

R Guide

version 0.1

Last generated: October 07, 2021



This work is licensed under a Creative Commons Attribution-ShareAlike 4.0 International License .

Table of Contents

Introduction

Overview
Using R
Getting Help
Printing and Setting the Working Directory5
Using Unix Commands from R6
Loading a Previous Session
Using External Data, Files, and Scripts
Directing Output to a File
Installing, Using, and Removing Packages11
Sample Data
Exiting R

Basic Math

Math in R

Variables and Matrix Algebra

Vectors	
Matrices	
Factors	

Central Tendency and Other Statistics

Mean	
Median	
Mode	
Standard Deviation	
Inner-Quartile Range	
Stem Plot	

Plots

Barplots	
Scatter Plots	
Histograms	
Boxplots	

Multiple Graphs	37

Hypothesis Testing

Hypothesis Testing Overview	. 39
Fisher's Exact Test	. 40
Chi-Square Test	. 41
Logistic Regression	. 42
Conditional Logistic Regression	. 44
Testing for Gene-Environment Interaction	. 46

PLINK Toolset

PLINK Data Files	. 51
PLINK Run Information	. 52

Introduction

This is a list of commands and their contexts for getting started with using R. R is a free programming language for statistical computing and graphics. To download R, visit cran and find the location closest to you. Then, follow the instructions based on your operating system. Many people who use R like the program RStudio, which has a console and interface for using R. That can be downloaded for free under an open source license from the RStudio website . A number of cheatsheets for common R packages have been created as a handy reference.

Using R

R can be loaded from the command line with R. Easy enough. If you remember, in Unix there is a \$ to tell you you're at the command line. For R's internal command line, that symbol is > . To clear the console in R or RStudio on a Windows or Linux operating system, use Cntrl+L. On a Mac, you can use either Cntrl+L or Command+Option+L.

PDF Download

Getting Help

R has some internal help commands. These include help(functionname) and example(functionname). An example is below.

Printing and Setting the Working Directory

The folder that you work through the terminal is known as the working directory. To know what your current working directory is, use:

> getwd()

which will return the path to where you are. To specify a working directory, you need to put the path to that directory in, which is shown in the example below.

> setwd("/Users/Path/Through/The/Computer")

Similarly, if a directory called Land within Computer needed to be accessed, then use:

> setwd("Land")

to access it.

Using Unix Commands from R

Another important thing to mention is that anything that needs to be run from the terminal can be done through the R command prompt using <code>system("")</code>. It will also print the information in the typical Unix format.

> system("pwd")
/Users/Path/Through/The/Computer

Loading a Previous Session

Previous R sessions are saved with the .RData extension.

> load("~/Path/Through/The/Computer/Example_old_session.RData")

Using External Data, Files, and Scripts

Files can be listed by using "list.files()", which will return possible files. The source command will run a script, and is shown below.

```
> source("bottle1.R")
[1] "This be a message in a bottle1.R!"
```

R can read a .csv file, which literally means comma separated values. The example below shows a csv, titled targets.csv, being read.

<pre>read.csv("schools.csv")</pre>		
University	Students	Tuition
1 Truman State University	6200	13500
2 University of Iowa	33300	29000
3 University of Michigan	44700	45000
4 University of North Texas	37900	20000

Text files (.txt) can also be read, but if the separator is a tab, the "read.table" command is a better fit. The following example shows the tab separator specification (sep="\t").

```
read.table("population.txt", sep="\t")
        V1 V2
1 City Population
2 Joliet 148262
3 Kirksville 17519
4 Denton 133808
5 New Lenox 26217
```

This example had V1 and V2 as headers. R isn't automatically aware that they are headers, so that needs to be specified when reading the table.

<pre>read.table("pop</pre>	ulation.txt",	sep="∖t",	header=TRUE)
City	Populati	on	
1 Joliet	148262		
2 Kirksville	17519		
3 Denton	133808		
4 New Lenox	26217		

Directing Output to a File

The **sink()** command is used direct output from an R job to a new file. Having nothing in the parentheses will return the output to the terminal. Using something like

```
> sink("myfile.txt", append=TRUE, split=TRUE)
```

will direct the output to a file titled **myfile.txt**, the append option will allow the output to be added to the existing file as opposed to overwriting it, and the split option will send the data to the screen and the output file.

Installing, Using, and Removing Packages

Packages are basically code bundles that enable certain functions and data sets to be used. If you know the name of the package that you wish to install, (in this example that package is car), then use:

> install.packages("car")

Once the package is installed, it can be referenced by using **library** to call it.

> library(car)

The active packages can be checked by using:

> (.packages())

Removing packages is known as "detaching" them. The following example removes the car package.

> detach("package:car")

Sample Data

R has several preloaded data packages that can be very helpful in learning R. The available packages can be loaded by using the command data(), which will then bring up a menu with options. To exit that menu, type **q**. To list what data is currently loaded, use the ls() command with empty parentheses. The following example loads the Orange data set (regarding the growth of orange trees).

> data(0range)

The commands head(), tail(), and rm() can be used similarly to how they would be in a Unix environment. To show the first line of the Orange dataset, the command would be head(Orange, n=1). If you wanted to unload the Orange dataset, use rm(Orange).

Exiting R

The R console can be exited by using **q()**.

```
> q()
Save workspace image? [y/n/c]:
```

The quit command prompts the ability to save your R session. Respond to the prompt, where y=yes, n=no, and c=cancel.

Math in R

You can use several different commands that are similar to how calculator programming works. The sum function would take the sum of all entered after it.

```
> sum(1, 3, 5)
[1] 9
```

The repeat function is fairly intuitive.

```
> rep("Yo ho!", times = 3)
[1] "Yo ho!" "Yo ho!" "Yo ho!"
```

As is acquiring the square root.

```
> sqrt(16)
[1] 4
```

Vectors

Vectors are a basic data structure that can include logical, integer, double, character, complex, or raw data. It can only hold one type of data at a time. A new vector can be created with the values 4, 7, and 9, where c stands for combine.

> c(4, 7, 9)
[1] 4 7 9

If the following command was used, all of the values would be coded as character values.

> c(1, TRUE, "three")
[1] "1" "TRUE"

"three"

Sequences and Variables

Sequences of numbers can also be saved as a vector, using a colon (start:end) or [seq(start, end)]. Using seq allows increments other than 1 to be used, when the increment is specified after end [seq(start, end, increment)].

> a <- 5:9							
[1]	5	6	7	8	9		
> alpha <-	seq(5, 9)					
[1]	5	6	7	8	9		
> quartersvector <- seq(5, 6, 0.25)							
[1]	5.00	5.25		5.50	5.75	6.00	
> variable	charlie <	- 9:5					
[1]	9	8	7	6	5		

The above example demonstrated saving sequences of numbers as variables. You can access the individual value in the vector through using the variable name and brackets. The following example will show that and include comments (shown by \#) about what's happening.

```
> variablecharlie[3] #I'm calling value 3 stored in the vector
[1] 7 #The third value is the number 7
> quartersvector[2]
[1] 5.25
> alpha[2] <-68 #replaces the second position of the vector wit
h 68
> print(alpha) #prints the values in alpha to command line
[1] 5 68 7 8 9
```

Names can be assigned to the variable with **names()**. After assignment, printing will show the names. Either single or double quotes can be used to assign names.

```
> limbs <- c(4, 3, 4, 3, 2, 4, 4, 4)</pre>
> names(limbs) <- c('One-Eye', 'Peg-Leg', 'Smitty', 'Hook', 'Sc</pre>
ooter', 'Dan', 'Mikey', 'Blackbeard')
> print(limbs)
0ne-Eye
                                Smitty
                                               Hook
                                                             Scoote
                Peg-Leg
                                    Blackbeard
r
         Dan
                     Mikey
          3
                             3
                                       2
4
                   4
                                                           4
                                                 4
4
```

To remove a variable (or dataset), use the rm() command, where the data you wish to remove is specified in the parentheses. Similarly, to list what data is available, use the ls() command with empty parentheses.

Saving R Data

Data can be saved with the save() command. Multiple data sets can be saved in one data file (in the example, these are x, y, and z). "R

```
save(x, y, z, file="xyz.rda")
```

Matrices

Filling Matrices

To make a 4x5 matrix filled with zeroes, it would look like:

> matrix	(0, 4, 5)					
	[,1]	[,2]	[,3]	[,4]	[,5]
[1,]	0	0	0	0	0	
[2,]	0	0	0	0	0	
[3,]	0	0	0	0	0	
[4,]	0	0	0	0	0	

If you wanted to have a matrix with values specified, store those values as a vector (page 15) first. The following example specifies a vector, **a**, as a sequence from 1 to 20, and then stores it as a matrix.

> a <- 1:20 > matrix(a, 4, 5)						
	[,1]	[,2]	[,3]		[,4]	[,5]
[1,]	1	5	9	13	17	
[2,]	2	6	10	14	18	
[3,]	3	7	11	15	19	
[4,]	4	8	12	16	20	

Determinants and Eigen-stuff

The determinant can be taken on a matrix. If the variable has been defined, but not yet placed in a matrix, both steps must be done at once.

```
> a<-1:25
> det(matrix(a,5,5))
[1] 0
```

The **eigen** function (I know, I'm laughing too) computes both eigenvalues and eigenvectors. If the matrix is symmetric, then include **symmetric=TRUE** to skip the symmetry check.

```
> eigen(matrix(a,5,5))
eigen() decomposition
$values
[1] 6.864208e+01+0.000000e+00i -3.642081e+00+0.000000e+00i
[3] 4.257350e-15+0.000000e+00i -1.270981e-16+4.588876e-16i
[5] -1.270981e-16-4.588876e-16i
$vectors
             [,1]
                            [,2]
[,3]
                      [,4]
[1,] 0.3800509+0i -0.76703416+0i 0.54621260+0i 0.1175132+0.04
59634i
[2,] 0.4124552+0i -0.48590617+0i -0.27228461+0i 0.4017692+0.00
65072i
[3,] 0.4448594+0i -0.20477817+0i -0.66418830+0i -0.7435988+0.00
00000i
[4,] 0.4772637+0i 0.07634982+0i -0.03961996+0i -0.1881630-0.20
33750i
[5,] 0.5096680+0i 0.35747782+0i 0.42988027+0i 0.4124793+0.15
09044i
                      [,5]
[1,] 0.1175132-0.0459634i
[2,] 0.4017692-0.0065072i
[3,] -0.7435988+0.0000000i
[4,] -0.1881630+0.2033750i
[5,] 0.4124793-0.1509044i
```

Matrix Multiplication

For element-wise multiplication, simply use the A*B format. To multiply matrices, you would use the ** symbol. The outer product (AB') can be obtained using the *0 symbol.

> > <	1.20	•									
- a <-	- 1.20	,									
> b <-	- 21:4	10									
> mat	rix(a,	4,	5)								
	[,1]		[,2]		[,	3]		[,4]		[,5]
[1,]		1		5		9		13		17	
[2,]		2		6		10		14		18	
[3,]		3		7		11		15		19	
[4,]		4		8		12		16		20	
> mat	rix(b,	5,	4)								
	[,1] [[,2]	[,3]	[,4]							
[1,]	21	26	31	36							
[2,]	22	27	32	37							
[3,]	23	28	33	38							
[4,]	24	29	34	39							
[5,]	25	30	35	40							
> a%*°	ы́р										
[,1]											
[1,]	7070										

Other Matrix Operations

Different formats are given, where A and B are matrices, and k is a scalar.

Table: Formats for Matrices

Command	Function
t(A)	transpose
<pre>crossprod(A,B)</pre>	A'B
crossprod(A)	A'A
solve(A)	Inverse of A (if A is a square matrix)
<pre>solve(A, b)</pre>	Returns x vector in b = Ax equation
diag(A)	Vector with elements of principal diagonal
diag(K)	Creates the k x k identity matrix
rowMeans(A)	Vector of row means

Command	Function
rowSums(A)	Vector of row sums
columnMeans(A)	Vector of column means
columnSums(A)	Vector of column sums
rowbind(A,B,)	Combines matrices or vectors vertically, returning a ma- trix
<pre>cbind(A,B,)</pre>	Combines matrices or vectors horizontally, returning a matrix

Writing Information to a File

Data, usually matrix data, are written to a file using the write command. In it, the original data vector is specified, then the file name, and then the delimiter. Tabs are specified through <u>\t</u>, but other separators can be used if placed in the quotes.

```
> write(mydata, "mydata.txt", sep="\t")
```

Files can also be written using write.csv() and write.table().

Factors

Factors are a new vector comprised of integer values with a corresponding character value. They are categorical variables. A good way to describe this is when thinking of an Olympic roster. There's hundreds of athlete names, and under "medal type" there would be a factor with four levels: gold, silver, bronze, and none. R can assign numbers to these factors when doing analysis (so, in the case of alleles, AA=3, Aa=2, aa=1 could be an assignment). To store a vector as a variable of a factor, use variable <- factor(vectorname). Their integer assignments can be checked with as.integer(variable), and the levels can be checked with levels(variable).

```
> allele <- c( 'aa', 'Aa', 'AA', 'aa', 'aa', 'aa', 'aa', 'Aa',
'Aa')
> types <- factor(allele)
> print(types)
[1] aa Aa AA aa aa aa aa Aa Aa
Levels: aa Aa AA
> as.integer(types)
[1] 1 2 3 1 1 1 1 2 2
> levels(types)
[1] "aa" "Aa" "AA"
```

Mean

Taking the average of a variable is as easy as using mean(variable).

```
> spidersconsumed <- c(0,0,0,0,0,0,0,0,0,30)
> mean(spidersconsumed)
[1] 3
```

The average person eats 3 spiders per year. The child of Spiders Georg, Spiders Meorg, ate 30 spiders, and kept the average in his class of 10 students. Clearly, this lead to a statistical error, and the data would be better represented another way.

Median

Similarly to mean, simply running median(variable) provides the median.

```
> median(spidersconsumed)
[1] 0
```

Thank goodness!

Mode

Unlike the other areas of central tendency, R doesn't have a straightforward way to calculate the mode. Never fear: the following has an example for both numeric and character vectors.

```
# Create the function.
getmode <- function(spidersconsumed) {</pre>
   uniq <- unique(spidersconsumed)</pre>
   uniq[which.max(tabulate(match(spidersconsumed, uniq)))]
}
# Create the vector with numbers.
spidersconsumed <- c(0,0,0,0,0,0,0,0,0,0,0,0)</pre>
# Calculate the mode using the user function.
result <- getmode(spidersconsumed)</pre>
print(result)
[1] "0"
# Create the vector with characters.
students <- c("Susie", "Jessica", "Sally", "Henry", "George",</pre>
"Alex", "Alex", "Olive", "Patrick", "Meorg")
# Calculate the mode using the user function.
result2 <- getmode(students)</pre>
print(result2)
[1] "Alex"
```

Standard Deviation

To get the standard deviation, use sd(variable).

```
> sd(spidersconsumed)
[1] 9.486833
```

This is almost worse, because it means people eat 3 ± 9.48 spiders per year!

Inner-Quartile Range

The command **summary(variable)** can be used to get the minimum, 1st IQR, median, mean, 3rd IQR, and maximum values.

>summary(spidersconsumed)							
	Min.	19	st Qu.	Median		Mean	3r
d Qu.		Max.					
	0	0	0	3	0	30	
	0	0	0	3	0	30	

Stem Plot

One way to visualize your data is with a stem and leaf plot, which gives a visual representation of where your data lie. The command to create a stem plot is stem(variable).

```
> stem(spidersconsumed)
The decimal point is 1 digit(s) to the right of the |
0 | 00000000
1 |
2 |
3 | 0
```

Wow, Spiders Meorg really does look like an outlier now!

Barplots

The following example shows what steps were taken to create a vector named **population** and then have a barplot with those values.

The command names assigned names to the values in the vector population. The working directory was set to specify where the file save location for a png file of the generated image populationbarplot.png should be saved (in this case, the Desktop). (Note: you can also select pdf() and jpg().) Then, the barplot was made with barplot. Finally, because you're no longer using the screen for image generation, you turn off the graphics device with dev.off(). The generated barplot is shown in the image below.



Example use of **barplot**, which creates a barplot.

To add a line at the mean (or median, etc.) of the population, the **abline** command can be used, which will update the current plot.

```
> abline(h = mean(population))
```

To show one standard deviation above the mean, use an argument of h = mean(population) + sd(population). The only way to remove a line by mistake is to regenerate the barplot and effectively start over. Multiple lines need to be added with individual abline commands.

Scatter Plots

This example will use the preloaded dataset Theoph, which has data from an experiment on the pharmacokinetics of the respiratory drug thephylline. By using head on this dataset, we can see what the columns are coded as.

> head(Theoph, $n = 1$)							
	Subject	Wt	Dose	Ti	me	conc	
1	1	79.6	4.02	0	0.74		

Knowing how the columns are coded allows you to reference certain columns in the data set using dataset\$columnname. With the plot command, you can designate where the data are (in the following example, the x-axis is Theoph\$Time and the y-axis is Theoph\$conc. Using xlab and ylab sets the axes labels, and pch chooses the marker type (seen here (page 32)).



Pharmacokinetics of Theophylline

A scatterplot created using the Theoph dataset.

Figure: Marker types

0 =		13 =	Ø
1 =	0	14 =	
2 =	\bigtriangleup	15 =	•
3 =	+	16 =	٠
4 =	×	17 =	۸
5 =	\diamond	18 =	٠
6 =	\bigtriangledown	19 =	٠
7 =		20 =	•
8 =	*	21 =	0
9 =	\Leftrightarrow	22 =	
10 =	\oplus	23 =	\diamond
11 =	XX	24 =	\bigtriangleup
12 =	⊞	25 =	\bigtriangledown

Different marker options available, with the default equal to pch=1.

Histograms

A histogram provides a visual depiction of a dataset's distribution. This example uses the dataset USArrests, which contains information on the violent crime rates by US state.



Histogram for US Murders

A histogram of the Murder column from the USArrests dataset.

Because there were multiple columns of information, the Data\$Column option was used to specify which column the histogram should be created from. The title was set using main=, colors for the border (border="gray") and columns (col="red") were set, and the x-axis boundries were chosen as 0-20 (xlim=c(0,20)). None of the options were necessary, except the initial Data\\$Column specification. There are also more options for histograms found in the documentation, including adding breaks (which can organize how data is grouped) or density curves.

Boxplots

Boxplots are one way of graphically orienting information from the summary(variable) (page 26) command.



A boxplot depicting the Murder, Assault, and Rape columns from the USArrests dataset.

Because specific columns of data were selected, their individual names had to be specified to create the image above. Otherwise, the default names of 1, 2, and 3, would have been used.

Multiple Graphs

R lets you graph multiple data sets in one composed figure. This is done through the par command. Using par, you can specify a matrix (in the examples case, 2x2), where images are added from top left to top right, bottom left to bottom right. Each one can be formatted as previously described. The example used three different sets of pre-loaded data, and the very first graph is the only one without a specified title.

```
> setwd("~/Desktop")
> png(filename="random_data_R.png")
> par(mfrow=c(2,2))
> plot(Theoph$Time, Theoph$conc, xlab="Time (min)", ylab="Conce
ntration (M)", pch=16)
> plot(Theoph$Wt, Theoph$Dose, xlab="Weight (kg)", ylab="Dose
(mM/L)", pch=2, main="Theoph Dosing")
> plot(Formaldehyde, xlab="Carb", ylab="optden", pch=16, mai
n="Formaldehyde Data")
> hist(AirPassengers)
> dev.off()
quartz
2
```



A random dataset, created using par.

Hypothesis Testing Overview

What would a statistics software be without hypothesis testing? Answer: Not a statistics software. Anyway, there are several options for hypothesis testing, which are all better explained in a stats class. So let's get on with the examples. First, we'll create our fake data set (which would be something we'd eventually want to test for Hardy-Weinberg equilibrium).

```
>Input =("
Genotype Observed Expected
AA
               200
                             160
               400
                             360
Aa
               400
                             480
aa
••)
>Matrix = as.matrix(read.table(textConnection(Input),
                    header=TRUE,
                    row.names=1))
> Matrix
   Observed Expected
AA
        200
                  160
        400
                  360
Aa
        400
                  480
aa
```

Fisher's Exact Test

Using our fake data, we'll run Fisher's exact test. This test is typically used in place of the chi-square on small samples, but it is valid for all sample sizes.

Hypotheses

- $H_0 = in HWE$
- $H_a = not in HWE$

```
>fisher.test(Matrix)
        Fisher's Exact Test for Count Data
data: Matrix
p-value = 0.0009992
alternative hypothesis: two.sided
```

Since the p-value is less than α (0.05 at the 95% level), we reject the (H_0) null hypothesis in favor of the alternative (H_a). There is enough evidence to say that the fake data is not in HWE.

Chi-Square Test

Using our fake data, we'll run a chi-square test. This test is used to evaluate how likely there is to be an observed difference between sets.

Hypotheses

- $H_0 = in HWE$
- $H_a = not in HWE$

```
> chisq.test(Matrix)
        Pearson's Chi-squared test
data: Matrix
X-squared = 13.822, df = 2, p-value = 0.0009965
```

Since the p-value is less than α (0.05 at the 95% level), we reject the (H_0) null hypothesis in favor of the alternative (H_a). There is enough evidence to say that the fake data is not in HWE.

Logistic Regression

Logistic regression is a statistical model where the independent variable is continuous and the dependent variable is binary. Some binary variables of interest would be health/sick, pass/fail, win/lose, or alive/dead.

R has a function to use the general linearized model to perform logistic regression. You're not going to need that fake data set here, but you will need a data set with several rows of individual data, specified by columns. Perhaps something like this:

StudentID	Passing	Age	Tutor	TotalCredits	Gender	<pre>matched_sets</pre>
1234	1	32	1	178	0	100
1235	0	12	0	12	1	300
1236	1	25	0	0	2	200
1237	1	24	1	123	0	200
1238	1		34	0 293	(0 100
1239	0	15	1	25	1	300

In the future example, **Passing** will be referred to as **STATUS** (where 1 = yes, 0 = no), and the columns to the right of **Passing** refer to the dummy variables A through D. Similarly, the data is stratified into the **matched_sets** strata. There can be more columns in the data, and the columns can be in any order; any formulas will reference columns by their header.

In the following example, the result of the test for the general linearized model is saved under logistical. The syntax for the glm command is thoroughly explained through using help(glm). A quick summary is that it tests a formula of specified data columns against a specific distribution model, such as binomial or Gaussian (family =).

```
> data = read.table("/path/to/text/file/with/data", header=TRU
E, na.strings = "NA")
> logistical <- glm(data$STATUS ~ data$VariableA + data$Variabl</pre>
eB + data$VariableC + data$VariableD, family = binomial)
> summary(logistical)
Call:
glm(formula = data$STATUS ~ data$VariableA + data$VariableB + d
ata$VariableC +
    data$VariableD, family = binomial)
Deviance Residuals:
    Min
              10
                  Median
                                30
                                       Max
-1.6684 -0.8207 -0.5998
                           1.1268
                                    2.0336
Coefficients:
                 Estimate Std. Error z value Pr(>|z|)
(Intercept)
              -2.5341507 0.5566591 -4.552 5.30e-06 ***
data$VariableA 0.0041581 0.0008622
                                      4.823 1.42e-06 ***
data$VariableB -0.0190579 0.0252128 -0.756 0.44972
data$VariableC 0.3291734 0.2707949 1.216 0.22414
data$VariableD 0.8224185 0.2691547 3.056 0.00225 **
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 366.95 on 293 degrees of freedom
Residual deviance: 330.86 on 289 degrees of freedom
  (1 observation deleted due to missingness)
AIC: 340.86
Number of Fisher Scoring iterations: 3
```

The level of significance for the *p*-value is given by the number of asterisks. Three asterisks means that the *p*-value for that result is below 0.001, but larger than 0. Significant results allow the null hypothesis to be rejected, and the significance code specifies whether this is done at the 90% (.), 95% (*), 99% (**), or 99.9% (***) level.

Conditional Logistic Regression

Conditional logistic regression is like logistic regression, but it can take stratification and matching into account.

One form of conditional logistic regression can be performed by loading the Epi package. An additional form can be performed by loading the survival package.

This example uses the Epi package. The individual components of the clogistic command syntax have been shown through the equals signs. The data\$ is not necessary because the data variable is specified in the equation (which was an option for glm). Conditional logistic regression requires the use of stratified data, where the column of the strata is specified (here as matched_sets). Unlike before, the definition must be entered after defining the test; summary will not work.

```
> library(Epi)
> data = read.table("/path/to/text/file/with/data", header=TRU
E, na.strings = "NA")
> EPI.clogistic <- clogistic(formula = STATUS ~ VariableA + Var
iableB + VariableC + VariableD, strata = matched_sets, data = d
ata)
> EPI.clogistic
Call:
cloqistic(formula = STATUS ~ VariableA + VariableB + VariableC
+ VariableD,
    strata = matched_sets, data = data)
                  coef exp(coef) se(coef)
                                               Ζ
                                                       р
VariableA
             -0.01002
                           0.99 0.03261 -0.307 7.6e-01
VariableB
              0.50098
                           1.65 0.36442 1.375 1.7e-01
VariableC
                           1.17 0.27884 0.554 5.8e-01
              0.15458
VariableD
              0.00486
                           1.00 0.00112 4.343 1.4e-05
Likelihood ratio test=26.5 on 4 df, p=2.51e-05, n=292
```

This example uses the **survival** package. It has two distinct differences from Epi's **clogit**. Firstly, **strata** are included into the formula, as opposed to being a second parameter. Second, the summary function can be used.

```
> library(survival)
> data = read.table("/path/to/text/file/with/data", header=TRU
E, na.strings = "NA")
> survival.clogit <- clogit(formula = STATUS ~ VariableA + Vari</pre>
ableB + VariableC + VariableD + strata(matched_sets) data = dat
a)
> summary(survival.clogit)
Call:
coxph(formula = Surv(rep(1, 295L), STATUS) ~ VariableA + Variab
leB + VariableC +
   VariableD + strata(matched_sets), data = data, method = "exa
ct")
  n= 294, number of events= 93
   (1 observation deleted due to missingness)
                   coef exp(coef)
                                   se(coef)
                                                 z Pr(>|z|)
VariableA
              -0.010025 0.990025
                                   0.032612 -0.307
                                                      0.759
VariableB
               0.500983 1.650342 0.364415 1.375
                                                      0.169
               0.154577 1.167164 0.278843 0.554
VariableC
                                                      0.579
VariableD
               0.004864 1.004876 0.001120 4.343 1.41e-05 ***
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
          exp(coef) exp(-coef) lower .95 upper .95
VariableA
              0.990
                        1.0101
                                  0.9287
                                             1.055
VariableB
                        0.6059
                                  0.8079
                                             3.371
              1.650
VariableC
                        0.8568
                                  0.6757
                                             2.016
             1.167
VariableD
              1.005
                        0.9951
                                  1.0027
                                             1.007
                 (max possible= 0.494 )
Rsquare= 0.086
Likelihood ratio test= 26.5
                              on 4 df,
                                         p=2.511e-05
Wald test
                     = 21.21
                              on 4 df,
                                         p=0.0002873
Score (logrank) test = 24.77
                              on 4 df,
                                         p=5.607e-05
```

The level of significance for the *p*-value is given by the number of asterisks. Three asterisks means that the *p*-value for that result is below 0.001, but larger than 0. Significant results allow the null hypothesis to be rejected, and the significance code specifies whether this is done at the 90% (.), 95% (*), 99% (**), or 99.9% (***) level.

Testing for Gene-Environment Interaction

Gene-environment interaction is when different genotypes have different responses to variation in the environment. To test for this, the data set needs genotypic and phenotypic information to be combined, which can be done using R's cbind function. If you're the type of person that like combining data in Excel, however, then you can get started once your input file looks clean.

Logistic Regression GxE

The only difference when performing logistic regression for gene by environment with the original is the addition of the terms of interest multiplied by the environment variable that you're testing.

The example tests the environmental VariableD on variables A through C (sometimes shortened like VA).

```
> data = read.table("/path/to/text/file/with/data", header=TRU
E, na.strings = "NA")
> logisticalGE <- glm(data$STATUS ~ data$VariableA + data$Varia</p>
bleB + data$VariableC + data$VariableD + data$VariableA*data$Va
riableD + data$VariableB*data$VariableD + data$VariableC*data$V
ariableD, family = binomial)
> summary(logisticalGE)
Call:
qlm(formula = data$STATUS ~ data$VariableA + data$VariableB + d
ata$VariableC +
    data$VariableD + data$VariableA * data$VariableD + data$Var
iableB *
    data$VariableD + data$VariableC * data$VariableD, family =
binomial)
Deviance Residuals:
    Min
              10
                  Median
                               30
                                       Max
-1.7205 -0.8537 -0.5189 1.0470
                                    2.5061
Coefficients:
                           Estimate Std. Error z value P
r(>|z|)
(Intercept)
                         -4.5268619 1.0737534 -4.216 2.49e-0
5 ***
data$VariableA
                     0.1690201 0.0530564 3.186 0.001444 **
datasVariableB
                     0.5921538 0.5478683 1.081 0.279772
data$VariableC
                     0.5843189 0.5787536 1.010 0.312679
data$VariableD
                     0.0136132 0.0038047 3.578 0.000346 ***
data$VA:data$VD
                    -0.0007583 0.0001849 -4.101 4.12e-05 ***
                    -0.0012611 0.0018368 -0.687 0.492361
data$VB:data$VD
data$VC:data$VD
                    -0.0003966 0.0018996 -0.209 0.834636
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 366.95 on 293 degrees of freedom
Residual deviance: 311.15 on 286 degrees of freedom
  (1 observation deleted due to missingness)
AIC: 327.15
Number of Fisher Scoring iterations: 5
```

The level of significance for the *p*-value is given by the number of asterisks. Three asterisks means that the *p*-value for that result is below 0.001, but larger than 0. Significant results allow the null hypothesis to be rejected, and the significance code specifies whether this is done at the 90% (.), 95% (*), 99% (**), or 99.9% (***) level.

Conditional Logistic Regression GxE

Again, like logistic regression's gene by environment interaction, the only change for conditional logistic regression is the addition of the terms of interest multiplied by the environment variable being tested.

The example tests the environmental VariableD on variables A through C (sometimes shortened like VA).

```
> survival.clogitGE <- clogit(formula = STATUS ~ VariableA+ Var</pre>
iableB + VariableC + totalanth + VariableA*VariableD + Variable
B*VariableD + VariableC*VariableD + strata(matched_sets),data)
> summary(survival.clogitGE)
Call:
coxph(formula = Surv(rep(1, 295L), STATUS) ~ VariableA + Variab
leB + VariableC +
  VariableD + VariableA * VariableD + VariableB * VariableD +
VariableC *
   VariableD + strata(matched_sets), data = data, method = "ex
act")
 n= 294, number of events= 93
  (1 observation deleted due to missingness)
                     coef exp(coef) se(coef)
                                                     zΡ
r(>|z|)
VariableA
                1.776e-01 1.194e+00 5.938e-02 2.991 0.00278
1 **
VariableB
                2.608e-01 1.298e+00 7.080e-01 0.368 0.71259
8
VariableC
                1.764e-01 1.193e+00 5.686e-01 0.310 0.75643
3
VariableD
                1.230e-02 1.012e+00 4.091e-03 3.005 0.00265
2 **
VA:VD
               -7.187e-04 9.993e-01 1.988e-04 -3.616 0.00029
9 ***
VB:VD
               -6.979e-05 9.999e-01 2.076e-03 -0.034 0.97318
1
VC:VD
               -7.048e-04 9.993e-01 1.976e-03 -0.357 0.72126
4
Signif. codes: 0 `***' 0.001 `**' 0.01 `*' 0.05 `.' 0.1 ` ' 1
            exp(coef) exp(-coef) lower .95 upper .95
VariableA
              1.1943
                         0.8373
                                   1.0631
                                             1.3418
VariableB
              1.2980
                         0.7704
                                   0.3240
                                             5.1991
VariableC
              1.1929
                         0.8383
                                   0.3914
                                             3.6356
VariableD
              1.0124
                         0.9878
                                   1.0043
                                             1.0205
VA:VD
              0.9993
                         1.0007
                                   0.9989
                                             0.9997
VB:VD
              0.9999
                         1.0001
                                   0.9959
                                             1.0040
VC:VD
              0.9993
                         1.0007
                                   0.9954
                                             1.0032
Rsquare= 0.133 (max possible= 0.494)
Likelihood ratio test= 41.97 on 7 df,
                                       p=5e-07
```

Wald test		=	27.16	on	7	df,	p=3e-04
Score (logrank)	test	=	36.31	on	7	df,	p=6e-06

The level of significance for the *p*-value is given by the number of asterisks. Three asterisks means that the *p*-value for that result is below 0.001, but larger than 0. Significant results allow the null hypothesis to be rejected, and the significance code specifies whether this is done at the 90% (.), 95% (*), 99% (**), or 99.9% (***) level.

PLINK Data Files

PLINK is a free computational package that can be used for genome association analyses.

Data for PLINK should be in PED and MAP files. PED files are space or tab delimited files, and need to start with the following 6 columns: Family ID, Individual ID, Maternal ID, Paternal ID, Sex, and Phenotype. Sex should be coded with 1=male; 2=female; and any other value meaning unknown. MAP files describe markers, and should contain only four columns: chromosome (1-22, X, Y, or 0 if unplaced), rs\# or SNP ID, Genetic distance (in the morgan unit), and Base-pair position (bp units).

There is more information about these file types on the PLINK website .

PLINK Run Information

Some of the summary statistics that PLINK can generate (through different commands) include missing genotype rate (missingness), Hardy-Weinberg equilibrium, minor allele frequency, and linkage disequilibrium. There are many more things PLINK can do (including family-based association testing for disease traits), which are all further described on the PLINK website.

PLINK Job Script

The following script would run the PLINK commands found in the **plink_frq** file on a SLURM scheduler.

```
#!/bin/bash
#SBATCH —p public
#SBATCH ——qos general
```

```
# partition aka allocation
# quality-of-service (priority)
```

```
module load plink/1.07
```

./plink_frq

If you get an error code, try **source plink_frq** instead of **./plink_frq**. The error dependent on where you're sourcing the file from.

Tests

A sample PLINK information file, **plink_frq**, specifying what PLINK needs to run to test for minor allele frequency (MAF) is below.

Where the actual test is will be changed for each different test. If the path were ~/home/euid123/R_jobs/ and the .ped and .map were both named example, then the line would be ~/home/euid123/R_jobs/example \.

Table: Test specifics for PLINK

Specifier	Test or Function	Test Statistic
freq	Minor Allele Frequency	MAF
het	Heterozygosity	F Value
hardy	Hardy Weinberg Equilibrium	HWE
r2	Linkage Disequilibrium	r ²
out snps	Linkage Disequilibrium	r ²
missingmind 1	Missingness	F_MISS
recodeAD	Recode	NA

- **Frequency**: test for minor allele frequency.
- Heterozygosity: test for inbreeding coefficients.
- Hardy-Weinberg Equilibrium: test for Hardy-Weinberg equilibrium.
- Linkage Disequilibrium: test for linkage disequilibrium.
- Missingness: test for missingness.
- Recode: change data coding to additive and dominance components.

Using awk on Data

The awk Unix command (page 0) can be used to parse out specified data. The following command would create the plinkawk.frq file from the generated plink.frq file, preserving the header, for values in column 5 that are less than 0.05.

awk 'NR == 1; NR > 1 {if (\$5<0.05) print}' plink.frq > plinkaw
k.frq