

Lab #3

Power and sample size

In this lab, we are working with a data set from Alizadeh et al. at Stanford. In this study, the investigators were evaluating diffuse large B-cell lymphoma (DLBCL). Using expression profiling and hierarchical clustering (a topic that we will visit later in this class), they were able to identify 2 distinct forms of DLBCL that indicate different stages of B-cell differentiation. “One type expressed genes characteristic of germinal centre B cells (‘germinal centre B-like DLBCL’); the second type expressed genes normally induced during in vitro activation of peripheral blood B cells (‘activated B-like DLBCL’).” They also found that the germinal centre B-like DLBCL patients had a better survival rate.

We will use this data set to evaluate the power and sample size in this experiment. We will also look for the necessary number of samples to appropriately power the study. First we will calculate the power and n required using a single gene calculation for illustration of the formula, then we will conduct a more multivariate summary that gives an idea of the power or n required for a specific percentage of genes/probes in the experiment. Remember that when we calculate these statistics for a microarray, we are dealing with more than a single variable, so general power formulas do not apply when attempting to summarize all genes/probes on an array.

The paper entitled “Distinct type of diffuse large B-cell lymphoma identified by gene expression profiling” can be found on the course website.

1.) Download the Eisen DLBCL data set and save as a text file (go to class web site or see syllabus for paper URL).

2.) Load into R, using read.table and arguments:

```
header=T
na.strings="NA"
blank.lines.skip=F
row.names=1
```

There are missing values in this data frame because we’re working with cDNA data.

```
data = read.table("c:\\temp\\eisen.txt", header=T,
                  na.strings="NA", blank.lines.skip=F,
                  row.names=1)
```

```
data[1:4, 1:4]
```

```
      DLCL.0001 DLCL.0002 DLCL.0003 DLCL.0004
1      1.042      0.419      1.285      0.883
2     -0.097     -0.072      0.563      0.151
3      0.339     -0.337     -0.372     -0.054
4      0.028      0.069      0.209     -0.225
```

3.) Get the class label file "eisenClasses.txt" from the class web site and read it into R. Use the header=T argument.

```
classes = read.table("c:\\temp\\eisenClasses.txt", header=T)
classes[1:4, 1:2]
```

```
  class sample
1     1 DLCL.0012
2     1 DLCL.0024
3     1 DLCL.0003
4     1 DLCL.0026
```

4.) Subset the data frame with the class labels and look at the positions so you know where one class ends and the other begins. Remember that 'subset' means to re-index (i.e. reorder) the column headers. If you look at the original column name order with `dimnames(dat)[[2]]` both before and after you reorder them, you will see what this has done.

```
dimnames(data)[[2]] #show column names
class1 <- as.character(classes[1:19,2])
data.class1 <- data[,class1]
dimnames(data.class1)[[2]] #show class1 column names
class2 <- as.character(classes[20:39,2])
data.class2 <- data[,class2]
dimnames(data.class2)[[2]] #show class2 column names
```

5.) Pick a gene, remove cells that have "NAs", and plot the values for both classes with a:

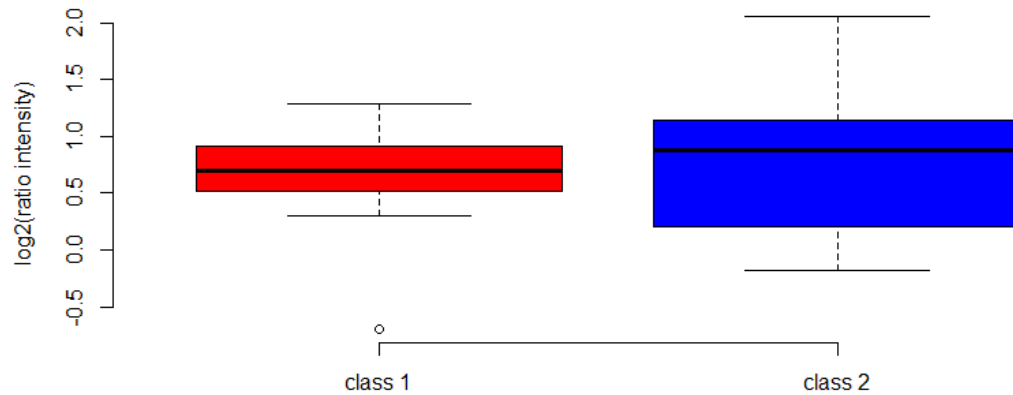
- boxplot (use the argument `col=c("red","blue")` to color separate boxes)
- histogram (this should have 2 separate histogram plots on 1 page; use the `par(mfrow=c(2,1))` function prior to plotting the first).

Color each class something different in the boxplot and histogram.

```
#pick a gene and remove NAs
gene1.class1 <- data.class1[1,]
gene1.class1.omitna <- gene1.class1[!is.na(gene1.class1)]
gene1.class2 <- data.class2[1,]
gene1.class2.omitna <- gene1.class2[!is.na(gene1.class2)]

#create boxplot of both classes
boxplot(gene1.class1.omitna, gene1.class2.omitna,
        col=c('red', 'blue'),
        names=c('class 1', 'class 2'),
        main='Example gene from DLBCL cDNA 2-channel dataset',
        axes=F, ylab='log2(ratio intensity)')
axis(2)
axis(1, at=c(1,2), c("class 1", "class 2"))
```

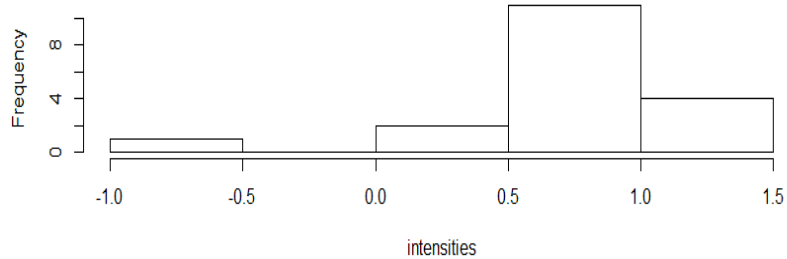
Example gene from DLBCL cDNA 2-channel dataset



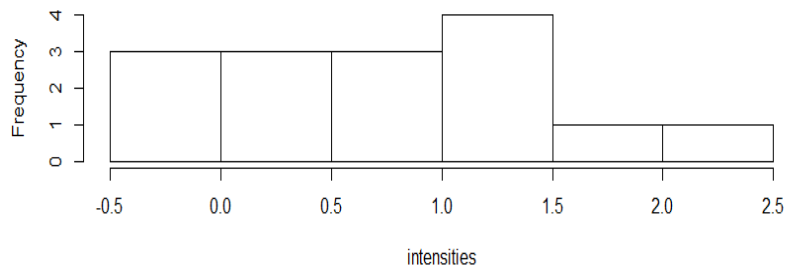
```
par(mfrow=c(2,1))
hist(gene1.class1.omitna,
     main='Example gene from DLBCL cDNA 2-channel dataset - class 1',
     xlab='intensities')

hist(gene1.class2.omitna,
     main='Example gene from DLBCL cDNA 2-channel dataset - class 2',
     xlab='intensities')
```

Example gene from DLBCL cDNA 2-channel dataset - class 1



Example gene from DLBCL cDNA 2-channel dataset - class 2



6.) Calculate the pooled variance as coded in the lecture notes, and calculate the minimum sample size necessary to detect a 1.5 fold difference (at 80% power and 99% confidence).

```
install.packages('pwr')
library(pwr)

# pooled variance
x <- gene1.class1.omitna
y <- gene1.class2.omitna
nx <- length(x)
ny <- length(y)
pool.var <- (((nx-1)*var(x)) + ((ny-1)*var(y)))/(nx+ny-2)
pool.var

[1] 0.2927542

#minimum sample size
dif.1.5fold <- log2(1.5)/sqrt(pool.var)
pl.ss1.5 <- pwr.t.test(d=dif.1.5fold,sig.level=.01,
                      power=0.8,type="two.sample")
pl.ss1.5
```

Two-sample t test power calculation

```
          n = 21.70052
          d = 1.081126
sig.level = 0.01
power = 0.8
alternative = two.sided
```

NOTE: n is number in *each* group

7.) Now calculate the sample size required for the same gene selected in #5 using the empirically determined delta between the two groups, assuming 99% confidence and 80% power.

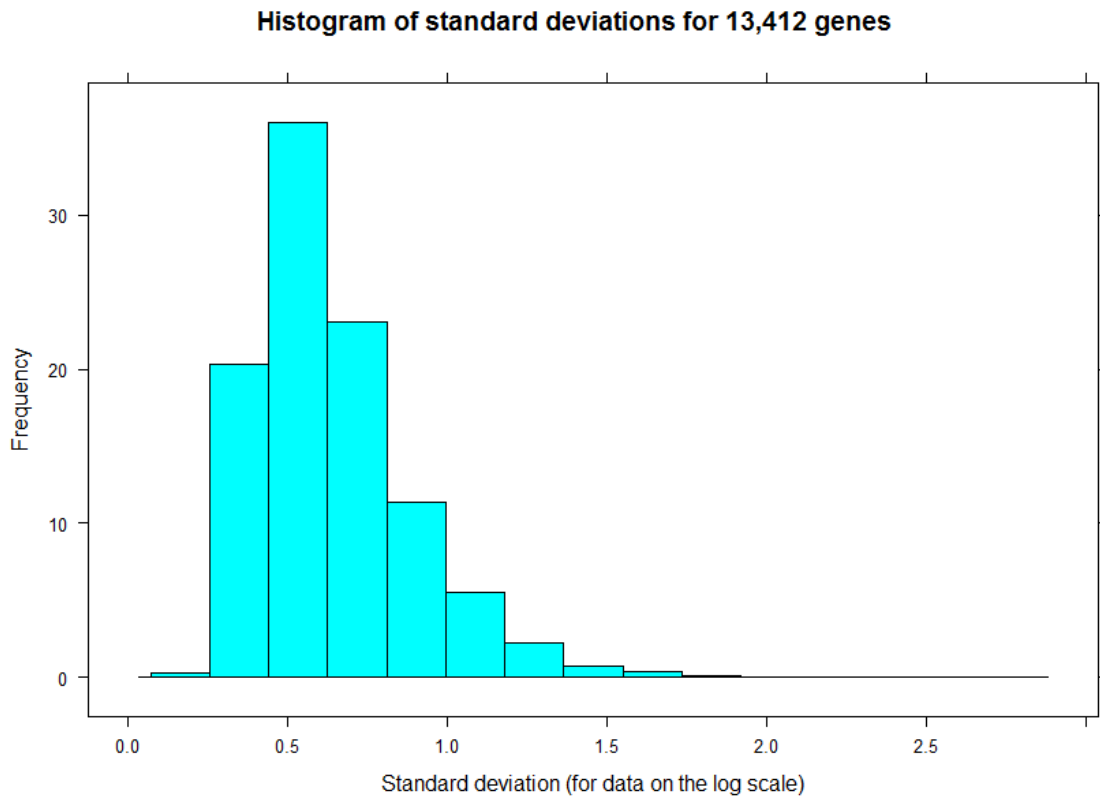
```
dif <- abs(mean(x)-mean(y))/sqrt(pool.var)
pl.ssdelta <- pwr.t.test(d=dif,sig.level=.01,
                        power=0.8,type="two.sample")
pl.ssdelta
```

Two-sample t test power calculation

```
          n = 2301.054
          d = 0.1007884
sig.level = 0.01
power = 0.8
alternative = two.sided
```

8.) Now load the ssize and gdata libraries, calculate the standard deviation for each gene in the matrix (Hint: use the na.rm=T argument), and plot a histogram of the standard deviations. Label the plot accordingly.

```
biocLite("ssize") # this also loaded gdata
library(ssize)
data.withsd <- transform(data, SD=apply(data,1,sd,na.rm=TRUE))
dim(data.withsd)
data.sd <- data.withsd[,41]
histogram(data.sd,
          main='Histogram of standard deviations for 13,412 genes',
          ylab='Frequency',
          xlab='Standard deviation (for data on the log scale)')
```



9.) Calculate and plot a proportion of genes vs. sample size graph to get an idea of the number of genes that have an adequate sample size for confidence=95%, effect size=3 (log2 transform for the function), and power=80%.

```
# plot a gene proportion vs. sample size plot using the same criteria
AND power=80%
```

```

all.size <- ssize(sd=data.sd, delta=log2(3.0), sig.level=0.05,
power=0.8)
ssize.plot(all.size, lwd=2, col="magenta", xlim=c(1,20))
xmax <- par("usr")[2]-1;
ymin <- par("usr")[3] + 0.05
legend(x=xmax, y=ymin, legend= strsplit( paste("fold change=",3.0,"",
"alpha=", 0.05, "", "power=",0.8,"", "# genes=", length(data.sd),
sep=''), ", " )[[1]], xjust=1, yjust=0, cex=1.0)
title("Sample Size to Detect 2-Fold Change")

```

