EGCG blood sample gene expression Analysis

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### This Rmarkdown document analyzes the gene expression data from blood samples of overweight people before being treated with EGCG pills twice a day for 70 days and after. There were some that were treated additionally with flavanoids, fish oil, and vitamin C in the quenterin samples. This data was pulled from the NCBI GEO gene expression data and combined with the HGNC names from BioMart of Ensembl using the Affymetrix human genome 1.0 version 1 ID names.

Read in all the data tables

#pre\_egcg <- read.csv('pre\_placebo\_EGCG.csv',sep=',', header=TRUE,  
# na.strings=c('',' '), skip=1)  
#pre\_egcg\_quer <- read.csv('pre\_vitC\_flavanoids\_fishOil\_EGCG.csv',   
# sep=',', header=TRUE, na.strings=c('',' '), skip=1)  
#post\_egcg <- read.csv('post\_placebo\_EGCG.csv', sep=',', header=TRUE,  
# na.strings=c('',' '), skip=1)  
#post\_egcg\_quer <- read.csv('post\_vitaminC\_flavanoids\_fishOil\_EGCG.csv',  
# sep=',', header=TRUE, na.strings=c('',' '), skip=1)  
#hgnc <- read.csv('HGNC\_bioMart\_affy\_1\_v1\_IDs.csv', sep=',',  
# header=TRUE, na.strings=c('',' '))

Combine the HGNC names to each of the four tables and make the table row names.

#HGNC <- hgnc[,4:5]  
#HGNC <- HGNC[!duplicated(HGNC$Affy\_HuGene\_1\_v1\_probe\_ID),]  
#colnames(HGNC)[2] <- 'Affy'

#post\_egcg2 <- merge(HGNC,post\_egcg, by.x='Affy',  
# by.y='ID\_REF')  
  
#post\_egcg\_quer2 <- merge(HGNC, post\_egcg\_quer,  
# by.x='Affy',by.y='ID\_REF')  
  
#pre\_egcg2 <- merge(HGNC, pre\_egcg, by.x='Affy',  
# by.y='ID\_REF')  
  
#pre\_egcg\_quer2 <- merge(HGNC, pre\_egcg\_quer,  
# by.x='Affy',  
# by.y='ID\_REF')

There are duplicate HGNC names because of the different Affymetrix IDs, this can be fixed by grouping by genes and getting the mean of those genes and creating a table for each gene by means using the dplyr package.This is time consuming to run because it groups all the genes then it gets the means for each group by sample and outputs that table. It takes 20-30 minutes for each table grouping by genes and then taking the means of each gene per sample.

library(dplyr)

Remove the Affymetrix ID from the four tables because it isn’t needed anymore.

#post\_eq <- post\_egcg\_quer2[,-1]  
#post\_eg <- post\_egcg2[,-1]  
#pre\_eq <- pre\_egcg\_quer2[,-1]  
#pre\_eg <- pre\_egcg2[,-1]

EGCG only pre-treatment samples

#samples <- colnames(pre\_eg)[2:8]  
#Samples <- as.vector(samples)  
#Pre\_E <- pre\_eg %>% group\_by(HGNC.Symbol) %>%   
# summarise\_at(vars(Samples), mean, na.rm=TRUE)

EGCG+quenterin pre-treatment samples

#samples1 <- colnames(pre\_eq)[2:8]  
#Samples1 <- as.vector(samples1)  
#Pre\_Q <- pre\_eq %>% group\_by(HGNC.Symbol) %>%   
# summarise\_at(vars(Samples1), mean, na.rm=TRUE)

EGCG samples and EGCG+flavanoids+fish oil+vitamin C can be combined into a pretreatment table.

#pre <- merge(Pre\_E, Pre\_Q, by.x='HGNC.Symbol', by.y='HGNC.Symbol')  
#row.names(pre) <- pre$HGNC.Symbol  
#PRE <- pre[,-1]

Post EGCG and Quercetin

#samples2 <- colnames(post\_eq)[2:8]  
#Samples2 <- as.vector(samples2)  
#Post\_Q <- post\_eq %>% group\_by(HGNC.Symbol) %>%   
# summarise\_at(vars(Samples2), mean, na.rm=TRUE)

Post EGCG only

#samples3 <- colnames(post\_eg)[2:8]  
#Samples3 <- as.vector(samples3)  
#Post\_E <- post\_eg %>% group\_by(HGNC.Symbol) %>%   
# summarise\_at(vars(Samples3), mean, na.rm=TRUE)

Give the post-treatment samples row names and remove the HGNC fields

#Post\_E <- as.data.frame(Post\_E)  
#Post\_Q <- as.data.frame(Post\_Q)  
#row.names(Post\_E) <- Post\_E$HGNC.Symbol  
#row.names(Post\_Q) <- Post\_Q$HGNC.Symbol  
#Post\_E <- Post\_E[,-1]  
#Post\_Q <- Post\_Q[,-1]

Write the new tables to csv files

#write.csv(PRE, 'pre\_combined.csv', row.names=TRUE)  
#write.csv(Post\_E, 'Post\_EGCG\_combine.csv', row.names=TRUE)  
#write.csv(Post\_Q, 'Post\_EGCG\_Quer\_combine.csv', row.names=TRUE)

### To save time, the above has been commented out, and the files can be opened to run the rest of the code.

PRE <- read.csv('pre\_combined.csv', sep=',', header=TRUE,  
 na.strings=c('',' '), row.names=1)  
Post\_E <- read.csv('Post\_EGCG\_combine.csv', sep=',', header=TRUE,  
 na.strings=c('',' '), row.names=1)  
Post\_Q <- read.csv('Post\_EGCG\_Quer\_combine.csv', sep=',', header=TRUE,  
 na.strings=c('',' '), row.names=1)

Get the means of the pre-treatment samples with rowMeans() per gene and the post-treatment samples of EGCG and EGCG+Quercentin, then attach to each of those tables as the mean of those samples of gene expression values.

PRE$PRE\_Means <- rowMeans(PRE)  
Post\_Q$Post\_Q\_Means <- rowMeans(Post\_Q)  
Post\_E$Post\_E\_Means <- rowMeans(Post\_E)

colnames(PRE)[1:14] <- paste('pre\_',sep='', colnames(PRE)[1:14])

colnames(Post\_E)[1:7] <- paste('post\_EG\_', sep='',colnames(Post\_E)[1:7])

colnames(Post\_Q)[1:7] <- paste('post\_EQ\_', sep='', colnames(Post\_Q)[1:7])

Combine all tables into one with the means of each type of sample at the start of the columns

All <- cbind(PRE[,15],Post\_E[,8],Post\_Q[,8],PRE[,1:14], Post\_E[1:7],  
 Post\_Q[1:7])  
colnames(All)[1:3] <- c('Pre\_Means','Post\_EGCG\_Means','Post\_EGCG\_Quercentin\_Means')

Now for the differential expression in Means using dplyr. Here the overweight is the initial value minus the value after treatment type.

DE <- All %>% mutate(DE\_EGCG=All$Pre\_Means-All$Post\_EGCG\_Means)  
DE2 <- DE %>% mutate(DE\_Quercentin=DE$Pre\_Means-All$Post\_EGCG\_Quercentin\_Means)  
  
DE3 <- DE2[,c(32,33,1:31)] #table with both differential values  
row.names(DE3) <- row.names(All)

Get the over and under expressed genes after treatment compared to before treatment of EGCG and EGCG+quercentin in overweight females.The negative DE values means the treatment made the gene over expressed, since the initial values minus treatment values would be negative only if treatment values are greater than initial values.These two tables are the top20 up regulated genes after treatment with either EGCG or EGCG+Quercentin

DE\_20up\_egcg <- DE3[order(DE3$DE\_EGCG, decreasing=FALSE)[0:20],]  
DE\_20up\_egcg\_quer <- DE3[order(DE3$DE\_Quercentin,decreasing=FALSE)[0:20],]

The top 20 up regulated genes after treatment with only EGCG

row.names(DE\_20up\_egcg)

## [1] "TMEM176A" "HBZ" "TMEM176B" "FAU" "HBZP1"   
## [6] "RNY3P16" "IGKV1-5" "RNA5SP63" "TREML4" "CYP4F35P"   
## [11] "ALOX15" "IGKV3-11" "IGKV2-30" "ZFP57" "RNA5SP449"  
## [16] "IGKV1-33" "CAV1" "RNU6-893P" "POU2AF1" "IGKV2-24"

The top 20 up regulated genes after treatment with EGCG and Quercenting

row.names(DE\_20up\_egcg\_quer)

## [1] "KARSP3" "GSTM1" "RNU4-51P" "RNU6-1024P" "COX7B"   
## [6] "OCLN" "CTNNAL1" "RNA5SP30" "C4BPA" "CD274"   
## [11] "LINC01356" "RNU6-522P" "HMGB3P24" "BBOF1" "RPL36AP26"   
## [16] "SLPI" "RNU6-943P" "SLC3A1" "RPS7" "RNU6-853P"

Now look at the top 20 down regulated genes after treatment with EGCG or EGCG and Quercentin

DE\_20down\_egcg <- DE3[order(DE3$DE\_EGCG, decreasing=TRUE)[0:20],]  
DE\_20down\_egcg\_quer <- DE3[order(DE3$DE\_Quercentin,decreasing=TRUE)[0:20],]

The top 20 genes down regulated in overweight females after treatment with only EGCG are:

row.names(DE\_20down\_egcg)

## [1] "RNU4-2" "TMTC1" "SLFN14" "SLC14A1" "GSTM1"   
## [6] "SCARNA10" "RNF182" "LINC00189" "XK" "TSPAN7"   
## [11] "SFRP2" "ORM1" "LRP1" "RPPH1" "ALDH5A1"   
## [16] "LGALS2" "IFIT1B" "RNA5SP242" "UBE2O" "OSBP2"

The top 20 genes down regulated in overweight females after treatment with EGCG and Quercentin are:

row.names(DE\_20down\_egcg\_quer)

## [1] "HBZ" "RNU6-1124P" "RNU6-921P" "IGKV2-26" "IGKV1D-27"   
## [6] "ALOX15" "TMEM176A" "RNU6-1315P" "KIAA1324" "IL1RL1"   
## [11] "SNORA38B" "HLA-DOA" "IGHV3-47" "VTRNA1-1" "SMPD3"   
## [16] "TMEM176B" "SNORA23" "RNU5A-1" "FOLR3" "RN7SL218P"

Now for the fold change in genes to see which genes changed the most in samples after treatment with EGCG or EGCG and Quercentin using the ratio of treatment type value/initial value.

FC <- DE3 %>% mutate(FC\_egcg = DE3$Post\_EGCG\_Means/DE3$Pre\_Means)  
FC <- FC[,c(34,1:33)]  
row.names(FC) <- row.names(DE3)  
  
FC2 <- FC %>% mutate(FC\_egcg\_quer = FC$Post\_EGCG\_Quercentin\_Means/FC$Pre\_Means)  
FC2 <- FC2[,c(35,1:34)]  
row.names(FC2) <- row.names(FC)

The top 20 genes that changed the most have a higher fold change value and are:

FC\_top20\_egcg <- FC2[order(FC2$FC\_egcg, decreasing=TRUE)[0:20],]   
FC\_top20\_egcg\_quer <- FC2[order(FC2$FC\_egcg\_quer, decreasing=TRUE)[0:20],]

The top 20 genes with the most fold change in EGCG treatment only are:

row.names(FC\_top20\_egcg)

## [1] "HBZ" "FAU" "RNA5SP449" "RNY3P16" "MTCO2P29"   
## [6] "TMEM176A" "TMEM176B" "PSG1" "CYP4F35P" "RN7SKP198"   
## [11] "RNA5SP63" "NPY5R" "MIR548AD" "RN7SL452P" "GPR87"   
## [16] "RNU6-1024P" "BUD31P2" "RNU6-893P" "ZNF578" "RNU6-263P"

The top 20 genes with the most fold change in EGCG and Quercentin treatment are:

row.names(FC\_top20\_egcg\_quer)

## [1] "KARSP3" "RNU4-51P" "RNU6-1244P" "RNU6-403P" "RNU6-1024P"  
## [6] "COX7B" "ZNF847P" "RNU6-853P" "RNU6-943P" "RNA5SP30"   
## [11] "OR5M10" "RNY1P5" "DDX18P6" "HMGB3P24" "LINC01356"   
## [16] "MTCO1P11" "RPS7" "RNU6-98P" "FAU" "GSTM1"

Create a data table of the statistical values only

Statistical <- FC2[,1:7] #all genes, not filtered for top 20

Create a table of all the samples with all the genes that are in some top 20 category from the statistical values produced earlier.

Fold1 <- FC\_top20\_egcg[,c(2,8:35)]  
colnames(Fold1)[1] <- 'change'  
Fold1$value <- rep('Fold Change EGCG',20)  
  
Fold2 <- FC\_top20\_egcg\_quer[,c(1,8:35)]  
colnames(Fold2)[1] <- 'change'  
Fold2$value <- rep('Fold Change EGCG+Quercentin',20)  
  
DE\_down\_1 <- DE\_20down\_egcg[,c(1,6:33)]  
colnames(DE\_down\_1)[1] <- 'change'  
DE\_down\_1$value <- rep('Down Regulated EGCG',20)  
  
DE\_down\_2 <- DE\_20down\_egcg\_quer[,c(2,6:33)]  
colnames(DE\_down\_2)[1] <- 'change'  
DE\_down\_2$value <- rep('Down Regulated EGCG+Quercentin',20)  
  
DE\_up\_1 <- DE\_20up\_egcg[,c(1,6:33)]  
colnames(DE\_up\_1)[1] <- 'change'  
DE\_up\_1$value <- rep('Up Regulated EGCG',20)  
  
DE\_up\_2 <- DE\_20up\_egcg\_quer[,c(2,6:33)]  
colnames(DE\_up\_2)[1] <- 'change'  
DE\_up\_2$value <- rep('Up Regulated EGCG+Quercentin',20)  
  
Top20 <- rbind(Fold1, Fold2, DE\_down\_1, DE\_down\_2, DE\_up\_1, DE\_up\_2)  
Top20 <- Top20[,c(30,1:29)]

Write the previous table out to csv with the change value and corresponding type measure each of the genes in a top 20 category produced.

write.csv(Top20, 'Top20\_all.csv', row.names=TRUE)

Now make the table of top genes (120) and the samples transposed with a new field for type of sample, so that it can be used to run machine learning on and predict the sample type the observation was from as in pre treatment, post treatment with EGCG or post treatment with EGCG and a combination of Vitatmin C, Niacin, and Quercentin fish oil.

Top20\_1 <- Top20[, -c(1:2)]  
  
ML\_Top20 <- t(Top20\_1)  
  
pre <- as.data.frame(rep('pre-treatment',14))  
post\_eg <- as.data.frame(rep('EGCG treatment',7))  
post\_eq <- as.data.frame(rep('EGCG and Quercentin treatment', 7))  
  
colnames(pre) <- 'Type'  
colnames(post\_eg) <- 'Type'  
colnames(post\_eq) <- 'Type'   
  
Type <- rbind(pre, post\_eg, post\_eq)  
  
genes <- Top20[,1:2]  
write.csv(genes, 'genes\_120\_top20.csv', row.names=TRUE)  
  
ML\_ready <- cbind(Type, ML\_Top20)  
  
write.csv(ML\_ready, 'ML\_ready.csv', row.names=TRUE)