## Analyze Protein Data

September 9, 2014

### 1 Cleaning the data

```
## Protein of interest for tech paper 01/27/2014
library("lme4")
library("plyr")
library("ggplot2")
library("MASS")
# read data
dat <- read.csv("239 tech paper_08_19_2014.csv")</pre>
dim(dat)
## [1] 241 66
x <- colnames(dat) # column name of data
X
    [1] "BVAExport"
                                              "Image"
##
   [3] "Gel_1_Run_1_Pig_1807_45034.CY3.gel" "X"
##
   [5] "Gel_1_Run_1_Pig_1807_45034.CY5.gel" "X.1"
   [7] "Gel_1_Run_2_Pig_1807_45046.CY3.gel" "X.2"
## [9] "Gel_1_Run_2_Pig_1807_45046.CY5.gel" "X.3"
## [11] "Gel_2_Run_1_4810_45035.CY3.gel"
## [13] "Gel_2_Run_1_4810_45035.CY5.gel"
## [15] "Gel_2_Run_2_Pig_4810_45047.CY3.gel" "X.6"
## [17] "Gel_2_Run_2_Pig_4810_45047.CY5.gel" "X.7"
## [19] "Gel_3_Run_1_Pig_1209_45036.CY3.gel" "X.8"
## [21] "Gel_3_Run_1_Pig_1209_45036.CY5.gel" "X.9"
## [23] "Gel_3_Run_2_Pig_1209_45048.CY3.gel" "X.10"
## [25] "Gel_3_Run_2_Pig_1209_45048.CY5.gel" "X.11"
## [27] "Gel_4_Run_1_Pig_1906_45037.CY3.gel" "X.12"
## [29] "Gel_4_Run_1_Pig_1906_45037.CY5.gel" "X.13"
## [31] "Gel_4_Run_2_Pig_1906_45049.CY3.gel" "X.14"
## [33] "Gel_4_Run_2_Pig_1906_45049.CY5.gel" "X.15"
## [35] "Gel_5_Run_1_Pig_3908_45038.CY3.gel" "X.16"
## [37] "Gel_5_Run_1_Pig_3908_45038.CY5.gel" "X.17"
## [39] "Gel_5_Run_2_Pig_3908_45050.CY3.gel" "X.18"
## [41] "Gel_5_Run_2_Pig_3908_45050.CY5.gel" "X.19"
## [43] "Gel_6_Run_1_Pig_3106_45039.CY3.gel" "X.20"
## [45] "Gel_6_Run_1_Pig_3106_45039.CY5.gel" "X.21"
## [47] "Gel_6_Run_2_Pig_3106_45051.CY3.gel" "X.22"
## [49] "Gel_6_Run_2_Pig_3106_45051.CY5.gel" "X.23"
```

```
## [51] "Gel_7_Run_1_Pig_2107_45040.CY3.gel" "X.24"
## [53] "Gel_7_Run_1_Pig_2107_45040.CY5.gel" "X.25"
## [55] "Gel_7_Run_2_Pig_2107_45052.CY3.gel" "X.26"
## [57] "Gel_7_Run_2_Pig_2107_45052.CY5.gel" "X.27"
## [59] "Gel_8_Run_1_Pig_2712_45041.CY3.gel" "X.28"
## [61] "Gel_8_Run_1_Pig_2712_45041.CY5.gel" "X.29"
## [63] "Gel_8_Run_2_Pig_2712_44495.CY3.gel" "X.30"
## [65] "Gel_8_Run_2_Pig_2712_44495.CY5.gel" "X.31"
y <- c( "Image", "X", paste("X.", 1:31, sep = "")) # Column name unsued
# Column name used in analysis including the id of protein and 16 gels
# of 8 animals, each gel runs twice
use_col <- setdiff(x,y)</pre>
use_col
## [1] "BVAExport"
                                              "Gel_1_Run_1_Pig_1807_45034.CY3.gel"
## [3] "Gel_1_Run_1_Pig_1807_45034.CY5.gel" "Gel_1_Run_2_Pig_1807_45046.CY3.gel"
## [5] "Gel_1_Run_2_Pig_1807_45046.CY5.gel" "Gel_2_Run_1_4810_45035.CY3.gel"
## [7] "Gel_2_Run_1_4810_45035.CY5.gel"
                                              "Gel_2_Run_2_Pig_4810_45047.CY3.gel"
## [9] "Gel_2_Run_2_Pig_4810_45047.CY5.gel" "Gel_3_Run_1_Pig_1209_45036.CY3.gel"
## [11] "Gel_3_Run_1_Pig_1209_45036.CY5.gel" "Gel_3_Run_2_Pig_1209_45048.CY3.gel"
## [13] "Gel_3_Run_2_Pig_1209_45048.CY5.gel" "Gel_4_Run_1_Pig_1906_45037.CY3.gel"
## [15] "Gel_4_Run_1_Pig_1906_45037.CY5.gel" "Gel_4_Run_2_Pig_1906_45049.CY3.gel"
## [17] "Gel_4_Run_2_Pig_1906_45049.CY5.gel" "Gel_5_Run_1_Pig_3908_45038.CY3.gel"
## [19] "Gel_5_Run_1_Pig_3908_45038.CY5.gel" "Gel_5_Run_2_Pig_3908_45050.CY3.gel"
## [21] "Gel_5_Run_2_Pig_3908_45050.CY5.gel" "Gel_6_Run_1_Pig_3106_45039.CY3.gel"
## [23] "Gel_6_Run_1_Pig_3106_45039.CY5.gel" "Gel_6_Run_2_Pig_3106_45051.CY3.gel"
## [25] "Gel_6_Run_2_Pig_3106_45051.CY5.gel" "Gel_7_Run_1_Pig_2107_45040.CY3.gel"
## [27] "Gel_7_Run_1_Pig_2107_45040.CY5.gel" "Gel_7_Run_2_Pig_2107_45052.CY3.gel"
## [29] "Gel_7_Run_2_Pig_2107_45052.CY5.gel" "Gel_8_Run_1_Pig_2712_45041.CY3.gel"
## [31] "Gel_8_Run_1_Pig_2712_45041.CY5.gel" "Gel_8_Run_2_Pig_2712_44495.CY3.gel"
## [33] "Gel_8_Run_2_Pig_2712_44495.CY5.gel"
dat_used <- as.matrix(dat[, use_col])</pre>
dim(dat_used)
## [1] 241 33
# The first row is sample classification, the second row is the name of Std. Abund,
# therefore the data actually in use is the dat_used except the first 2 rows
dat_final <- matrix(as.numeric(dat_used[-c(1:2),]),</pre>
                    nrow = nrow(dat_used) - 2,
                    ncol = ncol(dat_used),
                    byrow = F)
dim(dat_final)
## [1] 239 33
```

```
# sample type of each sample
group <- dat_used[1,-1]</pre>
group
## Gel_1_Run_1_Pig_1807_45034.CY3.gel Gel_1_Run_1_Pig_1807_45034.CY5.gel
##
                               "Whole"
                                                                 "Depleted"
   Gel_1_Run_2_Pig_1807_45046.CY3.gel Gel_1_Run_2_Pig_1807_45046.CY5.gel
##
##
                               "Whole"
                                                                 "Depleted"
##
       Gel_2_Run_1_4810_45035.CY3.gel
                                            Gel_2_Run_1_4810_45035.CY5.gel
##
                            "Depleted"
                                                                    "Whole"
   Gel_2_Run_2_Pig_4810_45047.CY3.gel Gel_2_Run_2_Pig_4810_45047.CY5.gel
##
##
                            "Depleted"
                                                                    "Whole"
##
   Gel_3_Run_1_Pig_1209_45036.CY3.gel Gel_3_Run_1_Pig_1209_45036.CY5.gel
##
                               "Whole"
                                                                 "Depleted"
## Gel_3_Run_2_Pig_1209_45048.CY3.gel Gel_3_Run_2_Pig_1209_45048.CY5.gel
                               "Whole"
##
                                                                 "Depleted"
## Gel_4_Run_1_Pig_1906_45037.CY3.gel Gel_4_Run_1_Pig_1906_45037.CY5.gel
                            "Depleted"
                                                                    "Whole"
##
## Gel_4_Run_2_Pig_1906_45049.CY3.gel Gel_4_Run_2_Pig_1906_45049.CY5.gel
##
                            "Depleted"
                                                                    "Whole"
## Gel_5_Run_1_Pig_3908_45038.CY3.gel Gel_5_Run_1_Pig_3908_45038.CY5.gel
                               "Whole"
                                                                 "Depleted"
##
## Gel_5_Run_2_Pig_3908_45050.CY3.gel Gel_5_Run_2_Pig_3908_45050.CY5.gel
##
                               "Whole"
                                                                 "Depleted"
## Gel_6_Run_1_Pig_3106_45039.CY3.gel Gel_6_Run_1_Pig_3106_45039.CY5.gel
                            "Depleted"
                                                                    "Whole"
##
## Gel_6_Run_2_Pig_3106_45051.CY3.gel Gel_6_Run_2_Pig_3106_45051.CY5.gel
##
                            "Depleted"
                                                                    "Whole"
## Gel_7_Run_1_Pig_2107_45040.CY3.gel Gel_7_Run_1_Pig_2107_45040.CY5.gel
##
                               "Whole"
                                                                 "Depleted"
## Gel_7_Run_2_Pig_2107_45052.CY3.gel Gel_7_Run_2_Pig_2107_45052.CY5.gel
##
                               "Whole"
                                                                 "Depleted"
## Gel_8_Run_1_Pig_2712_45041.CY3.gel Gel_8_Run_1_Pig_2712_45041.CY5.gel
##
                            "Depleted"
                                                                    "Whole"
## Gel_8_Run_2_Pig_2712_44495.CY3.gel Gel_8_Run_2_Pig_2712_44495.CY5.gel
##
                            "Depleted"
                                                                    "Whole"
\# Obtain data for each sample type: depleted and whole, the first column of dat_final
# contains name of protein
deplete <- dat_final[,-1][,group=="Depleted"]</pre>
dim(deplete)
## [1] 239 16
whole <- dat_final[,-1][,group=="Whole"]</pre>
dim(whole)
## [1] 239
            16
# Find out which Cy is used for each run
group [group == "Depleted"] # Cy for depleted sample : rep(c(5,5,3,3),4)
## Gel_1_Run_1_Pig_1807_45034.CY5.gel Gel_1_Run_2_Pig_1807_45046.CY5.gel
```

```
##
                            "Depleted"
                                                                "Depleted"
##
       Gel_2_Run_1_4810_45035.CY3.gel Gel_2_Run_2_Pig_4810_45047.CY3.gel
                                                                "Depleted"
##
                            "Depleted"
##
  Gel_3_Run_1_Pig_1209_45036.CY5.gel Gel_3_Run_2_Pig_1209_45048.CY5.gel
##
                            "Depleted"
                                                                "Depleted"
##
   Gel_4_Run_1_Pig_1906_45037.CY3.gel Gel_4_Run_2_Pig_1906_45049.CY3.gel
##
                            "Depleted"
                                                                "Depleted"
##
  Gel_5_Run_1_Pig_3908_45038.CY5.gel Gel_5_Run_2_Pig_3908_45050.CY5.gel
##
                            "Depleted"
                                                                "Depleted"
  Gel_6_Run_1_Pig_3106_45039.CY3.gel Gel_6_Run_2_Pig_3106_45051.CY3.gel
##
##
                            "Depleted"
                                                                "Depleted"
## Gel_7_Run_1_Pig_2107_45040.CY5.gel Gel_7_Run_2_Pig_2107_45052.CY5.gel
                            "Depleted"
## Gel_8_Run_1_Pig_2712_45041.CY3.gel Gel_8_Run_2_Pig_2712_44495.CY3.gel
                            "Depleted"
                                                                "Depleted"
group [group == "Whole"] # Cy for whole sample : rep(c(3,3,5,5),4)
## Gel_1_Run_1_Pig_1807_45034.CY3.gel Gel_1_Run_2_Pig_1807_45046.CY3.gel
##
                               "Whole"
                                                                   "Whole"
##
       Gel_2_Run_1_4810_45035.CY5.gel Gel_2_Run_2_Pig_4810_45047.CY5.gel
##
                               "Whole"
                                                                   "Whole"
   Gel_3_Run_1_Pig_1209_45036.CY3.gel Gel_3_Run_2_Pig_1209_45048.CY3.gel
##
                               "Whole"
                                                                   "Whole"
## Gel_4_Run_1_Pig_1906_45037.CY5.gel Gel_4_Run_2_Pig_1906_45049.CY5.gel
                               "Whole"
##
                                                                    "Whole"
## Gel_5_Run_1_Pig_3908_45038.CY3.gel Gel_5_Run_2_Pig_3908_45050.CY3.gel
                               "Whole"
##
                                                                   "Whole"
## Gel_6_Run_1_Pig_3106_45039.CY5.gel Gel_6_Run_2_Pig_3106_45051.CY5.gel
                               "Whole"
##
                                                                   "Whole"
## Gel_7_Run_1_Pig_2107_45040.CY3.gel Gel_7_Run_2_Pig_2107_45052.CY3.gel
                               "Whole"
                                                                   "Whole"
##
## Gel_8_Run_1_Pig_2712_45041.CY5.gel Gel_8_Run_2_Pig_2712_44495.CY5.gel
                               "Whole"
```

# 2 Function to fit a Linear Mixed Effect Model for each spot in each sample

Function to fit a linear mixed model for each spot, with fixed effects are Cy, RFI Line and the random effect is animal.

```
out_model <- function(x, depleted){ # x is the row of data (i.e., data of each protein spot)
  if (depleted == "TRUE"){
    cy <- as.factor(rep(c(5,5,3,3),4))
  } else {
    cy <- as.factor(rep(c(3,3,5,5),4))
  }
  animal <- as.factor(rep(1:8, each = 2))
  line <- as.factor(rep(c(1,2), each = 8))
  # check if all obsetvations for one Cy are missing or not
  if ((sum(is.na(x[cy==3]))==8|sum(is.na(x[cy==5]))==8) &</pre>
```

```
(sum(is.na(x[line==1]))==8|sum(is.na(x[line==2]))==8)){
    model <- lmer(x~ (1|animal), na.action="na.omit")</pre>
    #mean_est <- summary(model)$coeff[,1]</pre>
    sd_est <- as.vector(sqrt(summary(model)$vcov))</pre>
    #str(summary(model))
  }
  if ((sum(is.na(x[cy==3]))==8|sum(is.na(x[cy==5]))==8) &
        ((sum(is.na(x[line==1]))!=8)&(sum(is.na(x[line==2]))!=8))){
    model <- lmer(x~ line + (1|animal), na.action="na.omit")</pre>
    #mean_est <- summary(model)$coeff[1,1] + summary(model)$coeff[2,1]/2</pre>
    sd_est \leftarrow as.vector(sqrt(t(c(1,1/2)) %*%summary(model)$vcov%*%c(1,1/2)))
  }
  if ((sum(is.na(x[cy==3]))!=8|sum(is.na(x[cy==5]))!=8) &
        ((sum(is.na(x[line==1]))==8)&(sum(is.na(x[line==2]))==8))){
    model <- lmer(x~ cy + (1|animal), na.action="na.omit")</pre>
    \#mean\_est \leftarrow summary(model)\$coeff[1,1] + summary(model)\$coeff[2,1]/2
    sd_est \leftarrow as.vector(sqrt(t(c(1,1/2)) %*%summary(model)$vcov%*%c(1,1/2)))
  if ((sum(is.na(x[cy==3]))!=8& sum(is.na(x[cy==5]))!=8) &
        ((sum(is.na(x[line==1]))!=8)&(sum(is.na(x[line==2]))!=8))){
    model <- lmer(x~ cy + line + (1|animal), na.action="na.omit")</pre>
    #str(summary(model))
    mean_est <- summary(model)$coeff[1,1] +</pre>
       summary(model)$coeff[2,1]/2 +
      summary(model)$coeff[3,1]/2
    sd_est \leftarrow as.vector(sqrt(t(c(1, 1/2, 1/2))%*% summary(model)$vcov %*%c(1,1/2, 1/2))) }
 return( sd_est)
sd_depleted <- laply(1:dim(dat_final)[1], function(i)out_model(deplete[i,], depleted = "TRUE"))</pre>
sd_whole <- laply(1:dim(dat_final)[1], function(i)out_model(whole[i,], depleted = "FALSE"))</pre>
```

### 3 Results of Comparison of Standard Errors between Two Sample Types

## 3.1 Proportion of protein spots whose standard error in the depleted samples larger than that one in the whole samples

Proportion of protein spots whose standard error in the depleted samples larger than that one in the whole samples.

```
mean(sd_depleted > sd_whole)
## [1] 0.5941
```

Figure 1 show the log of standard error of all protein spots in 2 sample types.

```
log_sd <- data.frame(
  logsd = log(c(sd_whole, sd_depleted)),
  sample = rep(c("whole", "depleted"), each = length(sd_whole)))
# write.table(log_sd, file = "log_sd.txt")

p <- ggplot(log_sd, aes(sample, logsd))
p + geom_boxplot(aes(fill = sample)) +
  ggtitle("Standard Error for Each Sample Type") +
  scale_fill_discrete(name= "Sample Type") +
  xlab("Sample Type") +
  ylab("log(Standard Error)") +
  theme(text = element_text(size=11))</pre>
```

#### 3.2 One-sided Wilcoxon test

Consider a Wilcoxon signed-rank test H0: The distribution of the standard errors across protein is the same for depleted samples and whole samples, H1: the standard errors tend to be larger for depleted samples than for whole samples. The test using log-transformed standard errors has p-value = 0.004275.

```
# One-sided Test for the log-transformed standard errors
wilcox.test(log(sd_depleted/sd_whole), alternative = "greater")

##
## Wilcoxon signed rank test with continuity correction
##
## data: log(sd_depleted/sd_whole)
## V = 17154, p-value = 0.004275
## alternative hypothesis: true location is greater than 0
```

### 3.3 Boxplot for logsd(deplete/whole)

Figure 2 show the log ratio of the standard errors of two sample types.

```
logratiosd <- log(sd_depleted/sd_whole)
boxplot(logratiosd, main = "Log(sd_depleted/sd_whole)")
abline(h = 0)</pre>
```

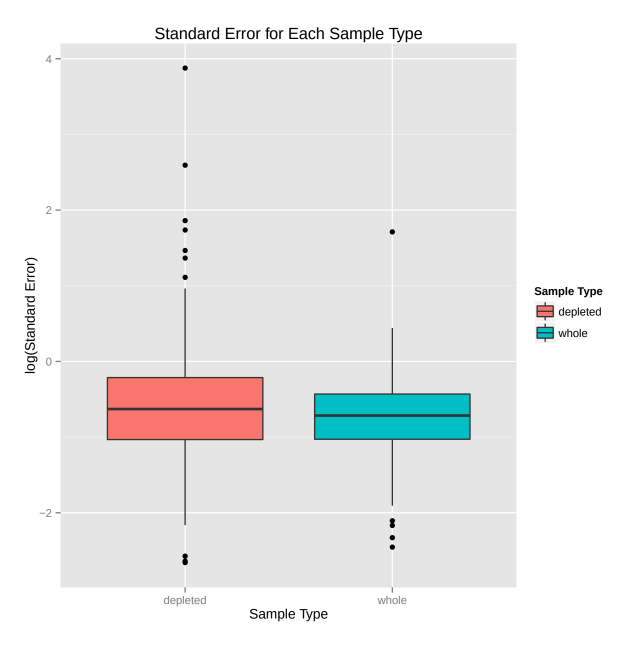


Figure 1: Log(Standard Error) for Each Sample Type

## Log(sd\_depleted/sd\_whole)

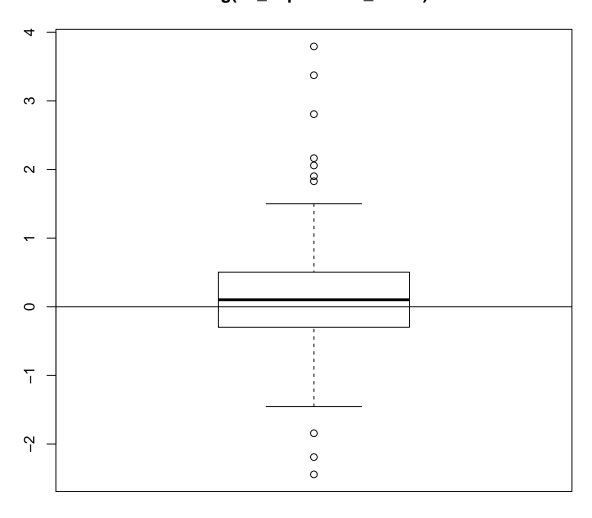


Figure 2: Log Ratio of Standard Error for Each Sample Type