# SPEAQeasy Differential Expression Analysis

# 9/18/2020

### Contents

1	Analysis		
	1.1	Load required libraries	1
	1.2	Load data and prepare directories to place outputs in $\ldots \ldots \ldots \ldots \ldots \ldots$	2
	1.3	statistics PCs	2
	1.4	Stats vs. race and brain region	2
	1.5	Explore and visualize gene expression	7
	1.6	Modeling	29
	1.7	Check plots	32
	1.8	Gene ontology	36
	1.9	Visualize differentially expressed genes	43
<b>2</b>	Rep	producibility	44

# 1 Analysis

The following analysis explores a RangedSummarizedExperiment object from the SPEAQeasy pipeline. Note that we will use a modified version of the object, which resolved sample identity issues which were present in the raw output from SPEAQeasy. This object also includes phenotype data added after resolving identity issues. Though SPEAQeasy produces objects for several feature types (genes, exons, exon-exon junctions), we will demonstrate an example analysis for just genes. We will perform differential expression across some typical variables of interest (e.g. sex, age, race) and show how to perform principal component analysis (PCA) and visualize findings with plots.

### 1.1 Load required libraries

```
library("SummarizedExperiment")
library("recount")
library("edgeR")
library("limma")
library("jaffelab") # GitHub: LieberInstitute/jaffelab
library("RColorBrewer")
library("clusterProfiler")
library("org.Hs.eg.db")
```

```
library("pheatmap")
library("here")
library("devtools")
library("BiocStyle")
```

### 1.2 Load data and prepare directories to place outputs in

For those who ran SPEAQeasy from the example FASTQ data set, the RangedSummarizedExperiment will have a different path, as specified with the --output flag.

```
# Load the RSE gene object
load(here("rse_speaqeasy.RData"), verbose = TRUE)

## Loading objects:
## rse_gene

# Create directories to organize outputs from this analysis
dir.create(here("DE_analysis", "pdfs"), showWarnings = FALSE)
dir.create(here("DE_analysis", "tables"), showWarnings = FALSE)
dir.create(here("DE_analysis", "rdas"), showWarnings = FALSE)
```

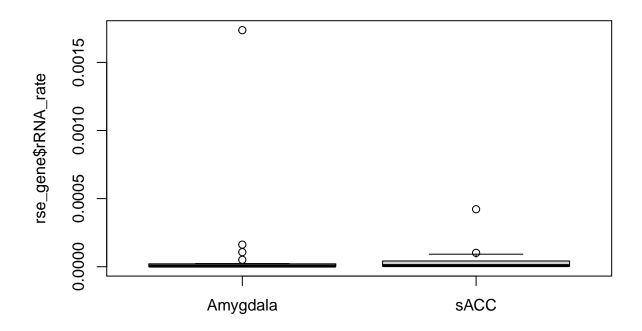
### 1.3 statistics PCs

Here we are using principal component analysis to control for the listed variables impact on expression. This will be later added into our linear model.

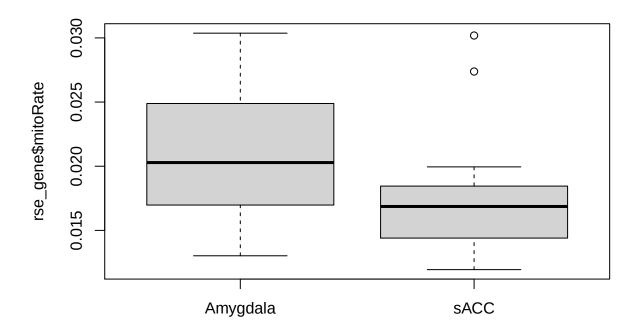
## [1] 87.3

### 1.4 Stats vs. race and brain region

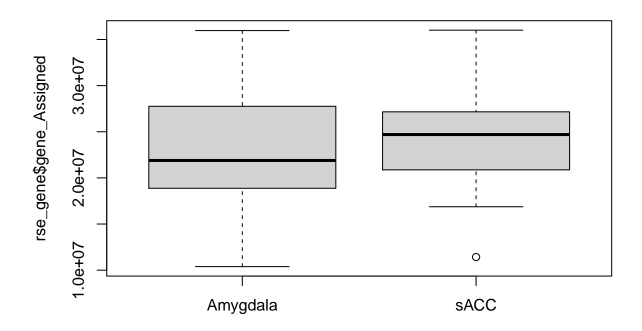
```
# Filter for expressed
rse_gene = rse_gene[rowMeans(getRPKM(rse_gene,"Length")) > 0.2,]
# Check if certain statistics changed by race or region
# Display box plots here
boxplot(rse_gene$rRNA_rate ~ rse_gene$BrainRegion,xlab="")
```



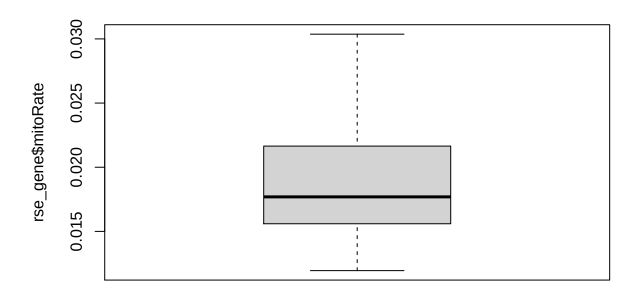
boxplot(rse\_gene\$mitoRate ~ rse\_gene\$BrainRegion,xlab="")



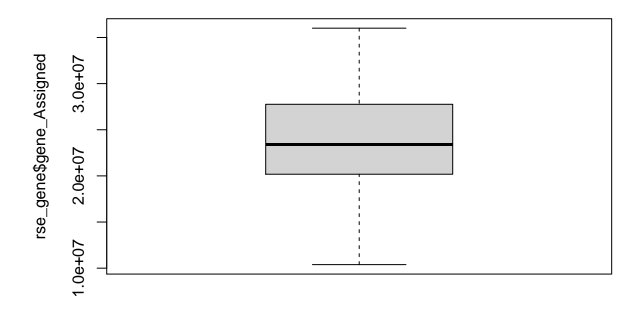
boxplot(rse\_gene\$gene\_Assigned ~ rse\_gene\$BrainRegion,xlab="")



boxplot(rse\_gene\$mitoRate ~ rse\_gene\$Race,las=3,xlab="")



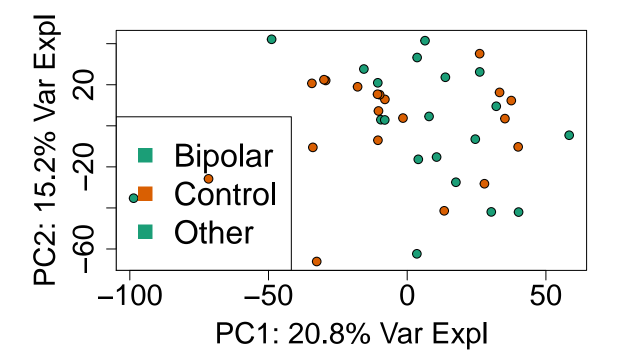
boxplot(rse\_gene\$gene\_Assigned ~ rse\_gene\$Race,las=3,xlab="")

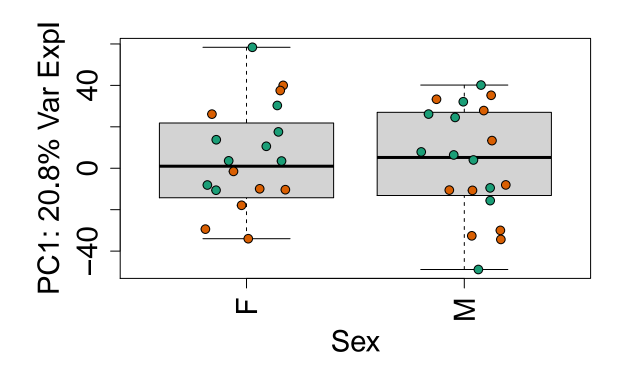


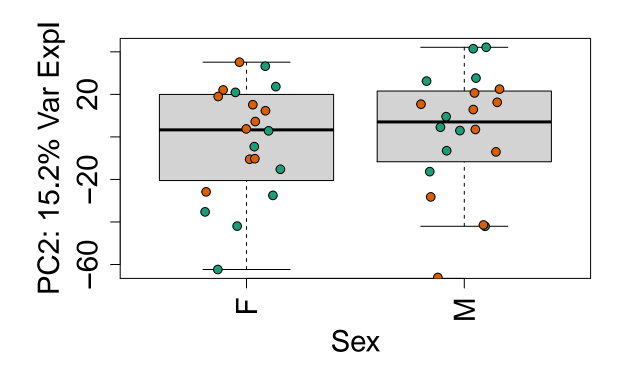
```
# Save box plots to PDF
pdf(file = here("DE_analysis", "pdfs", "Region_Race_cellcheck.pdf"))
boxplot(rse_gene$rRNA_rate ~ rse_gene$BrainRegion,xlab="")
boxplot(rse_gene$mitoRate ~ rse_gene$BrainRegion,xlab="")
boxplot(rse_gene$gene_Assigned ~ rse_gene$BrainRegion,xlab="")
boxplot(rse_gene$mitoRate ~ rse_gene$Race,las=3,xlab="")
boxplot(rse_gene$gene_Assigned ~ rse_gene$Race,las=3,xlab="")
dev.off()
## pdf
## pdf
## pdf
```

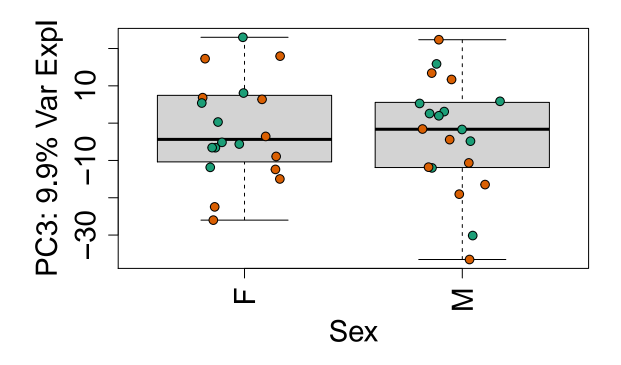
# 1.5 Explore and visualize gene expression

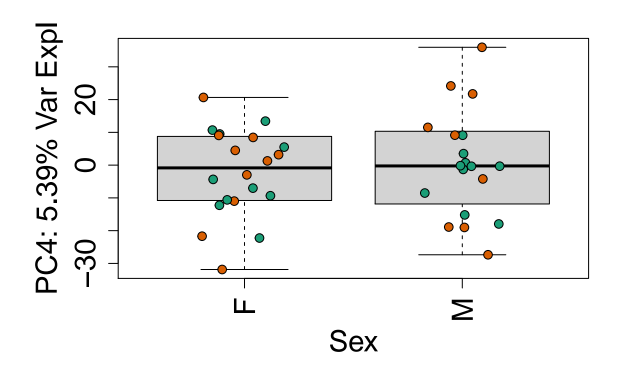
```
par(mar=c(8,6,2,2),cex.axis=1.8,cex.lab=1.8)
   palette(brewer.pal(4,"Dark2"))
    # PC1 vs. PC2
   plot(pca$x, pch=21, bg=factor(rse_gene$PrimaryDx),cex=1.2,
        xlab = pca_vars_lab[1], ylab = pca_vars_lab[2])
   legend("bottomleft", levels(rse_gene$PrimaryDx), col=1:2, pch=15,cex=2)
    # By line
   for(i in 1:10) {
       boxplot(pca$x[,i] ~ rse_gene$Sex,
            ylab=pca_vars_lab[i], las = 3,xlab="Sex",outline=FALSE)
       points( pca$x[,i] ~ jitter(as.numeric(factor(rse_gene$Sex))),
           pch = 21, bg = rse_gene$PrimaryDx,cex=1.2)
   }
    # By experiment
   for(i in 1:10) {
       boxplot(pca$x[,i] ~ rse_gene$Race,
            ylab=pca_vars_lab[i], las = 3,xlab="Race",outline=FALSE)
       points( pca$x[,i] ~ jitter(as.numeric(factor(rse_gene$Race))),
           pch = 21, bg = rse_gene$PrimaryDx,cex=1.2)
   }
}
# Display plots
generate_plots()
```

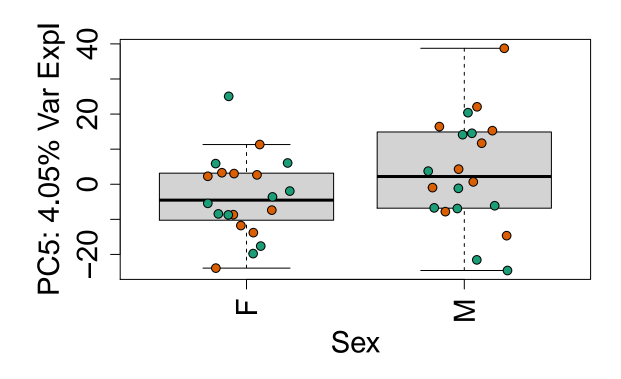


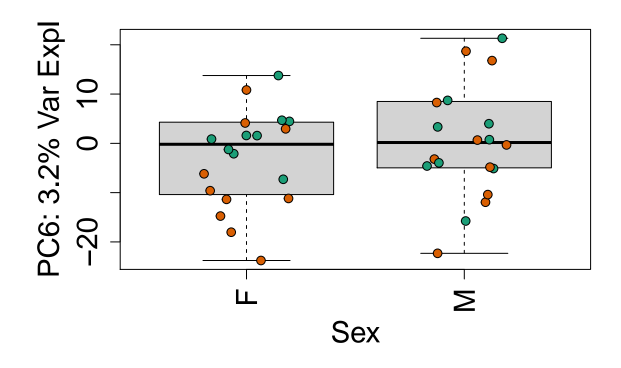


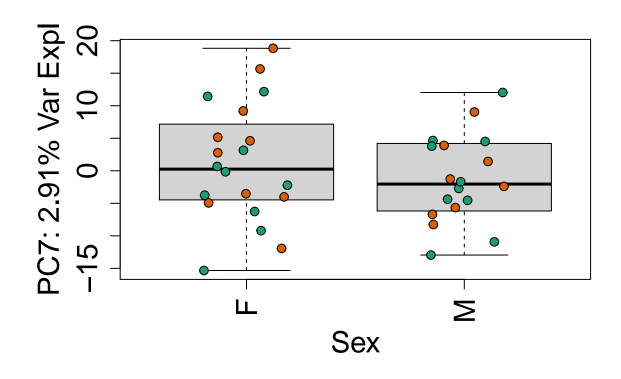


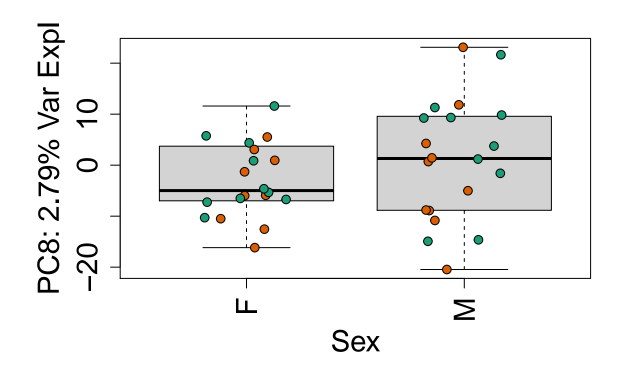


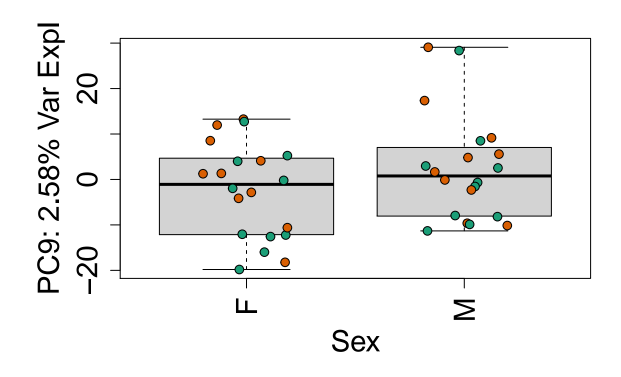


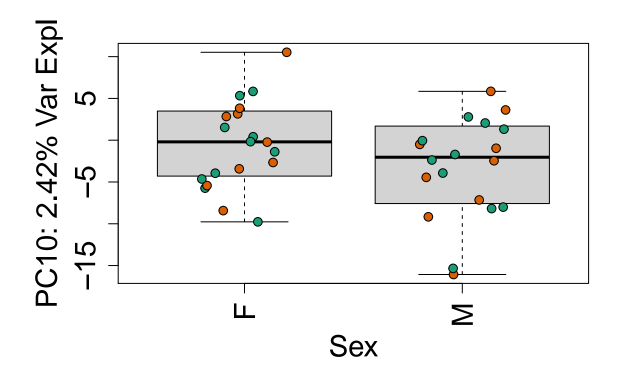


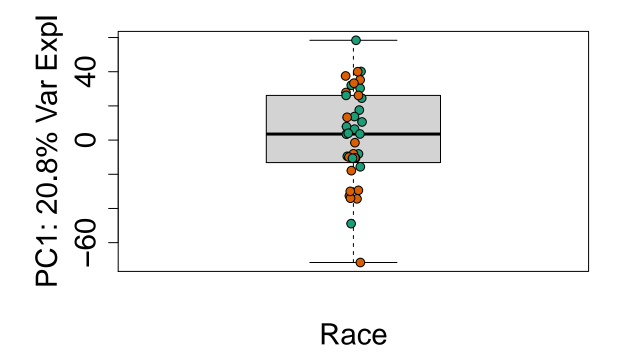


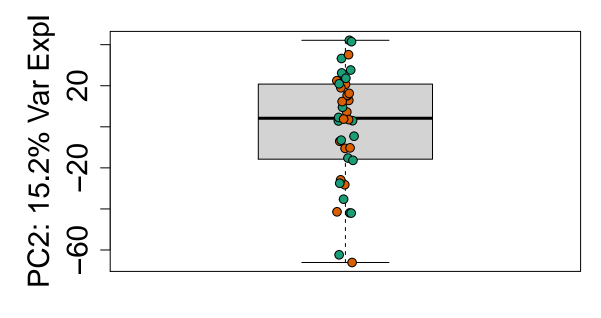




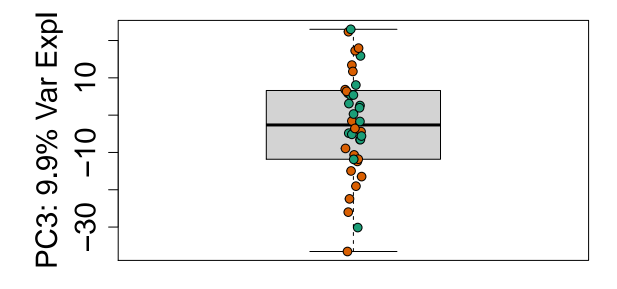




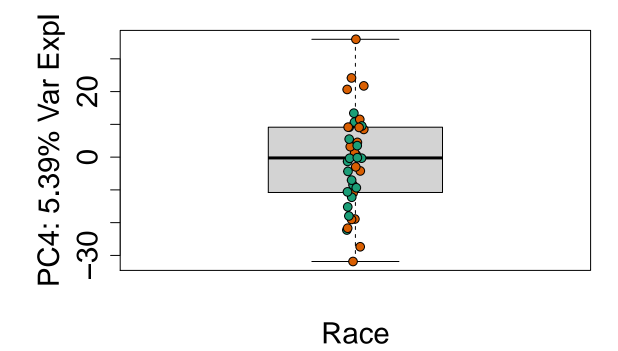


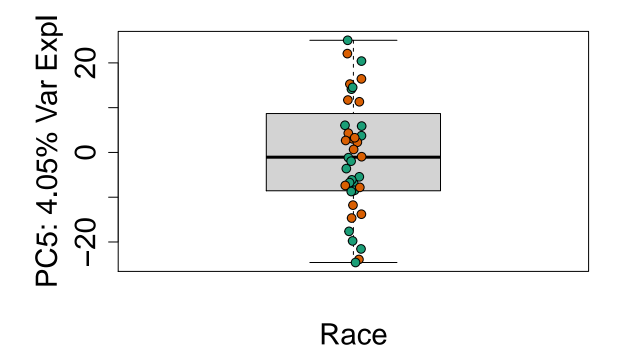


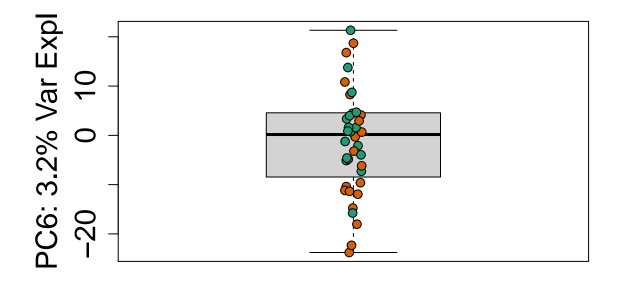
Race



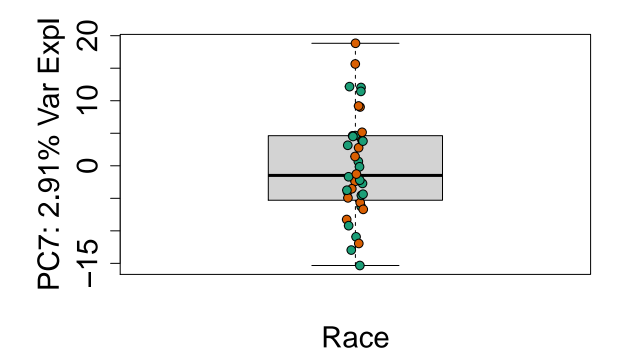
Race

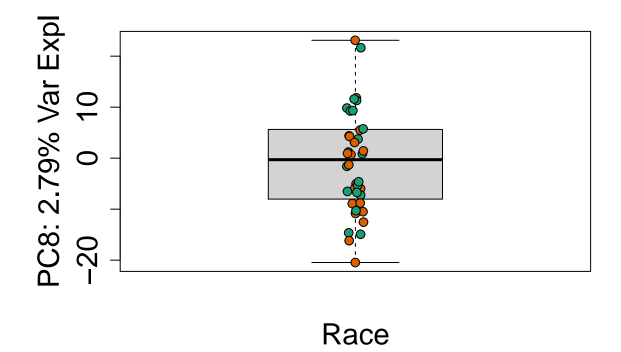


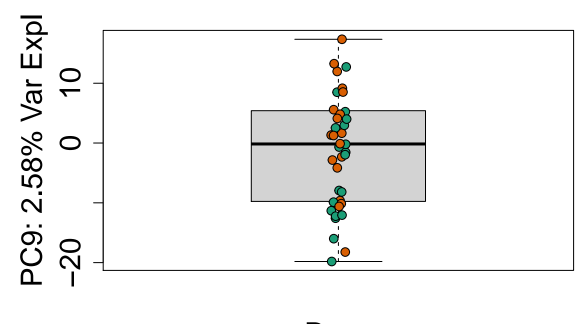


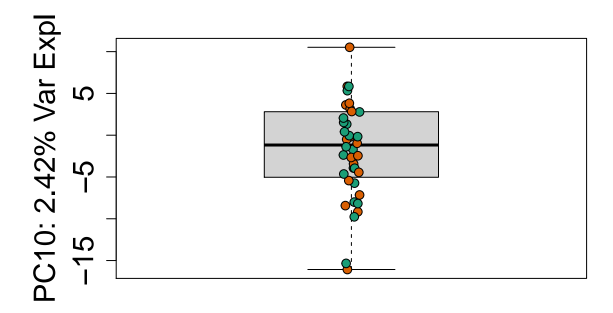


Race









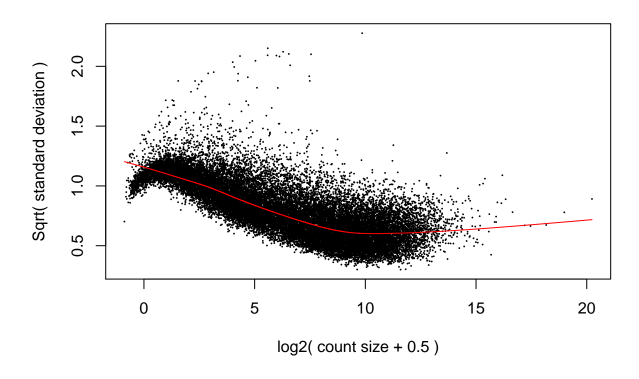
# Race

```
# Write plots to PDF
pdf(here("DE_analysis", "pdfs", "PCA_plotsExprs.pdf"), w=9)
generate_plots()
dev.off()

## pdf
## 2
```

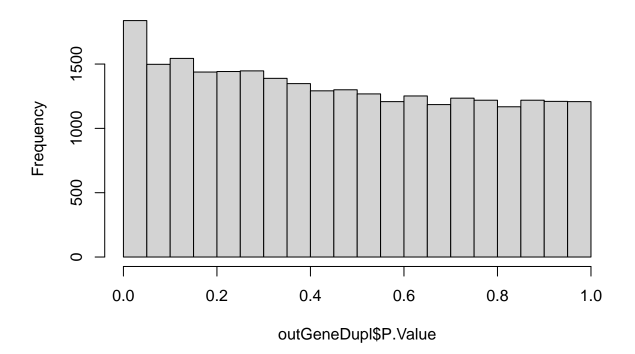
# 1.6 Modeling

# voom: Mean-variance trend



```
# Also write mean-variance plot to PDF
pdf(file = "DE_analysis/pdfs/vGene.pdf")
invisible(voom(dge,mod,plot=TRUE))
## Coefficients not estimable: PrimaryDxOther
## Warning: Partial NA coefficients for 26708 probe(s)
dev.off()
## pdf
##
# Get duplicate correlation
gene_dupCorr = duplicateCorrelation(vGene$E, mod,
    block=colData(rse_gene)$SAMPLE_ID)
## Coefficients not estimable: PrimaryDxOther
save(gene_dupCorr, file = here("DE_analysis", "rdas", "gene_dupCorr_neurons.rda"))
# Fit linear model
fitGeneDupl = lmFit(vGene,
    correlation=gene_dupCorr$consensus.correlation,
    block=colData(rse_gene)$SAMPLEID)
```

# Histogram of outGeneDupl\$P.Value



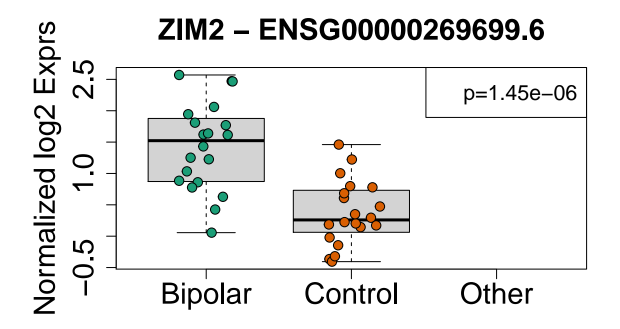
```
table(outGeneDupl$adj.P.Val < 0.05)</pre>
##
## FALSE TRUE
## 26707
table(outGeneDupl$adj.P.Val < 0.1)</pre>
##
## FALSE TRUE
## 26706
sigGeneDupl = topTable(ebGeneDupl,coef=2,
    p.value = 0.1,number=nrow(rse_gene))
sigGeneDupl[,c("Symbol","logFC", "P.Value","AveExpr")]
##
                     Symbol
                                 logFC
                                            P.Value
                                                      AveExpr
                      ZIM2 -1.1238665 1.448916e-06 0.8970203
## ENSG00000269699.6
## ENSG00000126368.6 NR1D1 -0.7637626 6.833687e-06 4.6128957
sigGeneDupl$logFC > 0,c("Symbol","logFC", "P.Value")]
## [1] Symbol logFC P.Value
## <0 rows> (or 0-length row.names)
sigGeneDupl[sigGeneDupl$logFC < 0,c("Symbol","logFC", "P.Value")]</pre>
##
                     Symbol
                                 logFC
                                            P. Value
                      ZIM2 -1.1238665 1.448916e-06
## ENSG00000269699.6
## ENSG00000126368.6 NR1D1 -0.7637626 6.833687e-06
write.csv(outGeneDupl, file = here("DE_analysis", "tables", "de_stats_allExprs.csv"))
write.csv(sigGeneDupl, file = here("DE_analysis", "tables", "de_stats_fdr10_sorted.csv"))
```

# 1.7 Check plots

```
exprs = vGene$E[rownames(sigGeneDupl),]

# Group together code for displaying boxplots
generate_plots = function() {
   par(mar=c(8,6,4,2),cex.axis=1.8,cex.lab=1.8, cex.main=1.8)
   palette(brewer.pal(4,"Dark2"))

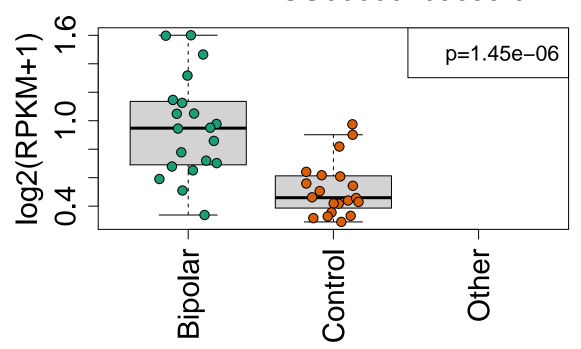
for(i in 1:nrow(sigGeneDupl)) {
   yy = exprs[i,]
   boxplot(yy ~ rse_gene$PrimaryDx, outline=FALSE,
```



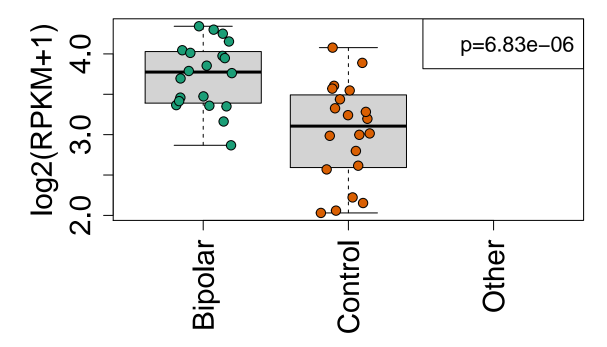
# NR1D1 – ENSG00000126368.6 S2.5 2.7 p=6.83e-06 Bipolar Control Other

```
# Write plots to PDF
pdf(here("DE_analysis", "pdfs", "DE_boxplots_byDiagnosis.pdf"), w=10)
generate_plots()
dev.off()
## pdf
##
e = geneExprs[rownames(sigGeneDupl),]
generate_plots = function() {
    par(mar=c(8,6,4,2),cex.axis=1.8,cex.lab=1.8, cex.main=1.8)
    palette(brewer.pal(4,"Dark2"))
    for(i in 1:nrow(sigGeneDupl)) {
        yy = e[i,]
        boxplot(yy ~ rse_gene$PrimaryDx, las=3,outline=FALSE,
              ylim=range(yy), ylab="log2(RPKM+1)", xlab="",
        main = paste(sigGeneDupl$Symbol[i], "-", sigGeneDupl$gencodeID[i]))
        points(yy ~ jitter(as.numeric(rse_gene$PrimaryDx)),
            pch = 21, bg= rse_gene$PrimaryDx,cex=1.3)
        11 = ifelse(sigGeneDupl$logFC[i] > 0, "topleft", "topright")
        legend(ll, paste0("p=", signif(sigGeneDupl$P.Value[i],3)), cex=1.3)
    }
}
```

# **ZIM2 - ENSG00000269699.6**



# NR1D1 - ENSG00000126368.6



```
# Write the same plots to PDF
pdf(here("DE_analysis", "pdfs", "DE_boxplots_byGenome_log2RPKM.pdf"), w=10)
generate_plots()
dev.off()

## pdf
## 2
```

# 1.8 Gene ontology

clusterProfiler is a gene ontology package we will use to see if our genes are specifically differentially expressed in certain pathways.

```
# Get significant genes by sign
sigGene = outGeneDupl[outGeneDupl$P.Value < 0.005,]
sigGeneList = split(as.character(sigGene$EntrezID), sign(sigGene$logFC))
sigGeneList = lapply(sigGeneList, function(x) x[!is.na(x)])
geneUniverse = as.character(outGeneDupl$EntrezID)
geneUniverse = geneUniverse[!is.na(geneUniverse)]

# Do GO and KEGG
goBP_Adj <- compareCluster(sigGeneList, fun = "enrichGO",
    universe = geneUniverse, OrgDb = org.Hs.eg.db,
    ont = "BP", pAdjustMethod = "BH", pvalueCutoff = 1,</pre>
```

```
qvalueCutoff = 1, readable= TRUE)
goMF_Adj <- compareCluster(sigGeneList, fun = "enrichGO",</pre>
   universe = geneUniverse, OrgDb = org.Hs.eg.db,
   ont = "MF", pAdjustMethod = "BH", pvalueCutoff = 1,
   qvalueCutoff = 1, readable= TRUE)
goCC_Adj <- compareCluster(sigGeneList, fun = "enrichGO",</pre>
   universe = geneUniverse, OrgDb = org.Hs.eg.db,
   ont = "CC", pAdjustMethod = "BH", pvalueCutoff = 1,
   qvalueCutoff = 1, readable= TRUE)
kegg_Adj <- compareCluster(sigGeneList, fun = "enrichKEGG",</pre>
   universe = geneUniverse, pAdjustMethod = "BH",
   pvalueCutoff = 1, qvalueCutoff = 1)
save(goBP_Adj, goCC_Adj, goMF_Adj, kegg_Adj,
   file = here("DE_analysis", "rdas", "gene_set_objects_p005.rda"))
goList = list(BP = goBP_Adj, MF = goMF_Adj, CC = goCC_Adj, KEGG = kegg_Adj)
goDf = dplyr::bind_rows(lapply(goList, as.data.frame), .id = "Ontology")
goDf = goDf[order(goDf$pvalue),]
write.csv(goDf, file = here("DE_analysis", "tables", "geneSet_output.csv"), row.names=FALSE)
options(width=130)
goDf[goDf$p.adjust < 0.05, c(1:5,7)]
```

	ID	Cluster	Ontology		##
tumor necrosis factor	GO:0032640	1	BP	1311	##
tumor necrosis factor superfamily cytokine	GO:0071706	1	BP	1312	##
regulation of cell	GO:0050865	1	BP	1313	##
interleukin-1 beta	GO:0032611	1	BP	1314	##
interleukin-1	GO:0032612	1	BP	1315	##
regulation of tumor necrosis factor	GO:0032680	1	BP	1316	##
regulation of tumor necrosis factor superfamily cytokine	GO:1903555	1	BP	1317	##
regulation of interleukin-1 beta	GO:0032651	1	BP	1318	##
positive regulation of cell	GO:0050867	1	BP	1319	##
leukocyte diff	GO:0002521	1	BP	1320	##
regulation of leukocyte	GO:0002694	1	BP	1321	##
regulation of interleukin-1	GO:0032652	1	BP	1322	##
response t	GD:0009617	1	BP	1323	##
defense response t	GO:0042742	_	BP	1324	
cellular response to inter	GO:0071346		BP	1325	
leukocyt	GD:0050900		BP	1326	
G protein-coupled purinergic nucleotide recept	GO:0045028		MF	3316	
positive regulation of leukocyt	GD:0002687		BP	1327	
response to inter	GO:0034341		BP	1328	
1-phosphatidylinositol-3-kinase regulat	GO:0046935		MF	3317	
regulation of defer	GO:0031347	1	BP	1329	##
regulation of leukocyt	GO:0002685		BP	1330	
cellular response to molecule of bacte	GO:0071219	1	BP	1331	##

positive regulation of tumor necrosis factor pa

## 1332

BP

1 GO:0032760

regulation of glial cell	903975	1 (	BP	1333	##
positive regulation of tumor necrosis factor superfamily cytokine p	903557	1 (	BP	1334	##
interleukin-6 p	032635		BP	1335	##
pha	006909	1 (	BP	1336	##
interferon-gamma p	032609	1 (	BP	1337	##
macrophage a	042116		BP	1338	
phosphatidylinositol 3-kinase regulator	035014	1 (	MF	3318	
positive regulation of leukocyte a	002696		BP	1339	
interleukin-1 beta	050702		BP	1340	
definitive he	060216		BP	1341	
purinergic nucleotide receptor	001614		MF	3319	
nucleotide receptor	016502		MF	3320	
cellular response to biotic	071216		BP	1342	
G protein-coupled receptor	004930	1 (	MF	3321	
interleukin-1	050701	1 (	BP	1343	##
positive regulation of macrophage a	043032	1 (	BP	1344	##
leukocyte tethering o	050901	1 (	BP	1345	##
positive regulation of interferon-gamma p	032729	1 (	BP	1346	##
phagocytosis, e	006911	1 (	BP	1347	##
chemokine-mediated signalin	070098	1 (	BP	1348	##
h	007599	1 (	BP	1349	##
negative regulation of interleukin-1 beta p	032691	1 (	BP	1350	##
E-bo	070888	-1 (	MF	3137	##
negative regulation of interleukin-1 p	032692	1 (	BP	1351	##
positive regulation of cytokine p	001819	1 (	BP	1352	##
positive regulation of defense	031349	1 (	BP	1353	##
positive regulation of pha	050766	1 (	BP	1354	##
Viral protein interaction with cytokine and cytokine	a04061	1	KEGG	3871	##
positive regulation of MAP	043410	1 (	BP	1355	##
cellular response to lipopolys	071222	1 (	BP	1356	##
plasma membrane inv	099024	1 (	BP	1357	##
response to	990868	1 (	BP	1358	##
cellular response to	990869	1 (	BP	1359	##
interleukin-8 p	032637	1 (	BP	1360	##
positive regulation of inflammatory	050729	1 (	BP	1361	##
positive regulation of interleukin-1 beta	050718	1 (	BP	1362	##
regulation of interleukin-6 p	032675	1 (	BP	1363	##
purinergic nucleotide receptor signalin	035590	1 (	BP	1364	##
positive regulation of interleukin-1	050716	1 (	BP	1365	##
positive regulation of response to external	032103	1 (	BP	1366	##
regulation of response to biotic	002831	1 (	BP	1367	##
myeloid cell activation involved in immune	002275	1 (	BP	1368	##
granulocyte differ	030851	1 (	BP	1369	##
positive regulation of lymphocyte	000403	1 (	BP	1370	##
myeloid leukocyte differ	002573	1 (	BP	1371	##
membrane inv	010324	1 (	BP	1372	##
leukocyte adhesion to vascular endothe	061756	1 (	BP	1373	##
regulation of interferon-gamma p	032649	1 (	BP	1374	##
	a04060	1	KEGG	3872	##
	005126	1 (	MF	3322	##
	032496	1 (	BP	1375	##
	903532	1 (	BP	1376	##
lymphocyte	072676	1 (	BP	1377	##
leukocyte cell-cell	007159	1 (	BP	1378	##

positive regulation of interleukin-6 p	GO:0032755	1	BP	1379	##
response to molecule of bacteri	GO:0002237	1	BP	1380	##
negative regulation of immune syste	GO:0002683	1	BP	1381	##
regulation of macrophage	GO:1905521	1	BP	1382	##
blood co	GO:0007596	1	BP	1383	##
cc	GO:0050817	1	BP	1384	##
regulation of interleukin-1 beta	GO:0050706	1	BP	1385	##
positive regulation of	GO:0051047	1	BP	1386	##
regulation of cytokine biosyntheti	GO:0042035	1	BP	1387	##
regulation of pha	GO:0050764	1	BP	1388	##
microglial cell a	GO:0001774	1	BP	1389	##
leukocyte activation involved in inflammatory	GD:0002269	1	BP	1390	##
macrophage differ	GO:0030225	1	BP	1391	##
regulation of interleukin-1	GO:0050704	1	BP	1392	##
RNA polymerase II cis-regulatory region sequence-specific DN	GO:0000978	1	MF	3323	##
positive regulation of cell	GO:0030335	1	BP	1393	##
GT	GO:0005525	1	MF	3324	##
myeloid cell differ	GO:0030099	1	BP	1394	##
interleukin-6	GO:0072604	1	BP	1395	##
purine ribonucleosid	GO:0032550	1	MF	3325	##
purine nucleosid	GO:0001883	1	MF	3326	##
ribonucleosid	GO:0032549	1	MF	3327	##
regulation of blood	GO:0008217	1	BP	1396	##
cytokine biosyntheti	GO:0042089	1	BP	1397	##
cytokine metaboli	GO:0042107	1	BP	1398	##
cis-regulatory region sequence-specific DN	GO:0000987	1	MF	3328	##
regulation of leukocyte differ	GO:1902105	1	BP	1399	##
positive regulation of ang	GO:0045766	1	BP	1400	##
positive regulation of cell	GO:2000147	1	BP	1401	##
alpha-beta T cell lineage o	GO:0002363	1	BP	1402	##
positive regulation of neuroinflammatory	GO:0150078	1	BP	1403	##
regulation of mast cell a	GD:0033003	1	BP	1404	##
nucleosid	GO:0001882	1	MF	3329	##
positive regulation of interleukin-1 beta p	GO:0032731	1	BP	1405	##
guanyl nucleotid	GD:0019001	1	MF	3330	##
guanyl ribonucleotid	GO:0032561	1	MF	3331	##
negative regulation of cytokine p	GO:0001818	1	BP	1406	
stem cell differ	GD:0048863	1	BP	1407	##
regulation of macrophage a	GD:0043030	1	BP	1408	##
regulation of immune effects	GD:0002697		BP	1409	
positive regulation of l	GO:0040017		BP	1410	
CD4-positive or CD8-positive, alpha-beta T cell lineage of	GO:0043369		BP	1411	
synaps	GD:0098883		BP	1412	
positive regulation of cellular component	GO:0051272		BP	1413	
E-bc	GO:0070888		MF	3332	
myeloid leukocyte	GO:0097529	1	BP	1414	##
regulation of leukocyte degr	GO:0043300		BP	1415	
hematopoietic progenitor cell differ	GD:0002244		BP	1416	
regulation of leukocyte mediated	GO:0002703		BP	1417	##
positive regulation of ERK1 and ERK	GO:0070374	1	BP	1418	
CCR chemokine receptor	GO:0048020		MF	3333	
positive regulation of he	GO:1903708		BP	1419	
positive regulation of interleukin-1 p	GO:0032732	1	BP	1420	
macrophage	GO:1905517	1	BP	1421	##

##	1422 B	P 1	GO	0:0070372	
##	1423 B	P 1	GO	0:1903037	
##	1424 B	P 1	GO	0:0071692	
##	1425 B	P 1	GO	0:0045088	
##	1426 B	P 1	GO	0:0061900	
##	1427 B	P 1	GO	0:0010919 re	gul
##	1428 B	P 1	GO	0:0033004	
##	1429 B	P 1	GO	D:0060100	
##	1430 B	P 1	GO	0:1905155	
##	1431 B	P 1	GO	0:0050878	
##	1432 B	P 1	GO	0:0002761	
##	1433 B	P 1	GO	0:1904018	
##	1434 B	P 1	GO	0:0008347	
##	1435 B	P 1	GO	0:0043551 r	egu
##	1436 B	P 1	GO	0:0072678	
##	3334 M	IF 1	GO	0:0019955	
##	1437 B	P 1	GO	0:0001562	
##	1438 B			0:0042832	
##	1439 B	P 1	GO	D:0060099	
##	1440 B	P 1	GO	0:1903978	
##	1441 B	P 1	GO	0:2000401	
##	1442 B	P 1	GO	0:0043270	
##	1443 B	P 1	GO	0:0042108 p	osi
##	1444 B	P 1	GO	0:0070371	
##	1445 B	P 1	GO	0:0002790	
##	1446 B			D:0002360	
##	1447 B	P 1	GO	0:0042104 po	sit
##	1448 B			0:1905153	
##	1449 B	P 1	GO	0:0002886	r
##	1450 B	P 1	GO	0:0045765	
##	1451 B	P 1	GO	0:0032677	
##	1452 B	P 1	GO	0:0045123	
##	1453 B			0:0050727	
##	1454 B			0:0002577	r
##	1455 B			0:0150146	
##	1456 B			0:0006954	
##	1457 B		GO	0:0031663	
##	-	value			
	1311 4.44284				
##	1312 5.95580				
##	1313 2.04203				
##	1314 2.76211				
##	1315 4.84655				
	1316 5.29114				
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	1319 2.34900				
	1320 3.60535				
##	1321 3.60535				
	1322 3.63754				
	1323 6.60498				
	1324 9.66358				
##	1325 2.18816	66e-05			

## 1326 2.555242e-05

regulation of ERK1 and ERK regulation of leukocyte cell-cell protein localization to extracellul regulation of innate immune glial cell a lation of inositol phosphate biosyntheti negative regulation of mast cell a positive regulation of phagocytosis, e positive regulation of membrane inv regulation of body flu regulation of myeloid leukocyte differ positive regulation of vasculature de glial cell : ulation of phosphatidylinositol 3-kinase T cell : cytokin response to defense response to regulation of phagocytosis, e regulation of microglial cell a regulation of lymphocyte positive regulation of ion sitive regulation of cytokine biosyntheti ERK1 and ERK peptide T cell lineage c tive regulation of activated T cell prol regulation of membrane inv regulation of myeloid leukocyte mediated regulation of ang regulation of interleukin-8 p cellular extr regulation of inflammatory regulation of antigen processing and pre cell junction di inflammatory

lipopolysaccharide-mediated signaling

- ## 3316 3.086309e-05
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- ## 1333 6.417224e-05
- ## 1334 6.438378e-05
- ## 1335 6.633435e-05
- ## 1555 0.055455e 00
- ## 1336 7.418961e-05
- ## 1337 7.924753e-05
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- ## 3872 7.205888e-04
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- ... 1004 0.0000000
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- ## 1393 1.367487e-03
- ## 3324 1.384455e-03
- ## 0024 1:0044000 00
- ## 1394 1.431051e-03
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- ## 3326 1.580520e-03
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- ## 1397 1.632240e-03
- ## 1398 1.632240e-03
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- ## 1399 1.635117e-03
- ## 1400 1.649303e-03
- ## 1401 1.654766e-03
- ## 1402 1.677685e-03
- ## 1403 1.677685e-03
- ## 1404 1.691385e-03 ## 3329 1.798052e-03
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- ## 1411 2.042202e-03
- ## 1412 2.042202e-03
- ## 1413 2.049387e-03 ## 3332 2.051128e-03
- ## 1414 2.052809e-03

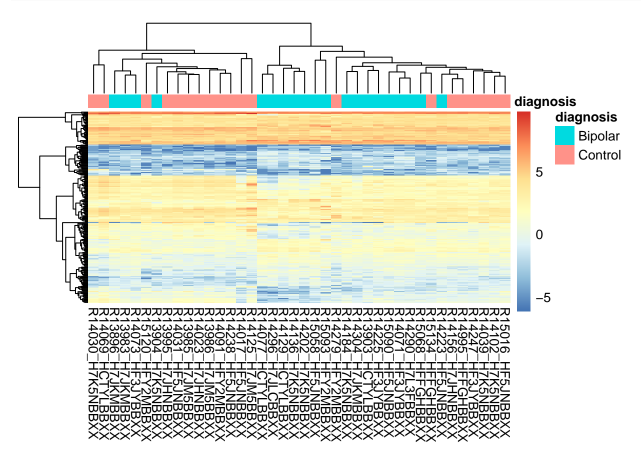
```
## 1415 2.108721e-03
## 1416 2.115847e-03
## 1417 2.180288e-03
## 1418 2.180288e-03
## 3333 2.213958e-03
## 1419 2.246151e-03
## 1420 2.260772e-03
## 1421 2.260772e-03
## 1422 2.292524e-03
## 1423 2.346125e-03
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## 1425 2.400666e-03
## 1426 2.419462e-03
## 1427 2.440725e-03
## 1428 2.440725e-03
## 1429 2.440725e-03
## 1430 2.440725e-03
## 1431 2.485301e-03
## 1432 2.602808e-03
## 1433 2.748562e-03
## 1434 2.757143e-03
## 1435 2.757143e-03
## 1436 2.757143e-03
## 3334 2.788894e-03
## 1437 2.872826e-03
## 1438 2.872826e-03
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## 1441 2.936322e-03
## 1442 3.000153e-03
## 1443 3.122514e-03
## 1444 3.132401e-03
## 1445 3.149398e-03
## 1446 3.338079e-03
## 1447 3.338079e-03
## 1448 3.338079e-03
## 1449 3.516285e-03
## 1450 3.629843e-03
## 1451 3.724031e-03
## 1452 3.724031e-03
## 1453 3.782205e-03
## 1454 3.836064e-03
## 1455 3.836064e-03
## 1456 3.902893e-03
## 1457 3.939125e-03
```

## 1.9 Visualize differentially expressed genes

Here we visualize DEGs with a heatmap.

```
exprs_heatmap = vGene$E[rownames(sigGene),]

df <- as.data.frame(colData(rse_gene)[,c("PrimaryDx")])</pre>
```



## pdf ## 3

## 2 Reproducibility

This analysis report was made possible thanks to:

- R (Kolde, 2019)
- BiocStyle (R Core Team, 2020)
- clusterProfiler (Allaire, Xie, McPherson, Luraschi, et al., 2020)

- devtools (Boettiger, 2019)
- edgeR
- here (Collado-Torres, Nellore, Kammers, Ellis, et al., 2017)
- jaffelab (Collado-Torres, Jaffe, and Burke, 2019)
- limma (Ellis, Collado-Torres, Jaffe, and Leek, 2018)
- knitcitations (Collado-Torres, Nellore, Kammers, Ellis, et al., 2020)
- org. Hs. eg. db (Frazee, Langmead, and Leek, 2011)
- pheatmap (Imada, Sanchez, Collado-Torres, Wilks, et al., 2020)
- RColorBrewer (McCarthy, Chen, and Smyth, 2012)
- recount
- rmarkdown
- SummarizedExperiment (Xie, Allaire, and Grolemund, 2018)
- voom (Yu, Wang, Han, and He, 2012)

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```
# Time spent creating this report:
diff(c(timestart, Sys.time()))
## Time difference of 1.453469 mins
# Date this report was generated
message(Sys.time())
## 2020-09-25 17:00:09
# Reproducibility info
options(width = 120)
devtools::session info()
## - Session info -----
##
   setting value
   version R version 4.0.2 Patched (2020-09-17 r79226)
## os
            Ubuntu 20.04.1 LTS
## system x86_64, linux-gnu
## ui
            X11
## language (EN)
## collate en_US.UTF-8
## ctype
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## tz
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## date
            2020-09-25
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                                     2017-09-09 [1] RSPM (R 4.0.0)
```

```
gridGraphics
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    Hmisc
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                                      2020-08-10 [1] RSPM (R 4.0.2)
##
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                                     2019-10-08 [2] RSPM (R 4.0.0)
##
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##
    httr
                            1.4.2
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                            1.2.5
##
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##
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                            1.7.1
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                            1.29
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    knitr
##
    lattice
                            0.20 - 41
                                     2020-04-02 [3] CRAN (R 4.0.2)
                            0.6 - 29
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##
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##
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##
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                            7.3 - 53
    Matrix
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    {\tt matrixStats}
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    org.Hs.eg.db
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##
                            2.0.3
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##
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##
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    qvalue
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##
    RColorBrewer
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##
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    Rcpp
##
   RCurl
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##
   readr
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## recount
                          * 1.15.1
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2019-04-03 [1] RSPM (R 4.0.0)
    RefManageR
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##
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##
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##
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##
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##
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                            1.5.3
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    stringr
                            1.4.0
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##
    SummarizedExperiment * 1.19.6
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##
    survival
                            3.2 - 3
                                     2020-06-13 [3] CRAN (R 4.0.2)
##
    testthat
                            2.3.2
                                     2020-03-02 [2] RSPM (R 4.0.0)
   tibble
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##
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                            1.1.2
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##
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##
  tidyselect
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  triebeard
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##
    tweenr
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##
    urltools
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##
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   {\tt VariantAnnotation}
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##
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##
    xfun
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  XML
##
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##
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##
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                                     2020-02-01 [2] RSPM (R 4.0.0)
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##
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    zlibbioc
##
## [1] /usr/local/lib/R/host-site-library
  [2] /usr/local/lib/R/site-library
## [3] /usr/local/lib/R/library
```