library(boot)
# function to obtain regression weights
coeffs <- function(formula, data, indices) {
    d <- data[indices,] # allows boot to select sample
    fit <- lm(formula, data=d)
    return(coef(fit))
}
# bootstrapping with 2000 replications</pre>
```

```
# for other kinds of statistics, `boot` may need a "maxit" argument to set the upper limit on iteratio
set.seed(1234) # for reproducibility within this document
system.time(results3 <- boot(data=cohen1, statistic=coeffs,</pre>
        R=2000, formula=salary~pubs+cits))
##
      user
            system elapsed
##
      1.36
              0.00
                      1.38
# view results
results3
##
## ORDINARY NONPARAMETRIC BOOTSTRAP
##
##
## Call:
## boot(data = cohen1, statistic = coeffs, R = 2000, formula = salary ~
##
       pubs + cits)
##
##
## Bootstrap Statistics :
         original
                               std. error
##
                      bias
## t1* 40492.9715 -92.356087
                               2481.39738
## t2*
         251.7500 -2.286656
                                 85.13455
## t3*
         242.2977
                    2.522431
                                 58.03025
plot(results3, index=1) # intercept
```




Histogram of t

plot(results3, index=3) # disp



```
# get 95% confidence intervals
boot.ci(results3, type="bca", index=1) # intercept
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 2000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = results3, type = "bca", index = 1)
##
## Intervals :
## Level
               BCa
         (35539, 45437)
## 95%
## Calculations and Intervals on Original Scale
boot.ci(results3, type="bca", index=2) # pubs
## BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
## Based on 2000 bootstrap replicates
##
## CALL :
## boot.ci(boot.out = results3, type = "bca", index = 2)
##
## Intervals :
## Level
               BCa
## 95%
         (86.9, 424.7)
## Calculations and Intervals on Original Scale
```

boot.ci(results3, type="bca", index=3) # cits
BOOTSTRAP CONFIDENCE INTERVAL CALCULATIONS
Based on 2000 bootstrap replicates
##
CALL :
boot.ci(boot.out = results3, type = "bca", index = 3)
##
Intervals :
Level BCa
95% (112.6, 340.1)
Calculations and Intervals on Original Scale

12.2 An alternative bootstrap approach from car package functions

John Fox has produce the **car** package that we have seen contains many useful functions for linear modeling (Fox, 2016). One function, Boot, provides a flexible approach to non-parametric bootstrapping that may be easier to use. I also find that the Confint function is a quicker way to get a full picture of bootstrapped confidence intervals.

```
# Bootstrap 95% CI for regression coefficients
#library(boot)
# function to obtain regression weights
coeffs <- function(formula, data, indices) {</pre>
  d <- data[indices,] # allows boot to select sample</pre>
  fit <- lm(formula, data=d)</pre>
  return(coef(fit))
}
# bootstrapping with 2000 replications
# for other kinds of statistics, `boot` may need a "maxit" argument to set the upper limit on iteratio
set.seed(1234) # for reproducibility within this document
system.time(results4 <- Boot(fit3, f=coef,</pre>
        R=2000, method=c("case"))) # can specify casewise or residual sampling with the method argument
##
      user
           system elapsed
##
      1.44
              0.00
                       1.45
# view results
summary(results4, high.moments=T)
##
## Number of bootstrap replications R = 2000
##
               original bootBias
                                    bootSE bootMed
                                                       bootSkew bootKurtosis
## (Intercept) 40492.97 -92.3561 2481.397 40430.85 -0.1058512
                                                                     0.314662
## pubs
                 251.75
                         -2.2867
                                    85.135
                                              248.05 0.0016417
                                                                     0.013766
                 242.30
                           2.5224
                                    58.030
                                              248.02 -0.1261283
                                                                     0.134424
## cits
```

Bootsrapped empirical distributions for each coefficient are produced with the application of the base system plot function to the bootstrapped results.



Histogram of t

plot(results4, index=2) # wt



plot(results4, index=3) # disp



And confidence intervals of multiple types are readily produced. Percentile ("perc") and "bca" are typically recommended.

```
# get 95% confidence intervals
Confint(results4, type="norm")
## Bootstrap normal confidence intervals
##
##
                 Estimate
                                 2.5 %
                                           97.5 %
## (Intercept) 40492.9715 35721.87804 45448.7770
## pubs
                 251.7500
                              87.17604
                                         420.8973
## cits
                 242.2977
                             126.03808
                                         353.5125
Confint(results4, type="perc")
## Bootstrap percent confidence intervals
##
##
                 Estimate
                                 2.5 %
                                           97.5 %
## (Intercept) 40492.9715 35280.22022 45274.3103
## pubs
                 251.7500
                              77.56269
                                         415.7893
## cits
                 242.2977
                             127.95169
                                         353.4045
Confint(results4, type="bca")
## Bootstrap bca confidence intervals
##
##
                 Estimate
                                 2.5 %
                                           97.5 %
```

```
## (Intercept) 40492.9715 35539.48858 45436.7048
```

##	pubs	251.7500	86.91784	424.6978
##	cits	242.2977	112.56818	340.1057

```
# obtain a useful set of histograms for each coefficient distribution
hist(results4, legend="separate")
```



vcov(results4)

##		(Intercept)	pubs	cits
##	(Intercept)	6157332.98	-47605.756	-112384.905
##	pubs	-47605.76	7247.891	-1593.342
##	cits	-112384.90	-1593.342	3367.510

12.3 Robust Regression for heteroscedastic data sets

Bootstrapping methods are primarily employed as a tool to handle situations where normality assumptions are violated. They are not necessarily insensitive to heteroscedasticity issues. In this section, I outline tools available from the **car**, **Imtest**, **MASS**, and **sandwich** packages for robust regression analyses. Also included is a brief introduction to a suite of robust methods for handling data sets with outliers and influential cases.

12.3.1 A robust Standard Errors approach from the car Package when heteroscedasticity is present.

The Salary-pubs-cits data set was not one that showed much heteroscedasticity, so submitting it to robust methods may not be necessary. Another data set we have see previously does have heteroscedasticity and this is what is used here. This is a data set provided by Dr. James Boswell on baseline scores of anxietydiagnosed patients on scales of depression severity (bods, treated as the DV), anxiety severity (boasev), and

term	estimate	std.error	statistic	p.value
(Intercept)	5.7363514	1.6676431	3.439796	0.0007105
boasev	0.5829990	0.0775505	7.517665	0.0000000
bpa	-0.2104816	0.0405446	-5.191359	0.0000005

Table 2: Regression Coefficients and Regular Parametric Std Errors

positive affect (bpa). A standard two-IV linear model is fit and then corrected.

boswell <- read.csv("data/baselinevars.csv")</pre>

The two-iv regression produced a multiple R-squared of about .42 and the standard tests of the two IVs are shown here. Note that the data set has a fairly large N (200) and thus substantial power for effects of the size seen.

The plot of residuals vs yhats reveals an odd pattern, that reflects a truncated variable (boasev). But it also reveals a modest amount of heteroscedasticity.

plot(fit10, which=1)



```
#qqPlot(fit10$residuals, distribution="norm", id=F)
```

Testing a null hypothesis of no heteroscedasticity with the Breusch-Pagan test finds a p value in the rejection range.

term	estimate	std.error	statistic	p.value
(Intercept)	5.7363514	1.9687184	2.913749	0.0039840
boasev	0.5829990	0.0984717	5.920470	0.0000000
bpa	-0.2104816	0.0454911	-4.626872	0.0000067

Table 3: Robust (HC0) standard errors and modified t-tests

#library(lmtest) bptest(fit10)

```
##
## studentized Breusch-Pagan test
##
## data: fit10
## BP = 16.966, df = 2, p-value = 0.0002069
```

Several different methods of estimating corrected covariance matrices based on heteroscedasticity are available. The one illustrated here ("HCO") is often simply called the White Std Error. See the help doc on hccm for more details. The hccm function is from the **car** package and **coeftest** is from the **lmtest** package. You can see that the standard errors are different than the regular standard errors, but they are not much larger owing to the fact that heteroscedasticity is not severe in this data set.

12.3.2 The Wald test and an alternative to adjusted t-tests for linear models with heteroscedasticity

The **lmtest** package has a function called **waldtest** that permits a commonly used inferential method, the Wald Test, which also employs covariance matrix correction for heteroscedasticity. It uses a model comparison strategy. First we created an intercept only model (mod0) and the the full two-IV model (modfull). **waldtest** then compares those two models in a standard model comparison methodology and the F test evalutes the improvement in fit, but using corrected standard errors of the White type (HC0). This F value is somewhat smaller than the one seen above for the regular OLS method, and is the alternate method for testing the null about the overall fit of the model. The vcovHC function comes from the **sandwich** package and is a common one for estimating the covariance structure in the presence of heteroscedasticity.

```
#library(lmtest)
#library(sandwich)
mod0 <- lm(bods~1 , data=boswell)</pre>
modfull <- lm(bods~boasev+bpa , data=boswell)</pre>
waldtest(mod0, modfull, vcov = vcovHC(modfull, type = "HCO"))
## Wald test
##
## Model 1: bods ~ 1
## Model 2: bods ~ boasev + bpa
     Res.Df Df
##
                     F
                          Pr(>F)
## 1
        199
## 2
        197 2 57.232 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

The **lmtest** package also has a function to test the individual regression coefficients with the adjusted standard errors. This approach is identical to the one shown above with the **car** package tools. The **vcovHC** function comes from the **lmtest** package. Adjusted standard errors and t's are the same as above since the same White Std. Errors are chosen.

```
#library("lmtest")
#library("sandwich")
# Robust t test
coeftest(modfull, vcov = vcovHC(modfull, type = "HCO"))
##
## t test of coefficients:
##
##
                Estimate Std. Error t value
                                             Pr(>|t|)
                5.736351
                           1.968718
                                     2.9137
                                              0.003984 **
## (Intercept)
## boasev
                0.582999
                           0.098472 5.9205 1.406e-08 ***
## bpa
               -0.210482
                           0.045491 -4.6269 6.719e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

12.4 Robust regression methods when outliers or highly influential cases are present

When outliers or highly influentical cases are detected via influence analysis it is possible to use that information to refine the linear model. The general idea is to take an influence statistic such as Cook's D and use it to weight the importance of cases in the regression. One commonly recommended method is called "Iteratively Reweighted Least Squares". The rlm function in the **MASS** package is a recommended method to perform such weighted/reweighted least squares. Rather than take additional space in this document, the reader is referred to a very good tutorial page on IRLS using rlm. This tutorial page is from the UCLA Statistical Consulting unit which has created many useful tutorials.

[https://stats.idre.ucla.edu/r/dae/robust-regression/]

13 Extensions to larger models and to categorical IVs

In this chapter, the goals are to extend the suite of methods to models with additional characteristics. This includes a model with more than two IVs but we will also consider using a categorical IV toward the end. At the same time, this chapter provides some emphasis on strong capabilities in rmarkdown and the "tidyverse" to generate nicer tables. Some of that has been seen in other chapters with the kable and gt functions, but it is expanded here.

The capability of **rmarkdown**, **bookdown** and **knitr** to produce a document such as this one is accompanied by increased options for formatting the output. The typical text output from R functions can be converted to style/fonts that are more readable. A few brief examples are illustrated in this section. The kable function permits enhancement of tables such as those coming from **summary** and **anova**. But capabilities in the **broom** package also permit enhancement of the standard output coming from 'summary. The reader should recognize that whole documents/manuscripts can be written in rmarkdown, even in APA style

13.1 Evaluate a 3-IV model

We will use the same data set that we used above with the enhanced output and formatting. However we will extend the analysis to three IVs. The reader should take note of the fact that adding a third variable is simply done in the model specification argument for the lm function and that such functions as summary and anova or Anova would be done the same way as previously executed:

```
fit6 <- lm(salary~pubs+cits+degree_yrs, data=cohen1) # add degree_yrs as a 3rd IV to our original pubs
summary(fit6)
##
## Call:
## lm(formula = salary ~ pubs + cits + degree_yrs, data = cohen1)
##
## Residuals:
##
                 1Q Median
       Min
                                  ЗQ
                                          Max
## -13907.1 -4313.8 -649.5 4366.1 21165.3
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 38967.85
                        2394.31 16.275 < 2e-16 ***
## pubs
                 93.61
                         85.35
                                  1.097 0.277273
## cits
                204.06
                           56.97 3.582 0.000699 ***
                          283.89 3.080 0.003160 **
## degree_yrs
                874.46
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 7031 on 58 degrees of freedom
## Multiple R-squared: 0.5011, Adjusted R-squared: 0.4753
## F-statistic: 19.42 on 3 and 58 DF, p-value: 7.727e-09
mrinfo(fit6, minimal=TRUE)
## [[1]]
## NULL
##
## $`supplemental information`
##
             beta wt structure r partial r semipartial r tolerances unique
## pubs
             0.13506
                       0.71500 0.14254
                                           0.10172
                                                          0.56721 0.01035
                                                          0.84665 0.11035
## cits
             0.36102
                        0.77662 0.42559
                                               0.33219
                        0.85873 0.37495
                                              0.28567
## degree_yrs 0.38541
                                                          0.54940 0.08161
              common total
##
## pubs
             0.24584 0.25618
             0.19190 0.30224
## cits
## degree_yrs 0.28793 0.36954
##
## $`var infl factors from HH:vif`
##
      pubs cits degree_yrs
    1.763010 1.181128 1.820156
##
##
## [[4]]
## NULL
Anova(fit6, type=3)
## Anova Table (Type III tests)
##
## Response: salary
##
                  Sum Sq Df F value
                                       Pr(>F)
## (Intercept) 1.3093e+10 1 264.8822 < 2.2e-16 ***
## pubs
             5.9458e+07 1 1.2029 0.2772728
              6.3412e+08 1 12.8291 0.0006989 ***
## cits
```

```
4.6897e+08 1
                               9.4878 0.0031599 **
## degree yrs
## Residuals
               2.8669e+09 58
## ---
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
anova(fit3,fit6)
## Analysis of Variance Table
##
## Model 1: salary ~ pubs + cits
## Model 2: salary ~ pubs + cits + degree_yrs
##
    Res.Df
                   RSS Df Sum of Sq
                                         F Pr(>F)
## 1
         59 3335822387
## 2
         58 2866853951 1 468968436 9.4878 0.00316 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

It is also useful to examine the Variance Inflation Factors in multiple regression models. Unlike the two-IV model, each IV now has a different VIF since each has a different tolerance. Note that the IV with the smallest tolerance has the largest VIF.

vif(fit6)

pubs cits degree_yrs
1.763010 1.181128 1.820156

Use the 'tidy' function from the 'broom' package to produce an alternative to the 'summary' table, and format it with 'kable':

```
fit6.tidy <- tidy(fit6) # tidy produces a nicer table than summary
kable(fit6.tidy) # kable converts the table to markdown format that is nicer than the default</pre>
```

term	estimate	std.error	statistic	p.value
(Intercept)	38967.84674	2394.30824	16.275201	0.0000000
pubs	93.60794	85.34837	1.096775	0.2772728
cits	204.06019	56.97179	3.581776	0.0006989
degree_yrs	874.46140	283.89492	3.080229	0.0031599

I actually prefer using the gt function for tables whenever I can. But sometimes gt doesn't play nice with rmarkdown when rendering to a pdf document. For example, using it in the code chunk here, with a tidy object, creates a problem. I show the code commented so that you can see how to use it in other formats. The same analysis is tabled here using gt.

#gt(fit6.tidy) # `gt` converts the table to markdown format that is nicer than the default

Use the glance function from the 'broom' package to obtain some additional information and then use 'kable' to format the output table. Note that I've relabeled the components of the glance object to fit better with our typically used notation schemes. I also broke the table into two components to better fit on the page. This latter strategy produced a printing of the pvalue that lost the exponential notation and I've not yet figured out how to prevent that. It appears to be an issue with kable, so I have printed the two tables without kable formatting.

```
#library(broom)
glancefit6a <- glance(fit6) # additional info on the model
glancefit6at1 <- glancefit6a[1:5]
#glancefit6at2
glancefit6at2 <- glancefit6a[6:11]
#glancefit6at2
kable(glancefit6at1,</pre>
```

col.n	ames=c("R.squa	ared", "ad	j.R.squar	ed", "RMS	SE","F value", "p.value"))		
R.squared	adj.R.squared	RMSE	F value	p.value	_		
0.5011234	0.4753195	7030.542	19.42041	0	-		
kable(glancefit6at2,							
<pre>col.names=c("df regression","logLik", "AIC", "BIC", "deviance", "df residual"),digits=3)</pre>							
df regression	n logLik	AIC	BIC	deviance	e df residual		

	df regression	logLik	AIC	BIC	deviance	df residual
3 -635.104 1280.208 1290.844 2866853951	3	-635.104	1280.208	1290.844	2866853951	58

Notes USING GLANCE:

1. Users of the **broom** package should be aware that what **glance** previously reported as "df" would not have been the expected regression df that would equal the number of IVs. Instead it reported one more than that, including the df for the intercept. In the model above with 3 IVs, that apparently erroneous df was reported as 4, which is actually the rank of the design matrix. But the F test was always the standard MSreg/MSresidual, so it was confusing. That has been corrected in the most recent version used here (ver 0.7.0) so that the 3 df seen in the table here is the correct DF regression for a three-IV model. Users who installed the package prior to the correction of this "feature" should update their installation.

This document is created with R4.0.2

- 2. The standard glance table is too wide it has too many columns/values. So, I extracted two different subsets of its components and produced two different tables in the commented out code lines using kable.
- 3. I also relabeled the labels (in those commented out kable code lines) for the columns of the first table to be more in line with our standard terminology (RMSE is root mean square error, or the square root of MS residual)

9.487811

NA

0.0031599

NA

Next, we use 'tidy' and 'kable' to obtain a nicely formatted anova summary table:

tidyanova6 ·	<- ti	.dy(anova(fit	t6))			
kable(tidya	nova6	5,				
col.na	ames=	c("Source",	"df", "SS",	"MS", "F v	value", "p	value"))
						_
Source	df	SS	MS	F value	p value	
pubs	1	1472195326	1472195326	29.784332	0.0000010	_
cits	1	938602110	938602110	18.989081	0.0000544	_

degree_yrs	1	468968436	468968436
Residuals	58	2866853951	49428516

Or use the 'Anova' function:

```
tidyAnova6 <- tidy(Anova(fit6, type=3))</pre>
kable(tidyAnova6,
```

col.names=c("Source", "SS", "df", "F value", "p value"))

Source	SS	df	F value	p value
(Intercept)	13092731852	1	264.882153	0.0000000
pubs	59458298	1	1.202915	0.2772728
cits	634124345	1	12.829119	0.0006989
degree_yrs	468968436	1	9.487811	0.0031599
Residuals	2866853951	58	NA	NA

Now add information from the 'mrinfo' function that was discussed above, extracting the additional useful information that we need for a fuller analysis than summary/anova/Anova gives us.

```
mrinfo(fit6, minimal=TRUE)
## [[1]]
## NULL
##
## $`supplemental information`
##
              beta wt structure r partial r semipartial r tolerances unique
              0.13506
                           0.71500
                                     0.14254
                                                    0.10172
                                                                0.56721 0.01035
## pubs
                                      0.42559
                                                    0.33219
                                                                0.84665 0.11035
## cits
              0.36102
                           0.77662
  degree_yrs 0.38541
                                      0.37495
                                                    0.28567
                                                                0.54940 0.08161
                           0.85873
##
##
               common
                         total
## pubs
              0.24584 0.25618
## cits
              0.19190 0.30224
## degree_yrs 0.28793 0.36954
##
##
   $`var infl factors from HH:vif`
##
         pubs
                     cits degree_yrs
##
     1.763010
                 1.181128
                            1.820156
##
## [[4]]
## NULL
```

13.1.1 Conclusions from the three-IV model

Several interesting outcomes emerged with the addition of degree_yrs to the model that already included pubs and cits. The multiple R-squared did increase to about .50 so degree_yrs did appear to improve predictability. But an interesting thing happened to the effect of pubs in this three-IV model. The regression coefficient was about 251 in the two-IV model, but it dropped to about 94 in this three-IV model. Pubs was not a significant predictor in the three-IV model and it's unique fraction of variance dropped to about 1% of the DV variation. Most of the overlap of pubs with salary was shared by the other IV's (common fraction of variance was about .25, or nearly all of its zero order overlap with salary). This is because degree_yrs must have been strongly correlated with pubs. Lets examine that correlation.

with(cohen1, cor(pubs,degree_yrs))

```
## [1] 0.6505472
```

A better model might be one that simply leaves pubs out since degree_yrs absorbed most of its shared variance.

```
fit6b <- lm(salary~cits+degree_yrs,data=cohen1)
fit6b.tidy <- tidy(fit6b) # tidy produces a nicer table than summary
kable(fit6b.tidy)</pre>
```

term	estimate	std.error	statistic	p.value
(Intercept)	39073.6747	2396.47360	16.304655	0.0000000
cits	212.1116	56.59392	3.747958	0.0004078
$degree_yrs$	1061.7642	227.17563	4.673759	0.0000176

```
mrinfo(fit6b, minimal=TRUE)
```

```
## [[1]]
## NULL
##
## $`supplemental information`
## beta wt structure r partial r semipartial r tolerances unique
## cits 0.37526 0.78476 0.43852 0.3482 0.86094 0.12124
```

```
## degree_yrs 0.46796
                            0.86773
                                       0.51981
                                                       0.4342
                                                                  0.86094 0.18853
##
               common
                        total
## cits
                0.181 0.30224
## degree_yrs 0.181 0.36954
##
## $`var infl factors from HH:vif`
##
         cits degree yrs
                 1.161517
##
     1.161517
##
## [[4]]
## NULL
glancefit6b <- glance(fit6b)</pre>
glancefit6bt1 <- glancefit6b[1:5]</pre>
#glancefit6bt1
glancefit6bt2 <- glancefit6b[6:11]</pre>
#glancefit6bt2
kable(glancefit6bt1,
      col.names=c("R.squared", "adj.R.squared", "RMSE", "F value", "p.value"))
 R.squared
            adj.R.squared
                             RMSE
                                       F value
                                                p.value
 0.4907768
                 0.473515
                           7042.621
                                      28.43137
                                                     0
kable(glancefit6bt2,
      col.names=c("df regression","logLik", "AIC", "BIC", "deviance", "df residual"),digits=3)
```

df regression	logLik	AIC	BIC	deviance	df residual
2	-635.74	1279.481	1287.989	2926312249	59

The revised two-IV model with degree_yrs and cits as IVs had an R-squared nearly as large as the three-IV model, and the AIC/BIC indices were slightly smaller indicating a better model. Both degree_yrs and cits were significant IVs as tested by the t-tests. Thus the most parsimonious model might be this revised two-IV model. Perhaps pubs was only a good IV because it was correlated so highly with degree_yrs and it is the latter that is the more important predictor.

13.2 Categorical IVs

We can do a quick preview of using categorical IVs in linear models. The cohen data set also had a categorical variable, gender, that could potentially serve as an IV. However, it is a factor and its values in the data set are the string values of "female" and "male". How could that be used as an IV in regression? The answer is that factors can be recoded to have numeric values with specially constructed coded values. For a factor such as gender with only two levels the code is simply a one and a zero. This is already built in to the characteristics of the factor that is stored in the data frame. These numeric codes are called contrasts in R:

contrasts(cohen1\$gender)

male ## female 0 ## male 1

With that knowledge, we now see that lm can handle such categorical variables by using the numeric contrast code. First I will illustrate with a simple regression using only gender as the IV.

```
fit7 <- lm(salary~gender, data=cohen1)
summary(fit7)</pre>
```

Call:

```
## lm(formula = salary ~ gender, data = cohen1)
##
##
  Residuals:
                        Median
                                     ЗQ
##
        Min
                   1Q
                                              Max
##
   -18660.3
             -5819.8
                          -3.7
                                 4769.2
                                          26903.7
##
##
   Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
                  52650
                               1810
                                     29.080
                                               <2e-16 ***
##
   (Intercept)
##
   gendermale
                    3949
                               2445
                                      1.615
                                                0.111
##
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 9580 on 60 degrees of freedom
## Multiple R-squared: 0.04168,Adjusted R-squared: 0.0257
## F-statistic: 2.609 on 1 and 60 DF, p-value: 0.1115
```

It appears that gender is not a strong predictor of salary (R-squared is small, and t-test is NS). But lets focus on two additional points. One is notation. Observe that the name of the gender variable in the coefficients table is not simply gender, the IV name. Rather it is found there as "gendermale". That notation reminds us that gender is a factor and the reason that it is called gendermale rather than genderfemale is that male was coded with a 1 as seen above. The zero and one assignments are arbitrary and could be reversed. We will soon extend this conversation with more complex categorical IVs that have more than two categories.

The second point of emphasis is the thinking that surrounds the question of what does it mean to think of gender as "correlated" with salary, or that it can "predict" salary? Perhaps a better phrasing might be that we have assessed whether gender is "associated" with salary. We have concluded that it is not, but what if it were? That could only mean that average salaries might differ between females and males. And that sounds just like a two-independent-samples t-test situation. So let's do that test.

t.test(salary~gender, var.equal=T, data=cohen1)

```
##
##
    Two Sample t-test
##
## data: salary by gender
## t = -1.6153, df = 60, p-value = 0.1115
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
    -8839.888
##
                941.241
## sample estimates:
##
  mean in group female
                          mean in group male
##
               52650.00
                                     56599.32
```

Note that the male mean is about \$4000 higher than the female mean. But the difference was not significant. A bit of a closer examination of the output reveals interesting coincidences. Note that the df for the t, the t-value itself, and the p value match what was reported for the regression coefficient in the above lm fit. This is not coincidence because the "association" question approached with regression is tantamount to the mean difference question posed by the independent samples t-test. Soon enough, we will progress through the technical reasons that these are the same inference.

For one final illustration in this chapter, we now build a regression model that adds gender to the earlier model that had degree_yrs and citations as IVs.

```
fit7b <- lm(salary~degree_yrs + cits + gender, data=cohen1)
summary(fit7b)</pre>
```

```
##
## Call:
##
  lm(formula = salary ~ degree_yrs + cits + gender, data = cohen1)
##
##
  Residuals:
##
        Min
                       Median
                                     ЗQ
                  10
                                             Max
                       -312.3
##
   -14214.6 -4590.3
                                 4014.9
                                         21917.9
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
##
  (Intercept) 38787.21
                           2478.78
                                     15.648 < 2e-16 ***
                             232.58
                                      4.473 3.64e-05 ***
##
  degree_yrs
                1040.41
## cits
                 210.14
                              57.09
                                      3.681 0.000512 ***
  gendermale
##
                 931.06
                            1859.70
                                      0.501 0.618513
##
  ___
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 7088 on 58 degrees of freedom
## Multiple R-squared: 0.493, Adjusted R-squared: 0.4667
## F-statistic: 18.8 on 3 and 58 DF, p-value: 1.227e-08
```

In this model gender is also not a significant IV and the regression coefficient is smaller than it was in simple regression indicating that any variance that it shared with salary was at least partially confounded with degree_yrs and cits.

One final comment about this type of modeling with a variable such as gender. It might be an interesting research question whether the predictability that IVs such as degree_yrs and cits have on salary differs in the two levels of a categorical variable such as gender. In other words, does the influence of degree_yrs and cits depend on whether we examine males or females? This is what we will come to call an interaction question. It can be modeled easily with lm. I will show the code/results for the three-IV model with all interactions as well as the results, but will save comment until we cover the interaction topic more explicitly.

```
fit7c <- lm(salary~degree_yrs*cits*gender, data=cohen1)
summary(fit7c)</pre>
```

```
##
## Call:
## lm(formula = salary ~ degree_yrs * cits * gender, data = cohen1)
##
##
  Residuals:
##
        Min
                  10
                        Median
                                     30
                                              Max
##
   -13966.3
            -4404.6
                        -773.2
                                 3657.9
                                         22513.1
##
## Coefficients:
##
                               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                               26877.05
                                           8716.17
                                                      3.084 0.00322 **
                                3715.49
## degree_yrs
                                            1591.34
                                                      2.335
                                                             0.02330 *
## cits
                                 632.12
                                             243.37
                                                      2.597
                                                             0.01208 *
                                                             0.28643
## gendermale
                               11287.69
                                           10484.27
                                                      1.077
## degree_yrs:cits
                                 -89.18
                                              45.11
                                                     -1.977
                                                             0.05315
## degree_yrs:gendermale
                               -2442.09
                                            1793.28
                                                     -1.362
                                                             0.17892
## cits:gendermale
                                -403.30
                                                     -1.464
                                                             0.14897
                                             275.46
## degree_yrs:cits:gendermale
                                              47.53
                                  86.42
                                                      1.818 0.07457 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
```

```
## Residual standard error: 7051 on 54 degrees of freedom
## Multiple R-squared: 0.5328,Adjusted R-squared: 0.4722
## F-statistic: 8.798 on 7 and 54 DF, p-value: 3.563e-07
```

14 Efficiency in OLS analysis using the olsrr package

The **olsrr** package has a collection of functions that streamline many of the piecemeal approaches oulined in earlier chapters. It makes it simple to obtain extensive/detailed information at once, with user-friendly functions. Not all of the topics covered in earlier chapters are included (e.g., semi-partial correlations), but extensive capabilities are also included for model criticism.

14.1 A basic analysis

The initial use of the ols_regress function can replace the individual uses of summary, anova, and confint functions. The model fit here is the three-IV model examined in the "Extensions" chapter.

```
fit6 <- lm(salary~pubs+cits+degree_yrs, data=cohen1)
ols_regress(fit6)</pre>
```

## ##			Model	Summary					
## ## ## ##	R R-Squared	0 . 0 . 0 .	.708 .501	RMSE Coef.	Var	7030	.542 .826		
	Adj. R-Square	R-Squared 0.475	MSE		49428516	. 393			
## ##	Pred R-Square	d 0.	420	MAE		5317	.619		
## ## ## ##	# # RMSE: Root Mean Square Error # MSE: Mean Square Error # MAE: Mean Absolute Error								
## ## ##		ANOVA							
##									
## ##	Squares		DF	Mean Square	e F	F Sig.	Sig.		
##	Regression	2879765872.6	503	3	959921957.534	l 19.4	42 0.0	0000	
##	Residual	2866853950.7	768	58	49428516.393	3			
## ##	Total	5746619823.3	371	61					
## ## ##				Pa	rameter Estima				
##									
## ##	model	Beta	Std.	Error	Std. Beta	t	Sig	lower	upper
## ##	(Intercept)	38967.847	23	94.308		16.275	0.000	34175.118	43760.575
##	pubs	93.608		85.348	0.135	1.097	0.277	-77.235	264.451
##	cits	204.060		56.972	0.361	3.582	0.001	90.019	318.102
## ##	degree_yrs	874.461	2	83.895 	0.385	3.080	0.003	306.184	1442.739
##									

14.2 Collinearity diagnostics

The ols_coll_diag function provides collinearity diagnostics:

ols_coll_diag(fit6)

```
## Tolerance and Variance Inflation Factor
##
  ____
        _____
##
     Variables Tolerance
                             VIF
## 1
          pubs 0.5672118 1.763010
## 2
          cits 0.8466483 1.181128
## 3 degree_yrs 0.5494035 1.820156
##
##
## Eigenvalue and Condition Index
##
  _____
##
    Eigenvalue Condition Index
                                intercept
                                                                   degree_yrs
                                                pubs
                                                            cits
## 1 3.55809347
                      1.000000 0.009810827 0.014093645 0.009215181 0.0110334455
                      3.728942 0.146119575 0.358711061 0.097741038 0.0590237875
## 2 0.25588579
## 3 0.10745243
                      5.754407 0.015671115 0.619788632 0.026076066 0.9299270069
                      6.729533 0.828398483 0.007406662 0.866967715 0.0000157601
## 4 0.07856832
```

14.3 Added-variable plots

We can also obtain added-variable plots. These depict the partial correlations of each IV with the DV, both adjusted for other IVs. Here, it can be seen that pubs has a weaker partial relationship and this reinforces the fact that it's test in the 3-IV model was non-significant. Recall that the partial correlation can be obtained with the mrinfo2 function. Unfortunately, partial and semi-partial correlations are not provided with the otherwise extensive information coming from ols_regress.

```
ols_plot_added_variable(fit6)
```

```
## `geom_smooth()` using formula 'y ~ x'
## `geom_smooth()` using formula 'y ~ x'
## `geom_smooth()` using formula 'y ~ x'
```



```
## [[4]]
## NULL
```

14.4 Residual Assumptions

A normal QQ plot of the residuals is available.

ols_plot_resid_qq(fit6)



Tests of the residual normality assumption are easily obtained.

```
ols_test_normality(fit6)
```

##			
##	Test	Statistic	pvalue
##			
##	Shapiro-Wilk	0.9782	0.3374
##	Kolmogorov-Smirnov	0.0759	0.8404
##	Cramer-von Mises	5.2312	0.0000
##	Anderson-Darling	0.4327	0.2946
##			

A histogram of the residuals with a normal curve overlaid for comparison is also available.

ols_plot_resid_hist(fit6)



And the standard plot of residuals against yhats is also available. ols_plot_resid_fit(fit6)



Studentized and Standardized residuals can be examined to look for sequential ordering effects in the data set by plotting them against the case number.

ols_plot_resid_stud(fit6)



ols_plot_resid_stand(fit6)



Several of the above plots plus other model diagnostic plots can be obtained more rapidly with the **ols_plot_diagnostics** function for which the code is shown here. The plots are not returned in order to save space.

ols_plot_diagnostics(fit6)

14.5 Plots for examination of Influence

Several plots are available for visualizing the influence statistics for a model.

First is examination of Cook's D values. It provides a visual indicators of which cases exceed a threshold for large influence and those cases are numerically labeled. I have not yet sorted out how this threshold is determined for this function and the following ones.

ols_plot_cooksd_chart(fit6)



The DFBeta index is visualized with a panel of graphs, one for each IV and one for the intercept, permitting identification of influential cases for each IV separately.

ols_plot_dfbetas(fit6)



And a comparable plot for DFfits is also avaiable. ols_plot_dffits(fit6)



Finally, two additional plots are common in model diagnostics. They examine studentized residuals and deleted studentized residuals against leverage and yhats, respectively.

ols_plot_resid_lev(fit6)



Outlier and Leverage Diagnostics for salary

ols_plot_resid_stud_fit(fit6)



Deleted Studentized Residual vs Predicted Values

The **olsrr* package has numerous other tools and is worth exploring. The reference manual and vignettes on the CRAN site are very helpful.

https://cran.r-project.org/web/packages/olsrr/index.html

15 Bayes Factor approach to multiple regression

There are many approaches and flavors of Bayesian inference. A major set of tools is found in the Stan libraries. In R, these are implemented in the **rstan** package. There are extensive ways of doing inference for linear models (and many other models) using **rstan** tools. These approaches are often laborious, and require far more background in Bayesian methods than most readers of this document will have at this point in time. An alternative approach involves BUGS algorithms. They are implemented in WinBugs for the Windows platform and OpenBugs. An R package wwritten by Andrew Gelman is an interface to WinBugs or OpenBugs and is called R2WinBUGS. These packages are not illustrated here, but links to useful sites are:

[https://mc-stan.org/users/interfaces/rstan]

[https://cran.r-project.org/web/packages/R2WinBUGS/index.html]

In recent years, another kind of bayesian inference has gained traction is social, behavioral, and life sciences. This approach involves a focus on Bayes Factors as an index of support for or against hypotheses. It is not the purpose of this document to provide a full background on the Bayes Factor logic or method. Nonetheless, it is fairly simple to implement a Bayes Factor analysis of the rudimentary multiple regression methods covered in this document.

15.1 Basic approaches with Bayes Factors

The **BayesFactor** package, developed by Richard Morey (Morey and Rouder, 2018), is well documented and extensive tutorials are available.

For example:

[https://richarddmorey.github.io/BayesFactor/]

This section provides a brief overview of rudimentary analyses that create Bayes Factor evaluations of the linear model. I will use the two-IV model that we settled on as a reasonable one, the one with degrees_yr and cits as IVs and salary as DV. First, the ordinary least squares model is repeated so that we can compare results.

olsfit <- lm(salary~degree_yrs+cits,data=cohen1)
kable(tidy(olsfit))</pre>

term	estimate	std.error	statistic	p.value
(Intercept)	39073.6747	2396.47360	16.304655	0.0000000
degree_yrs	1061.7642	227.17563	4.673759	0.0000176
cits	212.1116	56.59392	3.747958	0.0004078

The regressionBF function permits simultaneous comparison of several models. Evaluation of only one model can use the 1mBF function. The goal is the calculation of Bayes Factor indices permitting comparison of models and yeilding a metric that is interpreted as relative evidence for one hypothesis over another. Initially here, the regressionBF function is used to assess three different models against an "intercept-only" model. And then we will compare the three models against each other. Why three models? With two IVs there are three possible models of interest - one each with a single IV and a third with both IVs With the large sample size and the sizeable correlations of degree_yrs and cits with salary, it is not surprising that each model yields a Bayes Factor that is a very large value. The BF is interpreted as a ratio. For example, the degree_yrs - only model is found to be over 7000 times more likely than the intercept-only model. The two-IV model has extremely an extremely large BF. It is important to understand that there is substantial theory underlying the BF computation and assumptions have been made to create default shape and scaling of choices for the prior. The reader is expected to become skilled in these choices and the theory before use of these methods.

```
bf1 <- regressionBF(salary~degree_yrs+cits, data=cohen1)
bf1</pre>
```

```
## Bayes factor analysis
## ------
## [1] degree_yrs : 71125.54 ±0.01%
## [2] cits : 4126.233 ±0%
## [3] degree_yrs + cits : 4963640 ±0%
##
## Against denominator:
## Intercept only
## ---
## Bayes factor type: BFlinearModel, JZS
```

At this point we have seen that both the OLS and BF models offer strong support for the idea that the degree_yrs plus cits model is a strong one. But it might be informative to look at a model that has much weaker evidence.

We can reconsider the model that used gender as an IV. It was not a significant predictor in the OLS model, which is repeated here.

```
olsfit2 <- lm(salary~gender,data=cohen1)
kable(tidy(olsfit2))</pre>
```

term	estimate	std.error	statistic	p.value
(Intercept)	52650.000	1810.539	29.079734	0.0000000
gendermale	3949.324	2444.918	1.615319	0.1114884

In a BayesFactor analysis we find a very different sized BF than for the degree_yrs+cits model. A BF value of 1.0 would indicate equivalent support for the null hypothesis (β =0) and the alternative ($\beta \neq 0$). The BF value found here (.773) indicates equivocal support for each with only very slight evidence favoring the null. We can take the reciprocal of the BF to find the degree of relative evidence for the null. That value is 1/.773 or 1.29. So we would say that the null is 1.29 times more likely than the null

```
bf1b <- lmBF(salary~gender, data=cohen1)
bf1b</pre>
```

```
## Bayes factor analysis
## ------
## [1] gender : 0.7726611 ±0.01%
##
## Against denominator:
## Intercept only
## ---
## Bayes factor type: BFlinearModel, JZS
```

Returning to the original two-IV model, the **BayesFactor** package permits some interesting and useful approaches to model comparison. Above, we only compared each of the three possible models to an interceptonly model. Now, we will compare them to each other. The relative evidence for one particular model can be found by taking the ratio of the BF to those models. Here, I compare the "best" model (using max) to each of the others. First, see that comparing the best model (the two-IV model) to itself yields a ratio of 1, and this makes sense. Against the twomodel,degree_yrs-only has less than 2% the strength of evidence times the strength of evidence. Or take the reciprocal and find that the two-IV model is nearly 70 times more likely Against the cits-only model the two-IV model fares even better with the cits-only model garnering less than .1% support.

```
bf2 <- bf1/max(bf1)
bf2
## Bayes factor analysis
## -----
## [1] degree_yrs
                                        ±0.01%
                         : 0.01432931
## [2] cits
                         : 0.0008312918 ±0%
## [3] degree_yrs + cits : 1
                                        ±0%
##
## Against denominator:
##
    salary ~ degree_yrs + cits
## ---
## Bayes factor type: BFlinearModel, JZS
```

It is possible to draw some useful plots of competing models. In this simplistic system, these plots are redundant, but with more IVs and more possible models, the visualizations can be a rapid way of comparing models.

plot(bf2) # the ratio models



plot(bf1) # the comparison to intercept-only model



This brief exposition is a superficial treatment of BF methods, but may give an indication of the relative ease with which they can be performed in R. There are well-done tutorials on the Morey's web page cited above, and a blog provides well written background theory:

[http://bayesfactor.blogspot.com/2014/02/the-bayesfactor-package-this-blog-is.html]

15.2 An admonition regarding Bayesian Inference

It is important to understand that there is substantial theory underlying the BF computation and assumptions have been made to create default shape and scaling of choices for the prior. The reader is expected to become skilled in these choices and the theory before use of these methods. A voluminous literature is available for the logic, theory, and implementation of Bayesian methods. A substantial amount of that literature is available via the stattoolkit bibliography shared with the readers.

16 Documentation for Reproducibility

R software products such as this markdown document should be simple to reproduce, if the code file or code are available. But it is also important to document the exact versions of the R installation, the OS, and the R packages in place when the document is created.

sessionInfo()

```
## R version 4.0.2 (2020-06-22)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 10 x64 (build 18362)
##
## Matrix products: default
```

locale: ## [1] LC COLLATE=English United States.1252 ## [2] LC_CTYPE=English_United States.1252 ## [3] LC_MONETARY=English_United States.1252 ## [4] LC NUMERIC=C ## [5] LC_TIME=English_United States.1252 ## ## attached base packages: ## [1] grid stats graphics grDevices utils datasets methods ## [8] base ## ## other attached packages: UsingR_2.0-6 Hmisc_4.4-0 ## [1] yhat_2.0-2 ## [4] Formula_1.2-3 HistData_0.8-6 tseries_0.10-47 ## [7] sandwich_2.5-1 rmarkdown_2.3 rcompanion_2.3.25 ## [10] plyr_1.8.6 plot3Drgl_1.0.1 rgl_0.100.54 ## [13] plot3D_1.3 psych 2.0.7 olsrr 0.5.3 ## [16] nortest_1.0-4 moments_0.14 lmtest_0.9-37 ## [19] zoo 1.8-8 knitr 1.29 HH 3.1-40 multcomp_1.4-13 ## [22] gridExtra_2.3 TH.data_1.0-10 ## [25] MASS_7.3-51.6 survival 3.2-3 mvtnorm 1.1-1 ## [28] latticeExtra_0.6-29 lattice 0.20-41 gt_0.2.1 ## [31] gvlma_1.0.0.3 ggthemes 4.2.0 ggExtra_0.9 ## [34] GGally_2.0.0 ggplot2_3.3.2 car_3.0-8 ## [37] carData_3.0-4 broom_0.7.0.9000 boot 1.3-25 BayesFactor_0.9.12-4.2 Matrix_1.2-18 **##** [40] bcdstats_0.0.0.9002 ## [43] coda_0.19-3 ## ## loaded via a namespace (and not attached): ## [1] readxl_1.3.1 backports_1.1.8 miscTools_0.6-26 ## [4] splines_4.0.2 gmp_0.6-0 crosstalk_1.1.0.1 ## [7] digest_0.6.25 htmltools_0.5.0 yacca_1.1.1 ## [10] magrittr_1.5 checkmate_2.0.0 cluster_2.1.0 ## [13] openxlsx_4.1.5 matrixStats_0.56.0 xts 0.12-0 ## [16] jpeg_0.1-8.1 colorspace_1.4-1 haven_2.3.1 ## [19] xfun 0.16 dplyr 1.0.0 libcoin 1.0-5 ## [22] crayon_1.3.4 Exact_2.0 jsonlite_1.7.0 ## [25] glue_1.4.1 gtable_0.3.0 webshot_0.5.2 ## [28] MatrixModels_0.4-1 Rmpfr_0.8-1 quantmod_0.4.17 ## [31] abind 1.4-5 SparseM 1.78 scales 1.1.1 [34] miniUI 0.1.1.1 Rcpp_1.0.5 isoband 0.2.2 ## ## [37] plotrix_3.7-8 xtable_1.8-4 htmlTable_2.0.1 ## stats4_4.0.2 [40] tmvnsim_1.0-2 foreign_0.8-80 ## [43] vcd_1.4-7 htmlwidgets_1.5.1 RColorBrewer_1.1-2 ## [46] modeltools_0.2-23 acepack_1.4.1 ellipsis_0.3.1 ## [49] farver_2.0.3 pkgconfig_2.0.3 reshape_0.8.8 ## [52] multcompView_0.1-8 nnet_7.3-14 labeling_0.3 ## [55] tidyselect_1.1.0 rlang_0.4.7 manipulateWidget_0.10.1 ## [58] reshape2_1.4.4 later_1.1.0.1 munsell_0.5.0 ## [61] cellranger_1.1.0 tools_4.0.2 generics_0.0.2 ## [64] EMT_1.1 evaluate_0.14 shinyBS_0.61 ## [67] stringr_1.4.0 fastmap_1.0.1 yam1_2.2.1 ## [70] goftest_1.2-2 purrr_0.3.4 zip_2.0.4

##	[73]	coin_1.3-1	pbapply_1.4-2	nlme_3.1-148
##	[76]	mime_0.9	quantreg_5.61	leaps_3.1
##	[79]	compiler_4.0.2	rstudioapi_0.11	curl_4.3
##	[82]	png_0.1-7	e1071_1.7-3	tibble_3.0.3
##	[85]	DescTools_0.99.37	stringi_1.4.6	highr_0.8
##	[88]	forcats_0.5.0	vctrs_0.3.2	pillar_1.4.6
##	[91]	lifecycle_0.2.0	data.table_1.13.0	conquer_1.0.1
##	[94]	httpuv_1.5.4	R6_2.4.1	bookdown_0.20
##	[97]	promises_1.1.1	KernSmooth_2.23-17	rio_0.5.16
##	[100]	codetools_0.2-16	gtools_3.8.2	withr_2.2.0
##	[103]	mnormt_2.0.1	mgcv_1.8-31	expm_0.999-5
##	[106]	parallel_4.0.2	hms_0.5.3	quadprog_1.5-8
##	[109]	rpart_4.1-15	tidyr_1.1.0	class_7.3-17
##	[112]	misc3d_0.8-4	Cairo_1.5-12.2	TTR_0.23-6
##	[115]	shiny_1.5.0	base64enc_0.1-3	

16.1 Revision History

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Converted the document to bookdown

Added many sections and updated many code chunks

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