

Introduction to R

We will use R extensively in this course. R is a high level language especially designed for statistical calculations. R is free. You can get it at:

<http://www.cran.r-project.org/>

There are versions for Unix, Linux, Windows and Mac. There is a similar program called Splus. The commands are virtually identical. Splus has more stuff in it but R is free and it is faster. If you want to use Splus, it is available on Andrew (just type Splus) or you can purchase a copy from Insightful at <http://www.splus.mathsoft.com/>.

On my website I will often post R programs and examples.

Getting started

In Unix or Linux, you start R by typing: R. In windows, click on the R icon. You can now use R interactively. Just start typing commands. You can also use R in Batch mode. To do this, store your R commands in a file, say, file.r. In R type: `source("file.r")` which will execute the commands in file.r. In Unix, you can also do the following:

```
R BATCH file.r file.out &
```

which will execute the commands and store them in file.out.

IMPORTANT! Use the command: `q()` to quit from R.

Some Basics of R

Here is a simple R session. You should try these commands out. Also, feel free to experiment a bit. Note: the `#` symbol means “comment.” R ignores any command after `#`. I have added lots of comments below to explain what is going on. You do not need to type the comments.

```
x <- 5          ### assign x the value 5
x              ### print x
print(x)       ### another way to print x
x_5            ### you can also use _ to make assignments but <- is better
x
y <- "Hello there"
y
y <- sqrt(10)
z <- x+y
z
q()            ### Use this to quit
```

Scalars are treated by S-plus as vectors of length 1. That is why they print with a leading “[1]” indicating that we are at the first element of a vector.

- Vectors can be created using the `c()` command. `c()` stands for concatenate. Square brackets are used to get subsets of a vector. The colon is used for sequences. Start up R again then do this:

```
x <- 1:5
print(x)
x[1] <- 17
print(x)
x[1] <- 1
x[3:5] <- 0
print(x)
w <- x[-3]
print(w)
y <- c(1,5,2,4,7)
y
y[2]
y[-3]
y[c(1,4,5)]
i <- (1:3)
z <- c(9,10,11)
y[i] <- z
print(y)

y <- y^2
print(y)
y <- 1:10
y <- log(y)
y
y <- exp(y)
y
x <- c(5,4,3,2,1,5,4,3,2,1)
z <- x + y
z                                     ### R carries out operations on vectors, element by element

x <- 1:10
y <- c(5,4,3,2,1,5,4,3,2,1)
x == 2                                ### this is a logical vector
z <- (x == 2)
print(z)
z <- (x<5); print(z)                  ### You can out two commands on a line if you use a semi-colon
x[x<5] <- y[x<5]                      ### do you see what this is doing?
print(x)
```

- Two expressions can be written on the same line if separated by a semicolon. One expression can be written over several lines *as long as* a valid expression does not end a line.

- To create a “matrix”, use the `matrix()` function as follows:

```
junk <- c(1,2,3,4,5, 0.5, 2, 6, 0, 1, 1, 0)
m <- matrix(junk,ncol=3)
print(m)
m <- matrix(junk,ncol=3,byrow=T)
print(m)                                ### see the difference?
dim(m)
y <- m[,1]          ### y is column 1 of m
y
x <- m[2,]          ### x is row 2 of m
x
z <- m[1,2]
print(z)
zz <- t(z)          ### take the transpose
zz
new <- matrix( 1:9, 3 , 3)
print(new)
hello <- z + new
print(hello)
```

The square brackets are used to refer to the rows and columns of a matrix, similar to the way they are used for vectors.

```
m[1,3]
subm <- m[2:3, 2:4]
m[1,]
m[2,3] <- 7
m[,c(2,3)]
m[-2,]
```

```
x <- runif(100,0,1)          ### generate 100 numbers randomly between 0 and 1
mean(x)
y <- mean(x)
print(y)
help(mean)
min(x)
```

```
max(x)
summary(x)
help(summary)
```

- Lists are used to combine data of various types.

```
who <- list(name="Joe", age=45, married=T)
print(who)
print(who$name)
print(who[[1]])
print(who$age)
print(who[[2]])
print(who$married)
print(who[[3]])
names(who)
who$name <- c("Joe","Steve","Mary")
who$age <- c(45,23)
who$married <- c(T,F,T)
who
```

- A for loop is a statement that is used to repeat commands.

```
for(i in 1:10){
  print(i+1)
}

x <- 101:200
y <- 1:100
z <- rep(0,100)          ### rep means repeat
help(rep)
for(i in 1:100){
  z[i] <- x[i] + y[i]
}

w <- x + y
print(w-z)
```

```
### As this example shows, we can often avoid using loops since
### R works directly with vectors.
### Loops can be slow so avoid them if possible.
```

```
for(i in 1:10){
  for(j in 1:5){
    print(i+j)
  }
}
```

- **Reading in commands** To read in commands or functions from a file rather than typing them in, use `source()`. Put some R commands into a file called `hello`. Try `source('hello')`

- **Functions** You can create your own functions in R. Here is an example.

```
my.fun <- function(x,y){
  ##### This function takes x and y as input.
  ##### It returns the mean of x minus the mean of y
  a <- mean(x)-mean(y)
  return(a)
}
```

```
x <- runif(50,0,1)
y <- runif(50,0,3)
output <- my.fun(x,y)
print(output)
```

You can return more than one thing in a function. If you do, you should return a list.

```
my.fun <- function(x,y){
  mx <- mean(x)
  my <- mean(y)
  d <- mx-my
  out <- list(meanx=mx,meany=my,difference=d)
  return(out)
}
```

```
x <- runif(50,0,1)
y <- runif(50,0,3)
output <- my.fun(x,y)
print(output)
names(output)
output$difference
output[[3]]
```

- Here are some more R examples

```

### if statements
for(i in 1:10){
  if( i == 4)print(i)
}
for(i in 1:10){
  if( i != 4)print(i)      ### != means  'not equal to'
}
for(i in 1:10){
  if( i < 4)print(i)
}
for(i in 1:10){
  if( i <= 4)print(i)
}
for(i in 1:10){
  if( i >= 4)print(i)
}

###Plots

x <- 1:10
y <- 1 + x + rnorm(10,0,1)  ### rnorm(10,0,1) means 10 random Normals,
                           ### mean 0, standard deviation 1

plot(x,y)
plot(x,y,type="h")
plot(x,y,type="l")
plot(x,y,type="l",lwd=3)
plot(x,y,type="l",lwd=3,col=6)
plot(x,y,type="l",lwd=3,col=6,xlab="x",ylab="y")

par(mfrow=c(3,2))          ### put 6 plots per page, in a 3 by 2 configuration
for(i in 1:6){
  plot(x,y+i,type="l",lwd=3,col=6,xlab="x",ylab="y")
}

postscript("plot.ps")      ### put the plots into a postscript file
                           ### you have to do this if you use BATCH
plot(x,y,type="l",lwd=3,col=6,xlab="x",ylab="y")
dev.off()                  ### Now you can print the file our view it with
                           ### a ghostview previewer

par(mfrow=c(1,1))          ### return to 1 plot per page

```

```

x <- rnorm(100,0,1)      ### 100 random normals, mean 0, st.dev . 1
y <- rpois(500,4)        ### 500 random Poisson(4)
hist(y)                  ### histogram
hist(y,nclass=50)

pnorm(2,0,1)             ### P(Z < 2) where Z ~ N(0,1)
pnorm(2,1,4)             ### P(Z < 2) where Z ~ N(1,4^2)
qnorm(.3,0,1)            ### find x such that P(Z < x)=.3 where Z ~ N(0,1)

x <- seq(-3,3,length=1000) ### make a sequence of numbers
f <- dnorm(x,0,1)        ### normal density
plot(x,f,type="l",lwd=3,col=4)

```