Female tumor vs tumor adjacent

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5/22/2022

R Markdown

Title: "DEG Analysis Tumor vs Tumor Adjacent"

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Date: 05/22/2022

Purpose: This document is for the author to parse out the female tumor vs. tumor adjacent code from the "DEG changed comparison" file as well as demonstrate understanding of the code and theory ### **Libraries**

The first chunk of code is dedicated to installing the libraries. These libraries are to help execute the differential analysis and helps visualize the data. The code was not included for concision.

Environment parameters

This next section of code is dedicated to the environmental parameters. Environmental parameters are a series of variables and other code that will help make the rest of the script be easier to make and run later on.

Working Directory

A working directory is a code that iterates a file path on your computer th.t sets where the default location of any files that you read into R. Working directories work different in R files than R Markdowns. R Markdown files require directories to be defined at the end of each code chunk. Meaning from here on out you will see working directories being defined at the end of each code chunk.

```
setwd('~/R')
```

Defining Colors

This chunk defines color palette variables that are going to be used in plots later on the script. These variables are defined by conversiting BrewerCode palettes into palettes that can be used in R.

```
viralPalette <- brewer.pal(8, "Set1")
hbvColor <- viralPalette[1]
hcvColor <- viralPalette[2]
bothColor <- viralPalette[3]
neitherColor <- viralPalette[4]

sexTissuePalette <- brewer.pal(12, "Paired")
maleTumorColor <- sexTissuePalette[4]
maleAdjacentColor <- sexTissuePalette[3]
femaleTumorColor <- sexTissuePalette[6]
femaleAdjacentColor <- sexTissuePalette[5]
setwd('~/R')</pre>
```

Read in data

This code is where you read in all the data files that are going to be used in the script. The data is also converted into a variety of variables that makes the data easier to handle. The data is also cleaned up to make sure the analysis done later is accurate and precise.

```
metadata <- read.table("metadata_for_de.csv", row.names=1,header=TRUE, sep=",") #changing the na
me of the file
tumorAdjacentExp <- read.table("japan_all_samples_salmon_expression_counts.txt", row.names = 1,
header=TRUE) #changing the name of the file
colnames(tumorAdjacentExp) <- gsub("\\.", "-", colnames(tumorAdjacentExp)) #changing the column
names</pre>
```

Gene Length

This next code chunk is very similar However, it does calculate **gene length** which is the done by first defining a variable named "gene" and then changing the data type to a data frame. You then redefine "tumorAdjacentExp" (defined above) to have the rows of the previous "tumorAdjacentExp" and then the columns of "GENEID" that lies within "gene".

Gene length is then defined to have "with" of genes in the rows and 'end-start' as a column

Running Identical Function

We ran the identical function to see if the inputs of the match function are of length one. The function outputted a true value therefore they are identical.

```
genes <- read.table("gencodeTranscripts.txt", header=TRUE, sep="\t")
genes <- data.frame(genes)
tumorAdjacentExp <- tumorAdjacentExp[rownames(tumorAdjacentExp) %in% genes$GENEID ,]
genes <- genes[match(rownames(tumorAdjacentExp), genes$GENEID),]

identical(rownames(tumorAdjacentExp),genes$GENEID)</pre>
```

```
## [1] TRUE
```

```
# Calculating gene length, this is needed for calculating the FPKM values
genes$length <- with(genes, end - start)</pre>
```

xLow Quality

The next line shows a sample being removed due to low quality.

```
metadata<-metadata[!(metadata$ID == "RK023"), ]</pre>
```

Subsetting data

This next chunk of data is dedicated to sub-setting and organizing the data to make it easier to use going forward. Sub-setting means that the data is being organized to match a count matrix. In this specific case the count matrix is the sample ID attached to the tumors.

Running identical function

We ran the identical function to see if the columns are of length one in the match function. The function returned a TRUE value therefore they are of proper length.

```
tumorAdjacentExpSubset <- tumorAdjacentExp[,colnames(tumorAdjacentExp) %in% metadata$sampleid]
metadataSubset <- metadata[metadata$sampleid %in% colnames(tumorAdjacentExpSubset),]
metadataSubset <- metadataSubset[match(colnames(tumorAdjacentExpSubset), metadataSubset$sampleid),]
identical(colnames(tumorAdjacentExpSubset),metadataSubset$sampleid)</pre>
```

```
## [1] TRUE
```

```
rownames(metadataSubset) <- metadataSubset$sampleid</pre>
```

Tissue object

This next chunk of data is taking the meta data and subsetting it in such a way that converts a series of categorical variables into factors. This data also adds a tissue type.

```
metadataSubset$tumor <- as.numeric(grepl('tumor', metadataSubset$sampleid, ignore.case=T))
metadataSubset$gender_tissue <- paste(metadataSubset$Gender, metadataSubset$tumor, sep="_")
metadataSubset$gender_tissue_viral <- paste(metadataSubset$gender_tissue, metadataSubset$Virus_i
nfection, sep="_")
metadataSubset$library_type <- metadataSubset$strandedness
metadataSubset$library_type <- factor(metadataSubset$library_type)
metadataSubset$tumor <- factor(metadataSubset$tumor)
metadataSubset$Ta <- factor(metadataSubset$Ta)
metadataSubset$Portal_vein_invasion <- factor(metadataSubset$Portal_vein_invasion)
metadataSubset$Hepatic_vein_invasion <- factor(metadataSubset$Hepatic_vein_invasion)
metadataSubset$Bile_duct_invasion <- factor(metadataSubset$Bile_duct_invasion)
metadataSubset$Liver_fibrosisc <- factor(metadataSubset$Liver_fibrosisc)
metadataSubset$Prognosis <- factor(metadataSubset$Prognosis)</pre>
```

Creating DGE Object

This next chunk of code creates something called a DGEList Object. This object contains a dataset that is to be analyzed later in the script. Specifically the object contains:

- 1. Counts- numeric matrix containing read counts
- 2. group-vector giving the experimental conditiona for each sample
- 3. genes- data frame information for the genes for which we have count data
- 4. remove.zeros- whether to remove rows that have 0 total count

The last line of code takes the amount of samples and places them into a table for easy read and inspection.

```
##
##
   F 0 HBV F 0 HCV F 0 NBNC F 1 HBV F 1 HCV F 1 NBNC M 0 both M 0 HBV
##
                  34
                            3
                                     9
                                             36
                                                        3
                                                                 4
                                                                         33
          8
##
   M 0 HCV M 0 NBNC M 1 both M 1 HBV M 1 HCV M 1 NBNC
##
         59
                  25
                            4
                                    40
                                             71
                                                       25
```

Calculating fpkm values

This chunk of code takes the fpkm of all the genes in the dataset and calculates the mean. They also filter out genes that have a fpkm of 0.5 or lower.

Fpkm stands for fragments per kilo base of exon per million this term is interchangable with Rpkm (reads per kilobase of exon per million. This measure is a normalization method which allows us to compare gene expression levels by rescaling both library size and gene length.

Fpkm is calculated by multiplying the number of reads mapped to a gene by 1,000, 1,000,000 and then dividing that number by the total number of mapped reads to gene length in base pairs.

Please note that that calculation is done for RPKM which is analogous to Fpkm.

Here the fpkm is calculated from all the various tissue samples, are filtered for greater than 0.5 and put into a variable named "keep" which a cutoff point that is going to be used in limma/voom analysis

```
M_1_HBV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="M_1_HBV")],1,m
ean, na.rm=TRUE)
M_0_HBV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="M_0_HBV")],1,m
ean, na.rm=TRUE)
M_1_HCV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="M_1_HCV")],1,m
ean, na.rm=TRUE)
M 0 HCV mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="M 0 HCV")],1,m
ean, na.rm=TRUE)
M 1 HBVHCV mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="M 1 both"
)],1,mean,na.rm=TRUE)
M 0 HBVHCV mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="M 0 both"
)],1,mean,na.rm=TRUE)
M 1 NBNC mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="M 1 NBNC")],1
,mean,na.rm=TRUE)
M Ø NBNC mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="M Ø NBNC")],1
,mean,na.rm=TRUE)
F_1_HBV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="F_1_HBV")],1,m
ean, na.rm=TRUE)
F_0_HBV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="F_0_HBV")],1,m
ean, na.rm=TRUE)
F_1_HCV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="F_1_HCV")],1,m
ean, na.rm=TRUE)
F_0_HCV_mean_fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender_tissue_viral=="F_0_HCV")],1,m
ean, na.rm=TRUE)
F 1 NBNC mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="F 1 NBNC")],1
,mean,na.rm=TRUE)
F 0 NBNC mean fpkm <- apply(as.data.frame(fpkm)[(dge$samples$gender tissue viral=="F 0 NBNC")],1
,mean,na.rm=TRUE)
keep <- (M_1_HBV_mean_fpkm > 0.5 | M_0_HBV_mean_fpkm > 0.5 |
           M 1 HCV mean fpkm > 0.5 | M 0 HCV mean fpkm > 0.5 |
           M_1_HBVHCV_mean_fpkm > 0.5 | M_0_HBVHCV_mean_fpkm > 0.5 |
           M_1_NBNC_mean_fpkm > 0.5 | M_0_NBNC_mean_fpkm > 0.5 |
           F_1_HBV_mean_fpkm > 0.5 | F_0_HBV_mean_fpkm > 0.5 |
           F_1_HCV_mean_fpkm > 0.5 | F_0_HCV_mean_fpkm > 0.5 |
           F_1_NBNC_mean_fpkm > 0.5 | F_0_NBNC_mean_fpkm > 0.5)
```

DGE object organization

This chunk is further organizes and counts the libraries to be more tangible for later on as well as calculates the normalization factors (not normalizing the data) to use later on in the limma/voom DEG.

The normalization factors are calculated using Trimmed Mean of M-values (TMM). TMM is a between sample normalization that assumes that most genes are not differentially expressed. TMM normalizes the total RNA output among the samples, not considering gene length nor library size. TMM also considers the RNA population which makes it effective with samples that have diverse RNA repertoires.

TMM takes the library size normalized read count for each gene in each sample and calculates the log2 fold change between two samples (M-value). From there you calculate the absolute expression count (A values) which is the sum of the log2 fold change of treated sample count plus the log2 fold change of the control sample count divided by two.

M-values and A-values are double trimmed by 30% and 5% respectively. You then get the weight mean M after trimming and calculate the normalization factor.

```
dge <- dge[keep,,keep.lib.sizes=FALSE]
dge <- calcNormFactors(dge, method="TMM")
keep <- rowSums(dge$counts > 6) >= 10
dge <- dge[keep,,keep.lib.size=FALSE]
dge <- calcNormFactors(dge, method="TMM")</pre>
```

Counting number of FPKM genes

This code counts the number of genes that make it past the cutoff point.

```
# N of genes retained after filtering
dim(dge$genes)
```

```
## [1] 13384 7
```

DGE Analysis tumor vs. tumor- adjacent

This section is doing voom/limma and Go/KEGG analysis on all of the tumor vs. tumor adajcent samples

Design matrix

This code chunk is creating a design matrix to analyze tumor vs. tumor adjacent regardless of sex. This design matrix creates a model matrix that takes the inputs that we want to consider for later limma/voom analysis.

In this specific design matrix we are considering all tumor samples and all tumor adjacent samples.

The colnames code is just renaming columns of the design matrix to identify what they are for easier readability.

```
## [1] TRUE
```

```
colnames(design) <- gsub("dge\\$samples\\$tumor", "tumor", colnames(design))
colnames(design) <- gsub("dge\\$samples\\$library_typeunstranded", "library_type", colnames(design))
colnames(design) <- gsub("dge\\$samples\\$Ta2", "Ta2", colnames(design))
colnames(design) <- gsub("dge\\$samples\\$Ta3", "Ta3", colnames(design))
colnames(design) <- gsub("dge\\$samples\\$Ta4", "Ta4", colnames(design))
head(design)</pre>
```

```
##
    tumor0 tumor1 library_type Ta2 Ta3 Ta4
## 1
         1
                                     0
                0
                                 1
                                         0
## 2
                1
                             1
                                 1
                                         0
                                     0
## 3
         1
                0
                                 0
                                     0
                                         1
## 4
         0
                1
                             1
                                 0 0
                                         1
## 5
         1
                0
                             1
                                 1
                                     0
                                         0
         0
                1
                                 1
                                     0
                                         0
## 6
```

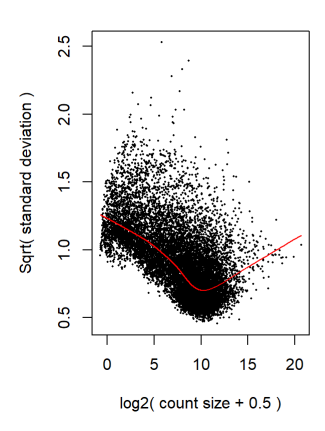
voom

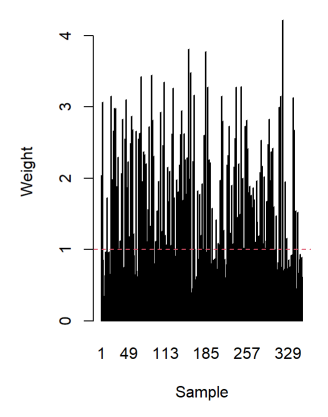
voom is a function that lies within a package called limma. limma/voom is used in DGE analysis. voom is a function that takes the counts in a metadata set and transforms them into log2 of TMM values calculated above in the normalization factors. A linear model is then fitted to the TMM for each gene and residuals are calculated. A smoothed curve is then fitted to the square root of the residual standard deviation by the average expression (this is the red line). This smooth curve is then used to obtain weights for each gene and sample that are passed into limma along with the TMM values.

```
# Running voom again with the new design matrix.
v <- voomWithQualityWeights(dge, design, plot=TRUE)</pre>
```

voom: Mean-variance trend

Sample-specific weights





limma

This sections marks the beginning of running limma. This section is dedicated to creating a linear fit to the data, making comparisons of the fitted data, and them apply Bayes smoothing. This first code chunk involves creating a variable that has all of the duplicate correlation values on v and design. These correlation values will be used later in a linear fit.

[1] 0.1595651

limma graph

This is the linear model with limma, notice that the correlation values were the duplicate correlations used earlier

```
# Fitting the linear model with limma.
# If using paired samples, the within-patient correlation and a block design
# for patient is used to account for pairwise samples
fit <- lmFit(v, design, block = v$targets$ID, correlation = corfit$consensus)</pre>
```

Coefficient vector

This code chunk involves extracting coefficients from the linear fit model and storing them in a vector for later use.

```
##
                  Contrasts
## Levels
                   Adjacent_vs_Tumor
##
     tumor0
                                    -1
##
                                    1
     tumor1
##
                                    0
     library_type
##
     Ta2
                                     0
##
     Ta3
                                     0
##
     Ta4
                                     0
```

```
# Assigning all comparisons to a vector for later allComparisons <- colnames(contrasts)
```

Contrast Analysis

This next code chunk reorients the linear model obtained earlier and obtains the coefficients and standard errors from the model. This step also sets us up to apply Empirical Bayes smoothing.

```
# Running contrast analysis
vfit <- contrasts.fit(fit, contrasts = contrasts)
# Look at N of DEGs with adj. p <0.01 and Log2FC>2
summary(decideTests(vfit, adjust.method = "BH", p.value = 0.05, lfc = 2))
```

```
## Adjacent_vs_Tumor
## Down 509
## NotSig 12717
## Up 158
```

Bayes smoothing

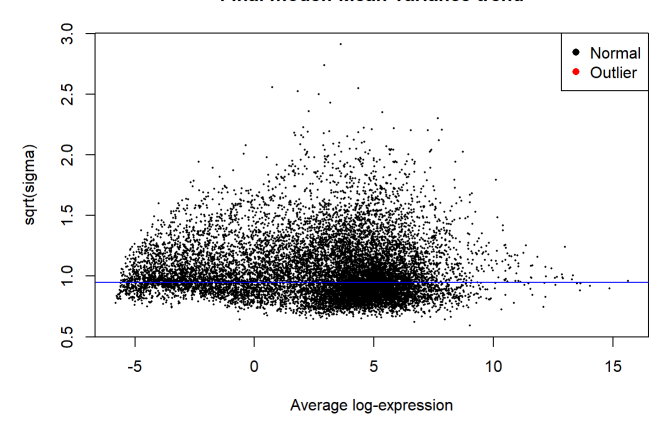
This code chunk uses Empirical Bayes smoothing to plot the final model after doing the limma and voom analysis.

Empirical Bayes smoothing is a way to account for uncertainty. The technique uses the population in a region as a measure of confidence. Meaning that areas with low margin of error are left untouched while estimates with higher margin of error are moved closer to the global average.

The write.csv is commented out because it is not needed for this file

```
# Computing differential expression based on the empirical Bayes moderation of
# the standard errors towards a common value. Robust = should the estimation of
# the empirical Bayes prior parameters be robustified against outlier sample
# variances?
veBayesFit <- eBayes(vfit, robust=TRUE)
plotSA(veBayesFit, main = "Final model: Mean-variance trend")</pre>
```

Final model: Mean-variance trend



```
vTopTable <- topTable(veBayesFit, n=Inf, p.value=1, lfc=0)
DEGs <- topTable(veBayesFit, n=Inf, p.value=0.05, lfc=2)
#DEGs_print <- data.frame(DEGs$GENEID, DEGs$gene_name, DEGs$adj.P.Val, DEGs$logFC)
#write.csv(DEGs_print, "~/R/gene_list_tumor_vs_tumor_adjacent.csv")</pre>
```

DGE Analysis Female tumor vs. tumor- adjacent

This section is doing voom/limma and Go/KEGG analysis on all of the Female tumor vs. tumor adajcent samples

Design matrix

This code chunk is setting up a second design matrix for a voom analysis of tumor vs. non-tumor differentiated by sex.

[1] TRUE

```
#Added target and library type column names
colnames(design) <- gsub("v\\$targets\\$gender_tissue", "", colnames(design))
colnames(design) <- gsub("v\\$targets\\$library_typeunstranded", "library_type", colnames(design))
colnames(design) <- gsub("v\\$targets\\$Ta2", "Ta2", colnames(design))
colnames(design) <- gsub("v\\$targets\\$Ta3", "Ta3", colnames(design))
colnames(design) <- gsub("v\\$targets\\$Ta4", "Ta4", colnames(design))
head(design)</pre>
```

```
F_0 F_1 M_0 M_1 library_type Ta2 Ta3 Ta4
##
## 1 0 0 1 0
                         1
                            1
                               0
## 2 0 0 0 1
## 3 0 0 1 0
     0 0 0 1
                         1
                            1
## 4 0 0 0 1
                       1 0 0 1
                      1 1 0 0
1 1 0 0
     0 0 1 0
## 5
## 6
```

head(design)

```
##
   F_0 F_1 M_0 M_1 library_type Ta2 Ta3 Ta4
## 1 0
       0
             0
                      1
                         1
                           0
          1
## 2
   0 0 0 1
                      1
                         1
                           0
                             0
    0 0 1 0
                     1 0
## 3
                           0 1
## 4
    0 0 0 1
                          0 1
    0 0 1 0
                    1 1
                           0 0
## 5
## 6
```

head(design)

```
##
    F_0 F_1 M_0 M_1 library_type Ta2 Ta3 Ta4
      0
             1
## 1
         0
## 2
      0
         0
             0
                1
                            1
                               1
                                   0
                                      0
## 3
      0 0 1
                0
                            1
                               0
                                   0
                                      1
      0 0 0 1
                            1
## 4
                               0
                                   0
                                     1
## 5
         0 1
                            1 1
                0
                                   0
## 6
             0
                1
                                   0
```

head(design)

```
F_0 F_1 M_0 M_1 library_type Ta2 Ta3 Ta4
##
## 1
             1
                0
                            1
                               1
                                   0
                                      0
      0
         0
                1
                           1
                               1
## 2
      0
                                   0
                                      0
## 3
      0 0 1 0
                           1
                               0
                                  0
                                      1
      0 0 0 1
                           1
                                     1
## 4
                               0
                                  0
## 5
      0
         0 1
                0
                           1
                               1
                                  0
                                      0
## 6
         0
                1
                           1
                                  0
```

head(design)

```
##
    F_0 F_1 M_0 M_1 library_type Ta2 Ta3 Ta4
                           1
                               1
## 1
     0
         0
            1
                0
                                  0
                                     0
         0
            0
               1
                           1
                              1
                                     0
## 2
      0
                                  0
## 3
     0 0 1
                0
                           1
                              0
                                  0
                                     1
## 4
     0 0 0 1
                           1
                              0
                                  0 1
## 5
         0 1
                           1
      0
                0
                              1
                                  0
                                     0
            0
                           1
## 6
      0
                1
                              1
                                  0
                                     0
```

Voom for Female Tumor vs. Tumor adjacent

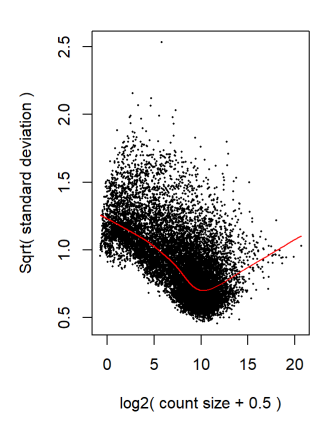
This code chunk is using the voom function and outputting the graph.

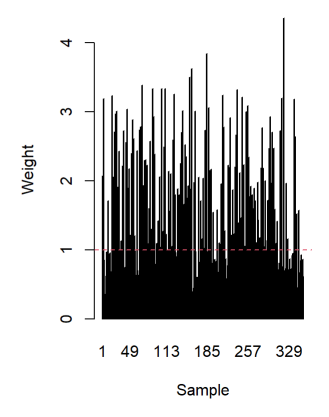
```
# Running voom with quality weights. Normalizes expression intensities so that
# the log-ratios have similar distributions across a set of samples.
# To quantile normalize, add normalize.method="quantile"
# Running parallel

v <- voomWithQualityWeights(dge, design, plot=TRUE)
```

voom: Mean-variance trend

Sample-specific weights





limma

This code chunk is for doing paired samples correlation tests. The value of the correlation should be positive.

```
# Block design for individual. This is used in tumor-normal comparisons with
# paired samples.
corfit <- duplicateCorrelation(v, design, block = v$targets$ID)
# This should give a positive correlation value. It represents the
# correlation between measurements made on the same person.
corfit$consensus</pre>
```

[1] 0.158629

limma graph This code chuck is doing a linear fit model with limma for female tumor vs tumor-adjacent.

```
# Fitting the linear model with limma.
# If using paired samples, the within-patient correlation and a block design
# for patient is used to account for pairwise samples
fit <- lmFit(v, design, block = v$targets$ID, correlation = corfit$consensus)</pre>
```

Coefficient vector

This code chunk does pairwise comparisons of the female tumor vs. tumor adjacent sample for contrast design for differential expressions and stores the comparisons into a vector for later.

```
##
                  Contrasts
## Levels
                   Adjacent_M_vs_Tumor_M Adjacent_F_vs_Tumor_F
     F_0
##
                                        0
                                                               -1
##
    F 1
                                        0
                                                                1
##
    M_0
                                       -1
                                                                0
##
     M 1
                                                                0
                                        1
##
     library_type
                                                                0
                                        0
                                        0
##
     Ta2
```

```
# Assigning all comparisons to a vector for later allComparisons <- colnames(contrasts)
```

Contrast Analysis

This code is running a contrast analysis using the data from the limma model.

```
# Running contrast analysis
vfit <- contrasts.fit(fit, contrasts = contrasts)
# Looking at N of DEGs with adj. p <0.01 and Log2FC>2
summary(decideTests(vfit, adjust.method = "BH", p.value = 0.01, lfc = 2))
```

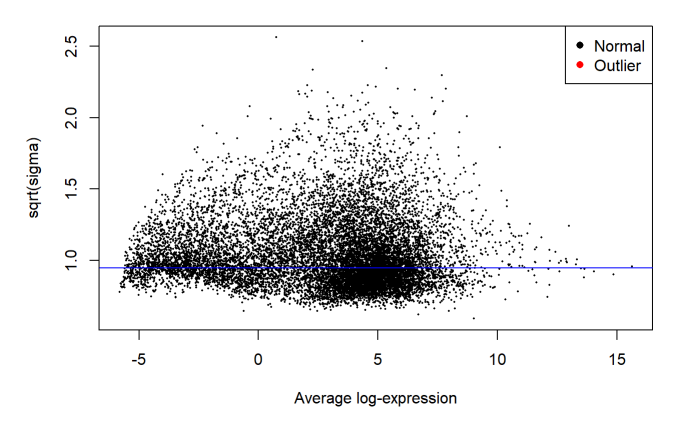
```
## Adjacent_M_vs_Tumor_M Adjacent_F_vs_Tumor_F
## Down 542 430
## NotSig 12678 12797
## Up 164 157
```

Bayes smoothing

Bayes smoothing on data from the limma model. Empirical Bayes smoothing increases the power of the limma model.

```
# Computing differential expression based on the empirical Bayes moderation of
# the standard errors towards a common value. Robust = should the estimation of
# the empirical Bayes prior parameters be robustified against outlier sample
# variances?
veBayesFit <- eBayes(vfit, robust=TRUE)
plotSA(veBayesFit, main = "Final model: Mean-variance trend")</pre>
```

Final model: Mean-variance trend



```
vTopTable_M <- topTable(veBayesFit, coef=1, n=Inf, p.value=1, lfc=0)
vTopTable_F <- topTable(veBayesFit, coef=2, n=Inf, p.value=1, lfc=0)

DEGs_M <- topTable(veBayesFit, coef=1, n=Inf, p.value=0.01, lfc=2)
DEGs_F <- topTable(veBayesFit, coef=2, n=Inf, p.value=0.01, lfc=2)
DEGs_F_relax_p <- topTable(veBayesFit, coef=2, n=Inf, p.value=0.1, lfc=2)</pre>
```

Data Visualization

This section creates Volcano plots for easy visualization of the gene lists.

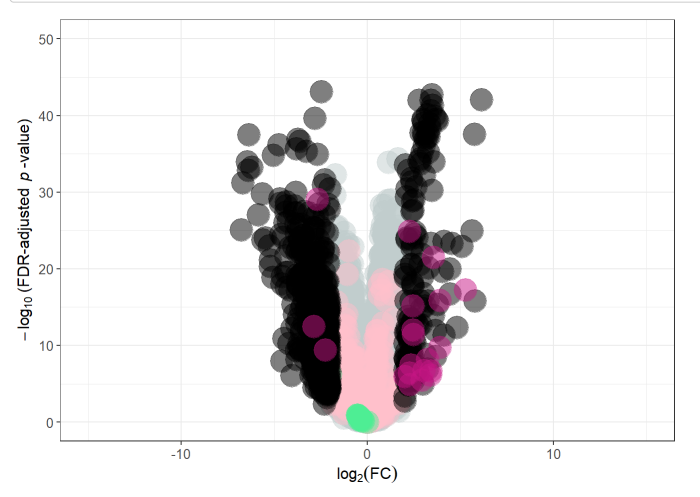
Volcano Plot Object This code chunk is developing the plot object for a volcano plot of female tumor vs. tumor adjacent samples.

```
#Volcano plot of female tumor vs tumor-adjacent
df <- data.frame(vTopTable_F$adj.P.Val, vTopTable_F$logFC, vTopTable_F$chr, vTopTable_F$GENEID,</pre>
 vTopTable F$gene name)
colnames(df) <- c("adj.P.Val", "logFC", "chr", "id", "name")</pre>
dfSig \leftarrow df[(abs(df$logFC) >= 2 \& df$adj.P.Val \leftarrow 0.01),]$id
dfAnons <- subset(df, chr != "chrX" & chr != "chrY" & !(id %in% dfSig))
dfAnons <- cbind(dfAnons , rep(1, nrow(dfAnons)))</pre>
colnames(dfAnons)[6] <- "Color"</pre>
dfXnons <- subset(df, chr == "chrX" & !(id %in% dfSig))</pre>
dfXnons <- cbind(dfXnons, rep(2, nrow(dfXnons)))</pre>
colnames(dfXnons)[6] <- "Color"</pre>
dfYnons <- subset(df, chr == "chrY" & !(id %in% dfSig))</pre>
dfYnons <- cbind(dfYnons, rep(3, nrow(dfYnons)))</pre>
colnames(dfYnons)[6] <- "Color"</pre>
dfA <- subset(df, chr != "chrX" & chr != "chrY" & id %in% dfSig)</pre>
dfA <- cbind(dfA, rep(4, nrow(dfA)))</pre>
colnames(dfA)[6] <- "Color"</pre>
dfX <- subset(df, chr == "chrX" & id %in% dfSig)</pre>
dfX <- cbind(dfX, rep(5, nrow(dfX)))</pre>
colnames(dfX)[6] <- "Color"</pre>
dfY <- subset(df, chr == "chrY" & id %in% dfSig)</pre>
dfY <- cbind(dfY, rep(6, nrow(dfY)))</pre>
colnames(dfY)[6] <- "Color"</pre>
dfPlot <- rbind(dfAnons, dfXnons, dfYnons, dfA, dfX, dfY)
dfPlot$Color <- as.factor(dfPlot$Color)</pre>
```

Volcano Plot

This code chunk plots the female tumor vs. tumor adjacent volcano plot.

```
p <- ggplot(data = dfPlot, aes(x = logFC, y = -log10(adj.P.Val), color=Color )) +
    geom_point(alpha = 0.5, size = 8) +
    theme_bw() +
    theme(legend.position = "none") +
    xlim(c(-15, 15)) + ylim(c(0, 50)) +
    scale_color_manual(values = c("azure3", "pink", "seagreen2", "black", "mediumvioletred", "spri
nggreen")) +
    labs(x=expression(log[2](FC)),
        y=expression(-log[10] ~ "(FDR-adjusted " ~ italic("p") ~ "-value)")) +
    theme(axis.title.x=element_text(size=12),
        axis.text.x=element_text(size=10)) +
    theme(axis.title.y=element_text(size=12),
        axis.text.y=element_text(size=10))
p</pre>
```



```
forLabel <- subset(dfPlot, adj.P.Val<=0.01 & abs(logFC)>=2)
```

Significance Thresholds This code chunk is setting the significance thresholds for the volcano plot. The thresholds can be seen via dotted lines on the graph.

```
# Adding lines for significance thresholds
pdf("~/R/Volcano_plot_female_DEGs.pdf", width=12, height=12)
p + geom_hline(yintercept = 2, colour="#000000", linetype="dashed"
) + geom_vline(xintercept = 2, colour="#000000", linetype="dashed"
) + geom_vline(xintercept = -2, colour="#000000", linetype="dashed"
) + geom_text_repel(data=forLabel, max.iter=1000, box.padding = 0.25, force=1, aes(x = logFC, y = -log10(adj.P.Val), label=name, color=Color), size=8)
```

```
## Warning: ggrepel: 551 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
```

```
#dev.off()
```

Venn Diagram

In this code chunk two Venn Diagrams are made. The commented out sections are the original code sections. The first two sections are creating the Venn Diagrams. Given that I have a different working directory and am doing this is an RMD file not an R file. I rewrote the code to create the Venn Diagrams and output them into the RMD file which you see below.

The third section is inputting data from a CSV file. I do not have the said CSV files so I have commented these sections out. These samples show the overlap between male and female samples.

```
library(VennDiagram)
library(grDevices)
#venn.diagram(List("Female"=DEGs_F$gene_name, "Male"=DEGs_M$gene_name), filename = "~/3.0 Hasti
ng Research/Liver Cancer Analysis/DEG Figures/Overlap_DEGs.png")

#venn.diagram(List("Female"=DEGs_F_relax_p$gene_name, "Male"=DEGs_M$gene_name), filename = "~/
3.0 Hasting Research/Liver Cancer Analysis/DEG Figures/Overlap_DEGs_Frelaxp.png")

#write.csv(DEGs_F, "~/3.0 Hasting Research/Liver Cancer Analysis/DEG Figures/Female_all_DEGs.cs
v")

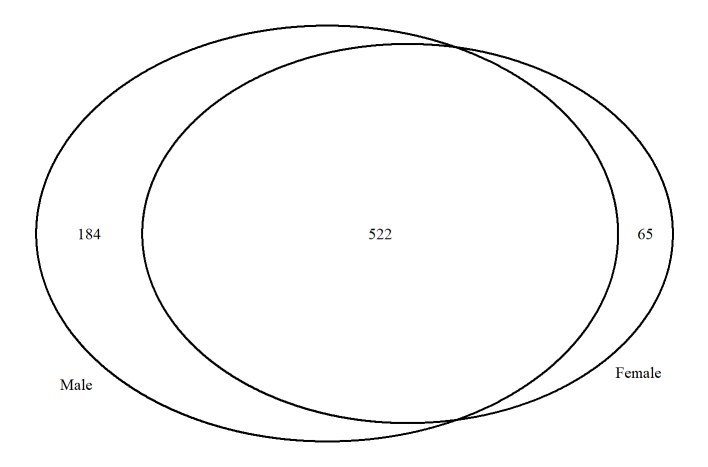
#write.csv(DEGs_M, "~/3.0 Hasting Research/Liver Cancer Analysis/DEG Figures/Male_all_DEGs.csv")

write.csv(DEGs_F, "~/R/Female_DEGs.csv")

write.csv(DEGs_F, "~/R/Male_all_DEGs.csv")

venn1<- venn.diagram(List("Female"=DEGs_F$gene_name, "Male"=DEGs_M$gene_name),filename = NULL)
grid.newpage()

grid.draw(venn1)</pre>
```



```
pdf(file="venn1.pdf")

venn2<-venn.diagram(List("Female"=DEGs_F_relax_p$gene_name, "Male"=DEGs_M$gene_name), filename
= NULL)
grid.newpage()
grid.draw(venn2)
pdf(file="venn2.pdf")
dev.off()</pre>
```

```
## png
## 2
```

Results

This sections shows the results of the DEG and Go/KEGG analysis

DEG results

This code chunk shows the results of our DEG analysis by retrieving specific data from a package called biomart. The combination of the DEG analysis results and the data from biomart is printed into a table.

The label "kegg-enzyme' was not valid so it has been deleted.

Ensembl site unresponsive, trying useast mirror

```
#biomart <- getBM(attributes=c('ensembl_gene_id', 'hgnc_symbol', 'entrezgene_id', 'kegg_enzym
e'), filters = 'hgnc_symbol', values = degResult_genes$gene_name, mart = ensembl)

#Deleted "kegg_enzyme" label beccause it does not work.
biomart <- getBM(attributes=c('ensembl_gene_id', 'hgnc_symbol', 'entrezgene_id'), filters = 'hgn
c_symbol', values = degResult_genes$gene_name, mart = ensembl)
colnames(degResult_genes) <- c("chr", "start", "end", "TXNAME", "GENEID", "hgnc_symbol", "lengt
h", "logFC", "AveExpr", "t", "P.Value", "adj.P.Val", "B")</pre>
```

GO/KEGG Analysis Male tumor vs. tumor-adjacent

```
degResult <- DEGs_F
degResult$hgnc_symbol <- degResult$gene_name
degResult_biomart <- merge(degResult, biomart, by="hgnc_symbol")
degResult_biomart <- degResult_biomart[complete.cases(degResult_biomart), ]
geneList <- degResult_biomart$entrezgene_id
geneList_quant <- degResult_biomart$logFC
ego <- enrichGO(geneList, OrgDb = "org.Hs.eg.db", ont="BP", readable=TRUE)
pdf("~/R/Goplot_all_female.pdf", width=12, height=12)
goplot(ego)
#dev.off()
pdf("~/R/dotplot_all_female.pdf", width=12, height=12)
dotplot(ego, showCategory=30)
#dev.off()</pre>
```