# Examine the association between gene expression and age\_group variable (fetus vs adult)

### Vy K Phung

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### Introduction

The purpose of this re-analysis is to examine the correlation of differential gene expression between fetal and adult brains, which is evaluated through RNA-sequencing. If it has correlation, then count how many up-regulated and down-regulated genes. I will do exploratory analysis and statistical analysis by using R, RStudio.

### Load library

```
library(tidyverse)
library(limma)
library(preprocessCore)
library(RColorBrewer)
library(org.Hs.eg.db)
library(AnnotationDbi)
library(edge)
library(sva)
library(DESeq2)
library(broom)
library(readxl)
```

### Data preprocessing

```
count <- read.delim("D:/word/bioinformatics/personal project/PRJNA245228/tidy data/count.csv")
pheno <- read.csv("D:/word/bioinformatics/personal project/PRJNA245228/sample_data/pheno_sample.csv")
head(count)</pre>
```

##		ENTREZID		ENSE	MBL	SYMBOL	SRR1554534	SRR1554535	SRR1554568	SRR1554561
##	1	1	EN	SG00000121	410	A1BG	444	378	328	650
##	2	10	EN	SG00000156	006	NAT2	11	8	4	0
##	3	100	EN	SG00000196	839	ADA	299	658	161	229
##	4	1000	EN	SG00000170	558	CDH2	7384	10623	43926	11016
##	5	10000	EN	SG00000117	020	AKT3	6837	7391	90930	13677
##	6	10000	EN	SG00000275	199	AKT3	6837	7391	90930	13677
##		SRR155456	37	SRR1554536	SRI	R1554541	SRR1554539	SRR1554538	SRR1554537	7
##	1	14	16	114		592	295	275	5 518	3
##	2		8	0		8	3	3 12	2	1
##	3	22	25	291		382	265	354	160	)
##	4	5274	16	3270		44244	10639	47354	52346	3
##	5	3624	16	937		74768	18216	48565	7968	5
##	6	3624	16	937		74768	18216	48565	7968	5

### head(pheno)

```
## Run age_group age sex
## 1 SRR1554534 adult 40.4200 male
## 2 SRR1554535 adult 41.5800 male
## 3 SRR1554568 fetus -0.4986 male
## 4 SRR1554561 adult 43.8800 male
## 5 SRR1554567 fetus -0.4027 male
## 6 SRR1554536 adult 44.1700 female
```

dup <- duplicated(count\$SYMBOL)</pre>

head(count\_symbol)

It is clear that there are some duplications in columns "SYMBOL"(gene symbol) and "ENTREZID", for example, in line 5,6 of "count" table. I will remove duplicated genes and use gene symbol as row name.

```
## dup
## FALSE TRUE
## 23741 4955

count_symbol<- count[!dup,-1:-2]
na <- is.na(count_symbol$SYMBOL)
count_symbol <- count_symbol[!na,]
row.names(count_symbol) <- count_symbol$SYMBOL
count_symbol <- count_symbol (-1]</pre>
```

##		SRR1554534	SRR1554535	SRR1554568	SRR1554561	SRR1554567	SRR1554536
##	A1BG	444	378	328	650	146	114
##	NAT2	11	8	4	0	8	0
##	ΔΠΔ	299	658	161	229	225	291

##	CDH2	7384	10623	43926	11016	52746	3270
##	AKT3	6837	7391	90930	13677	36246	937
##	GAGE12F	0	0	0	0	0	0
##		SRR1554541	SRR1554539	SRR1554538	SRR1554537		
##	A1BG	592	295	275	518		
##	NAT2	8	8	12	4		
##	ADA	382	265	354	160		
##	CDH2	44244	10639	47354	52346		
##	AKT3	74768	18216	48565	79685		
##	GAGE12F	0	0	0	0		

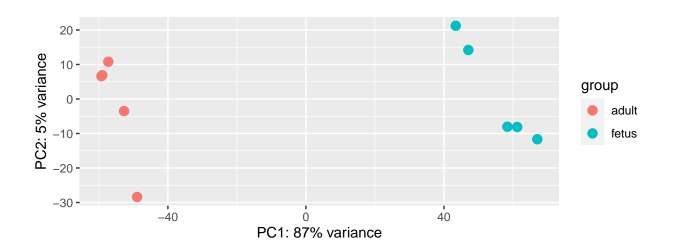
### Data exploration

Although the target of this exploratory analysis is figuring out if there is a correlation between differential gene expression in fetus vs adult brains, I still plot PCA for the sex group in this data exploration to have a brief overview if there might be an association between sex variable (female vs male) with gene expression in fetus vs adult.

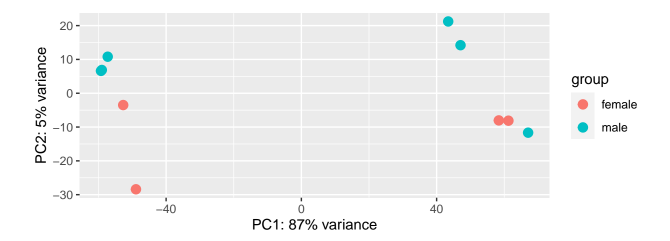
At first, I will explore unfiltered data which still has some genes' row names having 0 reads in "count\_symbol" table by using DESeq2.

### Unfiltered data

```
library(DESeq2)
edata <- DESeqDataSetFromMatrix(countData = count_symbol, colData = pheno, design = ~ age_group)
edata_tr <- rlog(edata, blind = FALSE)
plotPCA(edata_tr, intgroup = c("age_group"))</pre>
```



plotPCA(edata\_tr, intgroup = c("sex"))



According to plotPCA having intgroup "age\_group", there can be an association between differential gene expression and age\_group(fetus vs adult). However, in plotPCA having intgroup "sex", we can see that there might be no association between gender and gene expression.

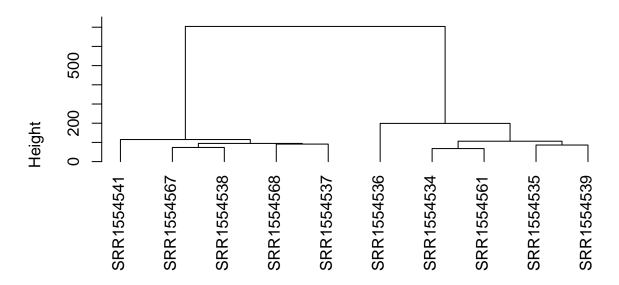
I will also show the table of data transform of the above count\_symbol table and the cluster of it in Dendogram so that we could easily visualize the correlation between those samples.

```
edata_tr <- assay(edata_tr)
head(edata_tr)</pre>
```

```
##
           SRR1554534 SRR1554535 SRR1554568 SRR1554561 SRR1554567 SRR1554536
## A1BG
             9.190681
                         8.591025
                                     8.087121
                                                9.266152
                                                            6.808106
                                                                        8.754521
## NAT2
             2.898727
                         2.564131
                                     2.169268
                                                1.840736
                                                            2.306952
                                                                        1.945286
                         9.225800
##
  ADA
             8.671241
                                     7.208636
                                                7.967730
                                                            7.255809
                                                                        9.862274
   CDH2
            13.536261
                        13.571832
                                    14.972885
                                               13.633542
                                                           14.836363
                                                                       13.764138
##
  AKT3
                        13.144329
                                    15.892417
                                                13.906920
                                                           14.381850
##
            13.453204
                                                                       12.281291
## GAGE12F
             0.000000
                         0.00000
                                     0.00000
                                                0.00000
                                                            0.000000
                                                                        0.00000
##
           SRR1554541 SRR1554539 SRR1554538 SRR1554537
## A1BG
             8.388780
                         8.418109
                                     7.553544
                                                8.518947
## NAT2
             2.289385
                         2.615617
                                     2.494186
                                                2.140643
## ADA
             7.824756
                         8.248514
                                     7.806678
                                                7.090937
  CDH2
                        13.699760
                                    14.722689
                                                15.072426
##
            14.563325
##
  AKT3
            15.224047
                        14.367595
                                    14.763178
                                               15.606567
## GAGE12F
             0.000000
                         0.000000
                                     0.00000
                                                0.000000
```

```
dist_samples <- dist(t(edata_tr))
gene_fit <- hclust(dist_samples, method="ward.D")
plot(gene_fit, hang=-1)</pre>
```

### **Cluster Dendrogram**



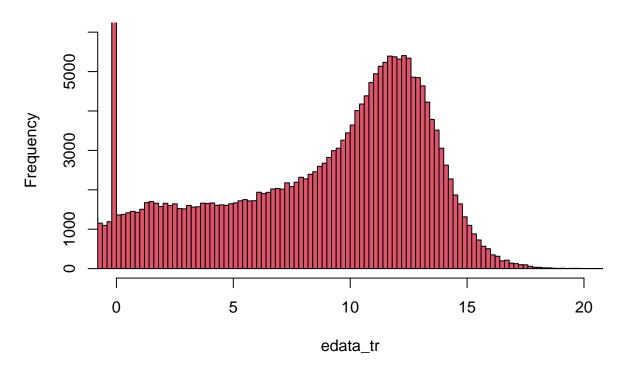
# dist\_samples hclust (\*, "ward.D")

According to cluster Dendogram, there are 2 main branches. The smaller branches SRR15545(41,67,38,68,37) on the main left are fetus group, while the others on the main right are in adult group. This visualization reinforces the hypothesis that there can be a correlation between differential gene expression in fetus vs adult.

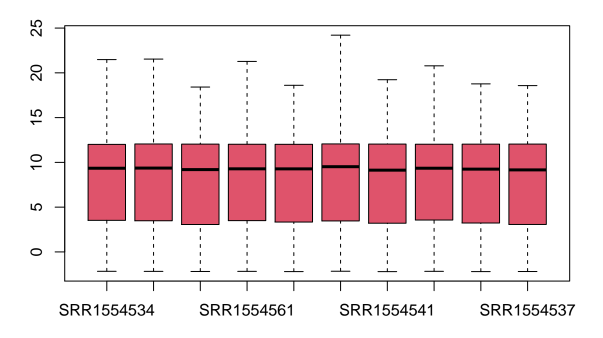
Additionally, I also visualize the frequency of number of reads in the count table when data has been transformed on histogram and boxplot. We can see that mostly the reads are below 20 and in the range between 5 to 15.

```
hist(edata_tr,breaks=100,col=2,xlim=c(0,20),ylim=c(0,6000))
```

# Histogram of edata\_tr



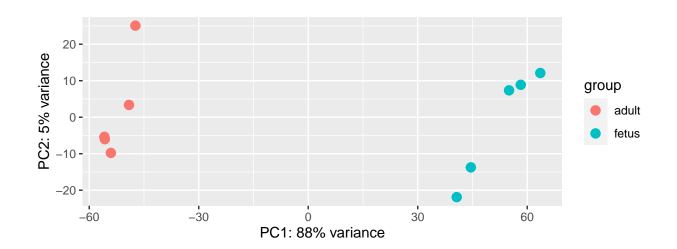
boxplot(edata\_tr,col=2,range=0)



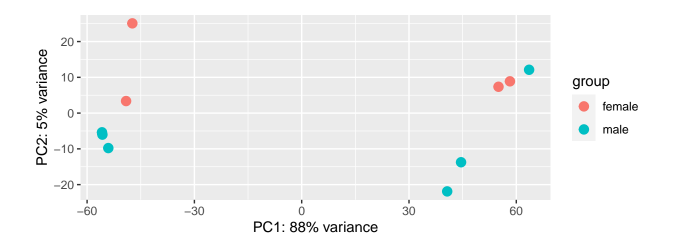
### Filtered data

Removing lowly expressed genes and using DESeq to explore data as the same way as above unfiltered data in order to see if there are any differences between filtered and unfiltered one.

```
count_filter = count_symbol[rowMeans(count_symbol) > 100,]
edata <- DESeqDataSetFromMatrix(countData = count_filter, colData = pheno, design = ~ age_group)
edata_tr <- rlog(edata, blind = FALSE)
plotPCA(edata_tr, intgroup = c("age_group"))</pre>
```



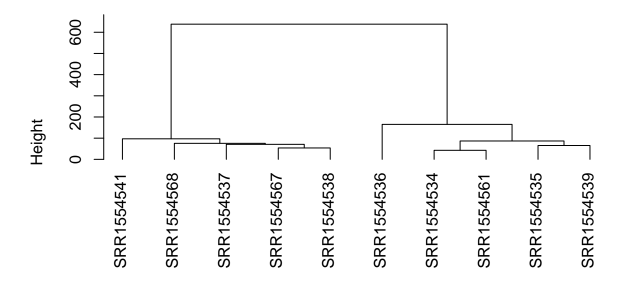
plotPCA(edata\_tr, intgroup = c("sex"))



```
edata_tr <- assay(edata_tr)
head(edata_tr)
```

```
##
              SRR1554534 SRR1554535 SRR1554568 SRR1554561 SRR1554567 SRR1554536
## A1BG
                9.143633
                            8.583227
                                        8.105891
                                                    9.206773
                                                               6.995598
                                                                           8.762864
                                                                7.351627
## ADA
                8.646385
                            9.145974
                                        7.303994
                                                    8.001912
                                                                           9.770115
                                                              14.829826
##
  CDH2
                13.562468
                           13.585906
                                       14.957846
                                                   13.652418
                                                                          13.812075
## AKT3
                13.479431
                           13.159273
                                       15.876052
                                                   13.925121
                                                              14.376621
                                                                          12.332018
## ZBTB11-AS1
                7.746129
                            7.539217
                                        7.489181
                                                    7.323988
                                                               7.706548
                                                                           7.731734
## MED6
                10.965760
                           10.953407
                                       11.062971
                                                   10.825176
                                                              11.161363
                                                                          10.689913
##
              SRR1554541 SRR1554539 SRR1554538 SRR1554537
                                        7.632873
## A1BG
                8.373665
                            8.429514
                                                    8.489938
## ADA
                7.843839
                            8.253862
                                        7.831596
                                                    7.197164
##
  CDH2
                14.545283
                           13.717079
                                       14.708753
                                                   15.051336
                15.204848
                           14.383520
                                       14.749481
## AKT3
                                                   15.584859
## ZBTB11-AS1
                7.684944
                            7.654166
                                        7.625887
                                                    7.705363
## MED6
                11.229573
                           11.095095
                                       11.270915
                                                  11.087431
dist_samples <- dist(t(edata_tr))</pre>
gene_fit <- hclust(dist_samples, method="ward.D")</pre>
plot(gene_fit, hang=-1)
```

## **Cluster Dendrogram**

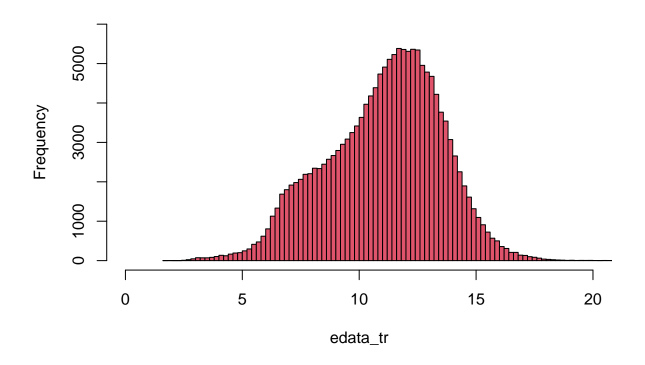


dist\_samples hclust (\*, "ward.D")

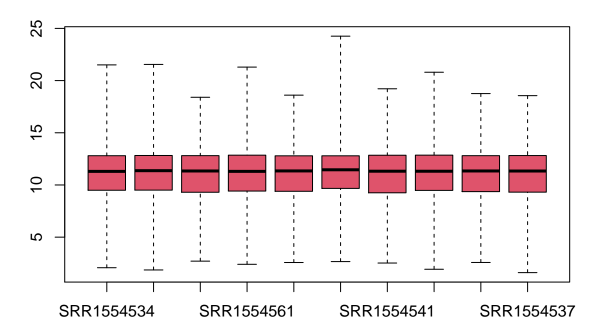
We can see that there is almost no difference in the results of unfiltered and filtered data except on the cluster Dendogram of filtered one, the relation between SRR1554568 and SRR1554537 are not close-related as same as that of unfiltered.

hist(edata\_tr,breaks=100,col=2,xlim=c(0,20),ylim=c(0,6000))

# Histogram of edata\_tr



boxplot(edata\_tr,col=2,range=0)



### Stratified analysis

## FALSE TRUE

6

##

Because I want to explore if the sex variable (male vs female) might effect the association between age\_group variable and gene expression, I get the female data in 10 samples above and analyse the factor age\_group in this gender and do the same with the male gender.

**Female** There are 4/10 samples having sex is female

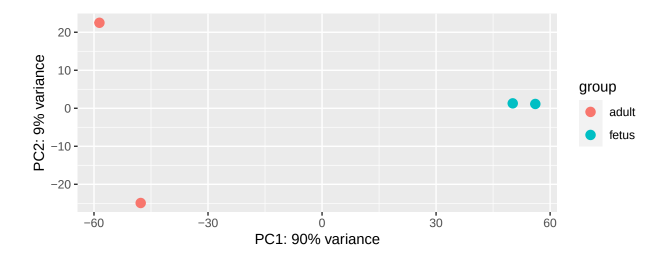
```
pheno_female = pheno[pheno$sex == "female",]
pheno_female
##
             Run age_group
                                 age
                                        sex
      SRR1554536
## 6
                      adult 44.1700 female
## 8
      SRR1554539
                      adult 36.5000 female
      SRR1554538
                      fetus -0.4027 female
## 10 SRR1554537
                      fetus -0.3836 female
female_run <- (colnames(count_symbol) %in% pheno_female$Run)</pre>
table(female_run)
## female_run
```

```
count_female = count_symbol[,female_run]
head(count_female)
```

```
##
           SRR1554536 SRR1554539 SRR1554538 SRR1554537
## A1BG
                              295
                  114
                                         275
                                                    518
## NAT2
                    0
                                8
                                          12
                  291
## ADA
                              265
                                         354
                                                     160
## CDH2
                 3270
                            10639
                                       47354
                                                   52346
## AKT3
                  937
                            18216
                                       48565
                                                   79685
## GAGE12F
                    0
                                                       0
```

edata\_fe <- DESeqDataSetFromMatrix(count\_female, pheno\_female, ~age\_group)</pre>

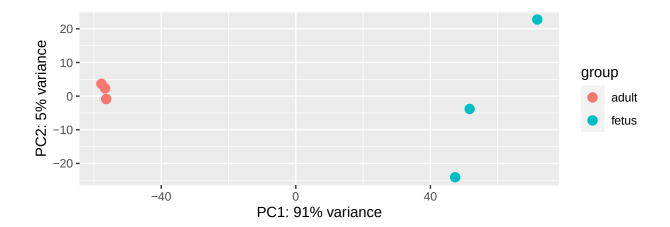
```
edata_fe_tr <- rlog(edata_fe, blind = FALSE)
plotPCA(edata_fe_tr, intgroup = c("age_group"))</pre>
```



```
pheno_male = pheno[pheno$sex == "male",]
pheno_male
```

Male

```
Run age_group
                           age sex
## 1 SRR1554534 adult 40.4200 male
## 2 SRR1554535
                    adult 41.5800 male
## 3 SRR1554568
                    fetus -0.4986 male
## 4 SRR1554561
                    adult 43.8800 male
## 5 SRR1554567
                    fetus -0.4027 male
## 7 SRR1554541
                    fetus -0.3836 male
male_run <- (colnames(count_symbol) %in% pheno_male$Run)</pre>
table(male_run)
## male_run
## FALSE TRUE
       4
count_male = count_symbol[,male_run]
head(count_male)
           SRR1554534 SRR1554535 SRR1554568 SRR1554561 SRR1554567 SRR1554541
##
## A1BG
                  444
                             378
                                        328
                                                   650
                                                               146
                                                                          592
## NAT2
                   11
                               8
                                          4
                                                    0
                                                                8
                                                                            8
## ADA
                  299
                             658
                                        161
                                                   229
                                                               225
                                                                          382
## CDH2
                 7384
                           10623
                                      43926
                                                  11016
                                                             52746
                                                                        44244
## AKT3
                 6837
                            7391
                                      90930
                                                  13677
                                                             36246
                                                                        74768
## GAGE12F
                    0
                               0
                                          0
                                                      0
                                                                            0
                                                                 0
edata_ma <- DESeqDataSetFromMatrix(count_male, pheno_male, ~age_group)</pre>
edata_ma_tr <- rlog(edata_ma, blind = FALSE)</pre>
plotPCA(edata_ma_tr, intgroup = c("age_group"))
```



For both female and male, the visualization of plotPCA shows that there still can be an association between gene expression with age\_group variable(fetus vs adult).

### Statistical analysis

The target of this statistical analysis is examining the correlation between age\_group variable (fetus or adult) and gene expression more clearly so that in the end I will count up and down regulated genes.

In this statistical analysis, I will use limma package and DESeq package to see if there are any different results between two methods.

I will use DEseq package to analyze edata (output of DESeq) and Limma package to analyze edata\_tr (edata has been transformed)

#### Unadjusted data

At first, I will analyze statistically variable age\_group without adjustment factor (sex variable).

```
# age_group
mod_age = model.matrix(~ pheno$age_group)
fit_limma_age = lmFit(edata_tr,mod_age)
ebayes_limma_age = eBayes(fit_limma_age)
re = topTable(ebayes_limma_age, number=dim(count_symbol)[1])
head(re)
```

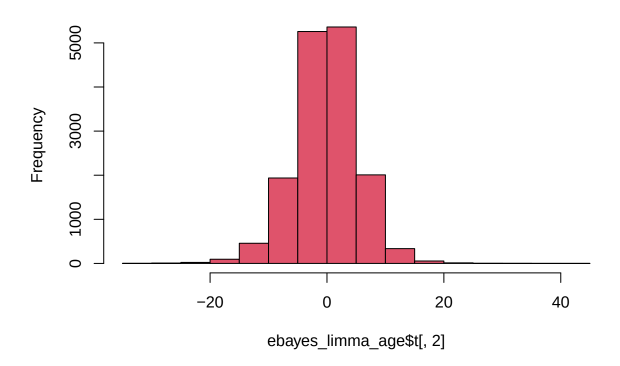
### Fit regression with limma package

```
##
              logFC
                      AveExpr
                                             P.Value
                                                        adj.P.Val
## ST8SIA2 6.264017 11.767199 40.30384 2.684402e-13 2.770718e-09 20.15611
## SOX11
            7.196578 13.549118 39.27425 3.561563e-13 2.770718e-09 19.94839
## TRIM54 -6.448248 6.397455 -34.32829 1.547426e-12 8.025467e-09 18.81527
                               33.37950 2.100303e-12 8.169655e-09 18.56884
            5.755159 12.718546
## SLA
## FBN3
            4.918986 11.616488 30.36669 5.882270e-12 1.382686e-08 17.71288
## SNCG
           -6.707473 10.307404 -30.34373 5.930876e-12 1.382686e-08 17.70589
```

#### # Statistics

hist(ebayes\_limma\_age\$t[,2], col=2)

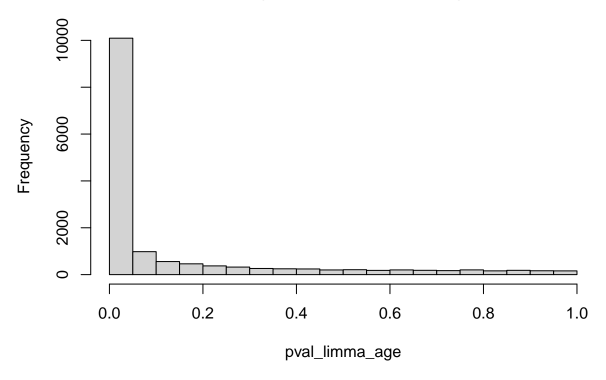
## Histogram of ebayes\_limma\_age\$t[, 2]



### # P-values

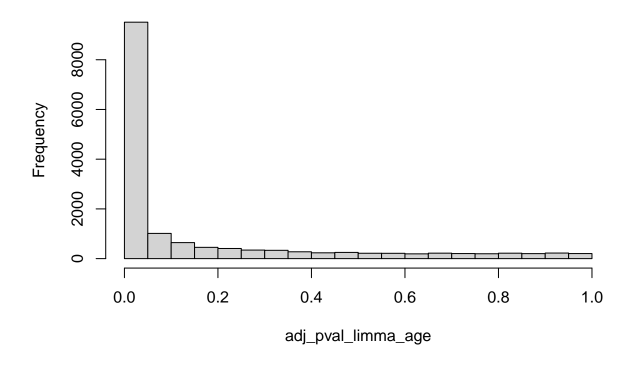
pval\_limma\_age = topTable(ebayes\_limma\_age, number=dim(edata\_tr)[1])\$P.Value
hist(pval\_limma\_age)

# Histogram of pval\_limma\_age



# # Adjusted p-values adj\_pval\_limma\_age = topTable(ebayes\_limma\_age,number=dim(edata\_tr)[1])\$adj.P.Val hist(adj\_pval\_limma\_age)

## Histogram of adj\_pval\_limma\_age



According to the histogram of adjusted p-value, it suggests that there might be an association between age\_group variable and gene expression, which means there is a differential gene expression between fetal and adult brains in these samples.

A number of genes have adjusted p-value less than 0.05

```
sum(re$adj.P.Val < 0.05)</pre>
```

## [1] 9511

```
dds <- DESeq(edata)
```

```
res <- results(dds)
res = as.data.frame(res)
head(res)</pre>
```

#### Fit regression with DESeq

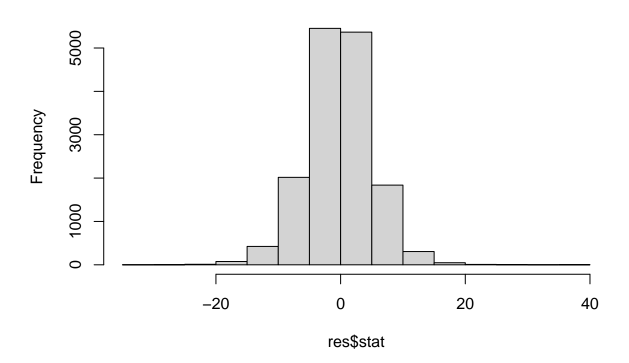
```
## baseMean log2FoldChange lfcSE stat pvalue
## A1BG 380.8469 -1.10293610 0.4076920 -2.7053167 6.823930e-03
## ADA 372.3584 -1.86297881 0.4452617 -4.1840088 2.864130e-05
```

```
## CDH2
              21681.2984
                             1.36767686 0.1515741 9.0231574 1.827447e-19
## AKT3
              27928.4725
                             1.91781744 0.4513954 4.2486416 2.150707e-05
## ZBTB11-AS1
                198.2623
                             0.06499846 0.2042175 0.3182805 7.502722e-01
## MED6
               2117.0608
                             0.29961394 0.1355707 2.2100194 2.710382e-02
                      padj
## A1BG
              1.190664e-02
## ADA
              7.382847e-05
## CDH2
              2.284087e-18
## AKT3
              5.653192e-05
## ZBTB11-AS1 7.941369e-01
## MED6
              4.215816e-02
```

### # Statistic

hist(res\$stat)

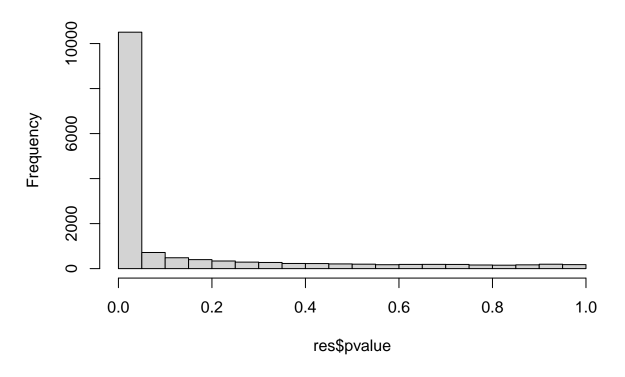
## Histogram of res\$stat



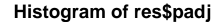
### # P-values

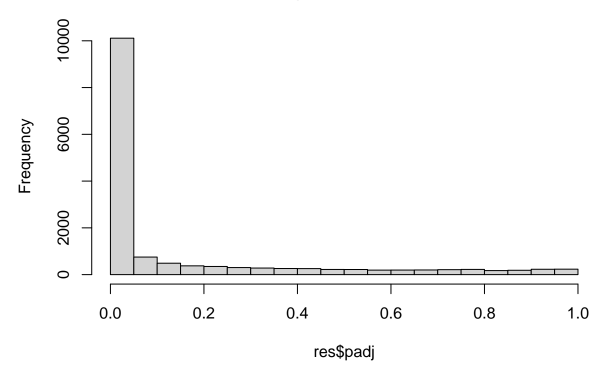
hist(res\$pvalue)

# Histogram of res\$pvalue



#Adjusted p-values hist(res\$padj)





We can see that there is also an association between gene expression with age\_group as same as the above result of limma package.

A number of genes have adjusted p-value less than 0.05

```
table(res$padj <0.05)
```

```
## ## FALSE TRUE ## 5349 10112
```

### Adjusted data

Because I suspect that sex variable can adjust the association between gene expression with age\_group factor, I will analyze statistically data with adjustment factor.

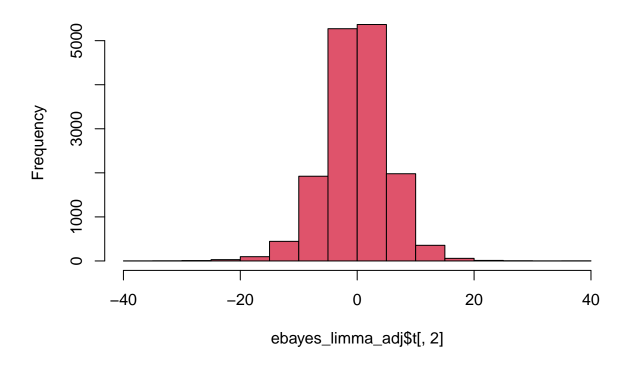
```
mod_adj = model.matrix(~ pheno$age_group+pheno$sex)
fit_limma_adj = lmFit(edata_tr,mod_adj)
ebayes_limma_adj <- eBayes(fit_limma_adj)
names(ebayes_limma_adj)</pre>
```

Fit regression with limma package with adjustment factor

```
[1] "coefficients"
                            "rank"
                                                "assign"
##
    [5] "df.residual"
                            "sigma"
                                                "cov.coefficients" "stdev.unscaled"
   [9] "pivot"
                            "Amean"
                                                                    "design"
                                                "method"
## [13] "df.prior"
                            "s2.prior"
                                                "var.prior"
                                                                    "proportion"
                            "t"
                                                "df.total"
                                                                    "p.value"
## [17] "s2.post"
  [21] "lods"
                            "F"
                                                "F.p.value"
# Statistics
```

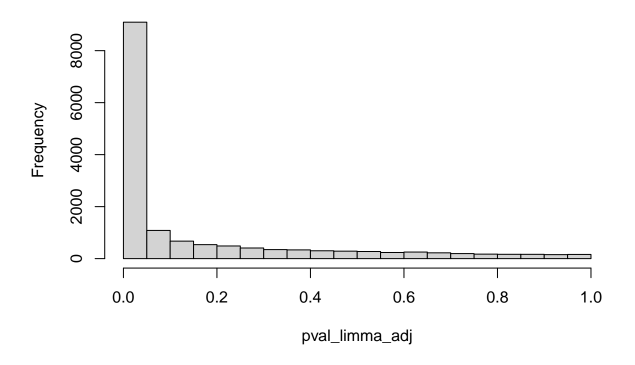
# # Statistics hist(ebayes\_limma\_adj\$t[,2], col=2)

## Histogram of ebayes\_limma\_adj\$t[, 2]



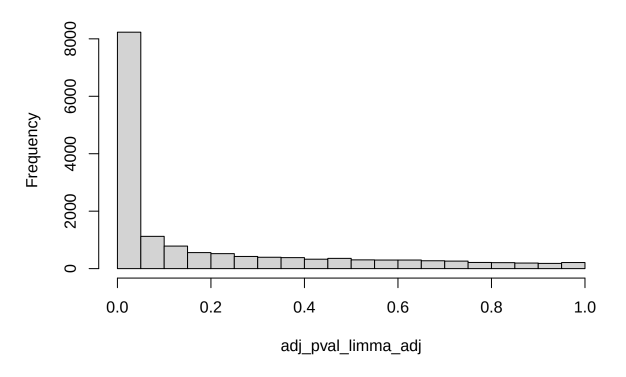
```
# P-values
pval_limma_adj = topTable(ebayes_limma_adj,number=dim(edata_tr)[1])$P.Value
hist(pval_limma_adj)
```

# Histogram of pval\_limma\_adj



# # Adjusted p-values adj\_pval\_limma\_adj = topTable(ebayes\_limma\_adj,number=dim(edata\_tr)[1])\$adj.P.Val hist(adj\_pval\_limma\_adj)

## Histogram of adj\_pval\_limma\_adj



```
re_adj = topTable(ebayes_limma_adj,number=dim(edata_tr)[1])
head(re_adj)
```

```
##
           pheno.age_groupfetus pheno.sexmale
                                                 AveExpr
                                                                F
                                  -0.09717847 11.767199 758.4099 1.177104e-11
## ST8SIA2
                       6.264017
## SOX11
                       7.196578
                                  -0.14784492 13.549118 739.4981 1.334594e-11
## SNCG
                      -6.707473
                                   0.44733611 10.307404 650.4990 2.524224e-11
## NKX6-2
                      -7.302594
                                   0.82472619 6.955729 588.5204 4.150116e-11
## TRIM54
                      -6.448248
                                   0.06076597 6.397455 537.9631 6.480537e-11
## SLA
                       5.755159
                                  -0.06607231 12.718546 509.9261 8.450284e-11
##
              adj.P.Val
## ST8SIA2 1.038247e-07
## SOX11
           1.038247e-07
           1.309147e-07
## SNCG
           1.614291e-07
## NKX6-2
## TRIM54
           2.016614e-07
## SLA
           2.191300e-07
sum(re_adj$adj.P.Val < 0.05)
```

## [1] 8232

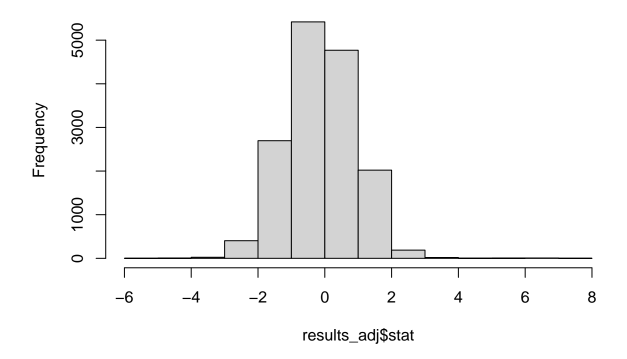
```
de_adj = DESeqDataSetFromMatrix(countData = count_filter, colData = pheno,~ age_group + sex)
glm_all_adj = DESeq(de_adj)

results_adj = results(glm_all_adj)
results_adj = as.data.frame(results_adj)
head(results_adj)
```

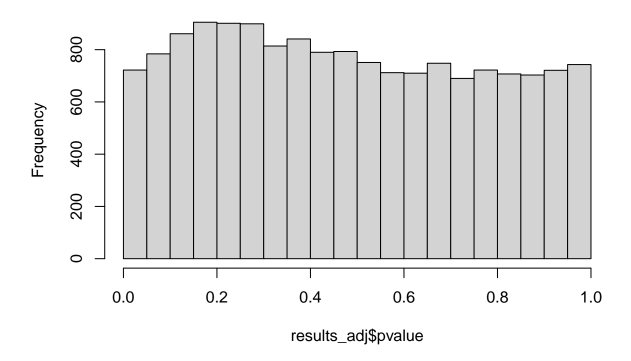
### Fit regression with DESeq with adjustment factor

```
##
                baseMean log2FoldChange
                                             lfcSE
                                                          stat
                                                                  pvalue
                                                                              padj
## A1BG
                380.8469
                             0.12906526 0.4352513 0.2965304 0.7668250 0.9776339
## ADA
                372.3584
                             -0.38152598 0.4613657 -0.8269492 0.4082658 0.9230627
## CDH2
              21681.2984
                             -0.15749391 0.1544875 -1.0194605 0.3079844 0.9171161
## AKT3
              27928.4725
                             -0.07403085 0.4782344 -0.1548003 0.8769787 0.9913890
                198.2623
                             \hbox{-0.12674096 0.2223552 -0.5699933 0.5686822 0.9462994}
## ZBTB11-AS1
## MED6
               2117.0608
                             -0.01682903 0.1510819 -0.1113901 0.9113070 0.9942778
# Statistic
hist(results_adj$stat)
```

## Histogram of results\_adj\$stat

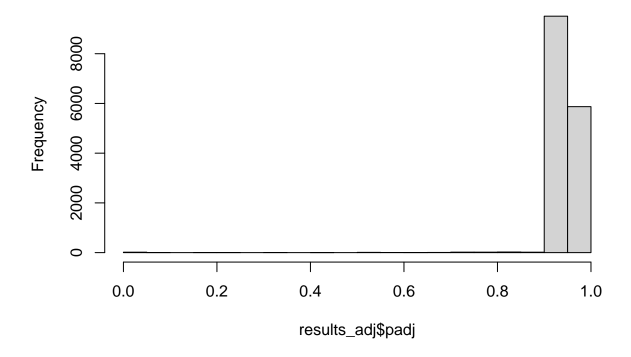


# Histogram of results\_adj\$pvalue



#Adjusted p-values
hist(results\_adj\$padj)

### Histogram of results\_adj\$padj



In the adjustment section, we can see that in the limma package, the results having adjusted p-value less than 0.05 are 8232, which also account for about 52.9% in the total of 15559 genes. However, in DESeq package, the results less than 0.05 are a few, and according to the the histogram of results\_adj pvalue from DESeq package, it is likely that there is no association between gene expression with adjusted data(including age group variable and sex variable).

### Count up and down regulated genes

Because of clear evidence of correlation when analyzing unadjusted data, which only have age\_group variable, we can see that there is a correlation between the age\_group factor and gene expression. Therefore, I will count the up-regulated genes, which are highly expressed and up-regulation in human especially for fetus, and down-regulated genes, which are down-regulation in human especially when getting older and as a result highly appear in the adult than the fetus

I will get up-regulated and down-regulated genes from unadjusted data of both limma package and DESeq package.

### Limma package

The number of up-regulated genes

```
sum(re$adj.P.Val <0.05 & re$logFC > 1)
```

## [1] 2560

```
head(up limma)
##
              logFC AveExpr
                                           P.Value
                                                      adj.P.Val
                                                                       В
## ST8SIA2 6.264017 11.76720 40.30384 2.684402e-13 2.770718e-09 20.15611
          7.196578 13.54912 39.27425 3.561563e-13 2.770718e-09 19.94839
## SOX11
## SLA
          5.755159 12.71855 33.37950 2.100303e-12 8.169655e-09 18.56884
          4.918986 11.61649 30.36669 5.882270e-12 1.382686e-08 17.71288
## FBN3
## VASH2
          5.011997 11.06801 29.96521 6.798356e-12 1.382686e-08 17.58961
          5.904494 14.73240 29.23579 8.886727e-12 1.382686e-08 17.35962
## DCX
The number of down-regulated genes
sum(re$adj.P.Val < 0.05 & re$logFC < -1)
## [1] 3080
down_limma <- re %% filter (logFC < -1 & adj.P.Val < 0.05) %>% arrange(adj.P.Val)
head(down limma)
             logFC
                                             P.Value
                     AveExpr
                                                        adj.P.Val
                                      t
## TRIM54 -6.448248 6.397455 -34.32829 1.547426e-12 8.025467e-09 18.81527
         -6.707473 10.307404 -30.34373 5.930876e-12 1.382686e-08 17.70589
## OPALIN -8.692659 8.768713 -29.51529 8.013662e-12 1.382686e-08 17.44869
## SOHLH1 -7.503042 7.626357 -29.37968 8.424990e-12 1.382686e-08 17.40562
## KRT17 -6.499428 5.955451 -27.47641 1.743578e-11 2.466212e-08 16.77083
## UAP1L1 -4.556272 8.839835 -26.51408 2.566615e-11 3.135364e-08 16.42680
DESeq package
The number of up-regulated genes
sum(res$padj < 0.05 & res$log2FoldChange > 1, na.rm=TRUE)
## [1] 3178
up_de <- res %>% filter (log2FoldChange > 1 & padj < 0.05) %>% arrange(padj)
head(up_de)
##
            baseMean log2FoldChange
                                         lfcSE
                                                   stat
## ST8SIA2 21470.986
                            7.442980 0.2003001 37.15915 3.119957e-302
## SOX11
          105896.449
                            8.543693 0.2330527 36.65991 3.180749e-294
                            6.781894 0.2163388 31.34849 1.020083e-215
## SLA
           33778.704
## FBN3
           11490.654
                            5.816179 0.2007956 28.96568 1.781244e-184
                            4.157591 0.1503316 27.65614 2.354317e-168
## MEX3B
            12548.477
## VASH2
            8164.861
                            5.920934 0.2269659 26.08732 5.077630e-150
##
                    padj
## ST8SIA2 4.823766e-298
          2.458878e-290
## SOX11
```

up\_limma <- re %>% filter (logFC > 1 & adj.P.Val < 0.05) %>% arrange(adj.P.Val)

```
## SLA 5.257168e-212
## FBN3 5.507963e-181
## MEX3B 6.066682e-165
## VASH2 9.813154e-147
```

The number of down-regulated genes

```
sum(res$padj < 0.05 & res$log2FoldChange < -1, na.rm=TRUE)</pre>
## [1] 3853
\label{localization} $\operatorname{down\_de} < -\text{res \%>\% filter (log2FoldChange} < -1 \& padj < 0.05) \%>\% arrange(padj)$
head(down_de)
##
           baseMean log2FoldChange
                                          lfcSE
                                                                   pvalue
                                                      stat
## BCL2L2 21968.220 -2.676407 0.08831507 -30.30521 9.788276e-202
## SNCG
           9473.167
                        -8.005651 0.29376717 -27.25169 1.587048e-163
## CLMN
           3227.245
                         -3.760479 0.14437310 -26.04695 1.456668e-149
## UAP1L1 1538.034
                         -5.591225 0.23073417 -24.23232 1.015679e-129
## ITPKA
           5882.483
                          -7.047788 0.29600941 -23.80934 2.673017e-125
## OPALIN 7380.398
                         -10.922203 0.45988078 -23.75008 1.096772e-124
##
                    padj
## BCL2L2 3.783413e-198
## SNCG 3.505336e-160
## CLMN
        2.502394e-146
## UAP1L1 1.427583e-126
## ITPKA 3.443960e-122
## OPALIN 1.304400e-121
```

The number of common up-regulated and also down-regulated genes from both DESeq package and Limma package  ${\bf P}$ 

```
up <- rownames(up_de) %in% rownames(up_limma)
table(up)

## up
## FALSE TRUE
## 638 2540

down <- rownames(down_de) %in% rownames(down_limma)
table(down)

## down
## FALSE TRUE
## 827 3026

up = up_de[up,]
head(up)</pre>
```

```
baseMean log2FoldChange
                                          lfcSE
                                                    stat
                                                                pvalue
## ST8SIA2 21470.986
                            7.442980 0.2003001 37.15915 3.119957e-302
## SOX11
           105896.449
                            8.543693 0.2330527 36.65991 3.180749e-294
## SLA
            33778.704
                            6.781894 0.2163388 31.34849 1.020083e-215
## FBN3
            11490.654
                            5.816179 0.2007956 28.96568 1.781244e-184
                            4.157591 0.1503316 27.65614 2.354317e-168
## MEX3B
            12548.477
                            5.920934 0.2269659 26.08732 5.077630e-150
## VASH2
             8164.861
##
## ST8SIA2 4.823766e-298
## SOX11
           2.458878e-290
## SLA
           5.257168e-212
## FBN3
           5.507963e-181
## MEX3B
           6.066682e-165
## VASH2
           9.813154e-147
down = down_de[down,]
head(down)
##
           baseMean log2FoldChange
                                         lfcSE
                                                    stat
                                                                pvalue
## BCL2L2 21968.220
                         -2.676407 0.08831507 -30.30521 9.788276e-202
## SNCG
           9473.167
                         -8.005651 0.29376717 -27.25169 1.587048e-163
                         -3.760479 0.14437310 -26.04695 1.456668e-149
## CLMN
           3227.245
## UAP1L1
           1538.034
                         -5.591225 0.23073417 -24.23232 1.015679e-129
## ITPKA
           5882.483
                         -7.047788 0.29600941 -23.80934 2.673017e-125
## OPALIN
                        -10.922203 0.45988078 -23.75008 1.096772e-124
           7380.398
##
## BCL2L2 3.783413e-198
## SNCG
          3.505336e-160
## CLMN
          2.502394e-146
## UAP1L1 1.427583e-126
## ITPKA 3.443960e-122
## OPALIN 1.304400e-121
```

As we can see, there are 2540 common up-regulated genes and 3026 common down-regulated genes between limma package and DESeq package.

In addition to analyzing the correlation in R, I also want to predict and classify some characteristics of those 10 samples such as gender, or age by using Python. Below is the preparation for that process.

### Preparing data for prediction and classification

```
up_down_reg = rbind(up,down)
dim(up_down_reg)

## [1] 5566 6

up_down_tr <- (rownames(edata_tr) %in% rownames(up_down_reg))
table(up_down_tr)

## up_down_tr
## FALSE TRUE
## 9993 5566</pre>
```

```
up_down = edata_tr[up_down_tr,]
head(up_down)
          SRR1554534 SRR1554535 SRR1554568 SRR1554561 SRR1554567 SRR1554536
##
## ADA
                      9.145974
                               7.303994
                                           8.001912
                                                     7.351627
          8.646385
                                                                9.770115
## CDH2
          13.562468 13.585906 14.957846 13.652418 14.829826 13.812075
## AKT3
          13.479431 13.159273 15.876052 13.925121 14.376621 12.332018
## ACOT8
           11.946038 11.757942 10.652184 12.014095 10.470291 11.250633
## ZBTB33
          11.724310 12.137270 13.056753 11.858819 12.975194 11.759541
## ZSCAN30 11.187060 11.583213 12.612542 11.040663 12.383769 11.699403
          SRR1554541 SRR1554539 SRR1554538 SRR1554537
## ADA
          7.843839 8.253862 7.831596
                                          7.197164
## CDH2
          14.545283 13.717079 14.708753 15.051336
## AKT3
           15.204848 14.383520 14.749481 15.584859
## ACOT8
           10.522167 11.461451 10.601639 10.700495
## ZBTB33
           12.988798 12.410141 13.263124 13.212136
## ZSCAN30 12.182915 11.462477 12.586344 12.277068
# df is saved in a name "data for regulated gene.csv"
df <- merge(up_down_reg,up_down, by =0)</pre>
row.names(df) <- df$Row.names</pre>
df = df[,-1]
head(df)
##
           baseMean log2FoldChange
                                     lfcSE
                                                           pvalue
                                                 stat
                                                                         padj
## A2M
         13185.3832 -1.670680 0.4204894 -3.973179 7.091974e-05 1.717291e-04
## A2ML1
          484.6328
                        -2.748295 0.5692828 -4.827644 1.381576e-06 4.322248e-06
## A4GALT
          412.3115
                       -2.790862 0.5719644 -4.879432 1.063917e-06 3.379052e-06
## AARD
                       -2.105662 0.5372000 -3.919699 8.865947e-05 2.115377e-04
           124.8856
## AARS1 33645.6493
                        -1.564934 0.2709131 -5.776517 7.626273e-09 3.150984e-08
## AATK
       21134.6178
                        -3.684225 0.3588961 -10.265435 1.008658e-24 1.959154e-23
##
         SRR1554534 SRR1554535 SRR1554568 SRR1554561 SRR1554567 SRR1554536
## A2M
         13.657416 13.795969 12.584194 13.003245 12.597035 15.235020
                              6.909735 8.644313
## A2ML1
          8.848175 9.062477
                                                   7.457088 10.622862
## A4GALT 8.868851 8.344972 6.895125 8.568670 6.710507 10.362960
## AARD
          7.347104 7.314591
                              5.402734 6.636227
                                                    6.395016
                                                             7.306073
## AARS1 15.757594 15.345289 14.133470 15.908997 14.037013 14.566318
## AATK
          15.412314 15.065392 11.532492 15.465764 11.828298 13.675639
##
         SRR1554541 SRR1554539 SRR1554538 SRR1554537
## A2M
         12.938130 13.881265 12.930901 12.535831
## A2ML1
          6.885452 7.708383 7.500564 7.351208
## A4GALT
         7.260562 7.533763 7.292369 6.723675
## AARD
          6.484457 7.470666
                              5.264764 5.737391
## AARS1
          14.312565 15.532491 14.157486 14.182287
          12.056377 14.628722 11.641881 12.151114
## AATK
# save file
```

# write.csv(df, file = "D:/word/bioinformatics/personal project/genomic-data-science-project-about-fetu