### main

July 10, 2021

### 1 Differential Expression with limma-voom pipeline

#### 1.1 Functions

#### 1.1.1 Simple functions

```
[2]: # Function from jaffelab github
     merge_rse_metrics <- function(rse) {</pre>
         stopifnot(is(rse, 'RangedSummarizedExperiment'))
         rse$overallMapRate = mapply(function(r, n) {
             sum(r*n)/sum(n)
         }, rse$overallMapRate, rse$numReads)
         rse$mitoRate = mapply(function(r, n) {
             sum(r*n)/sum(n)
         }, rse$mitoRate, rse$numMapped)
         rse$rRNA_rate = mapply(function(r, n) {
             sum(r*n)/sum(n)
         }, rse$rRNA_rate, rse$numMapped)
         rse$totalAssignedGene = mapply(function(r, n) {
             sum(r*n)/sum(n)
         }, rse$totalAssignedGene, rse$numMapped)
         rse$numMapped = sapply(rse$numMapped, sum)
         rse$numReads = sapply(rse$numReads, sum)
         rse$numUnmapped = sapply(rse$numUnmapped, sum)
         rse$mitoMapped = sapply(rse$mitoMapped, sum)
         rse$totalMapped = sapply(rse$totalMapped, sum)
         return(rse)
     }
```

```
save_volcanoPlot <- function(top, label, feature){</pre>
    pdf(file=paste0(feature, "/volcanoPlot_", label, ".pdf"), 8, 6)
    with(top, plot(logFC, -log10(P.Value), pch=20, cex=0.6))
    with(subset(top, adj.P.Val<=0.05), points(logFC, -log10(P.Value),</pre>
                                                pch=20, col='red', cex=0.6))
    with(subset(top, abs(logFC)>0.50), points(logFC, -log10(P.Value),
                                                pch=20, col='orange', cex=0.6))
    with(subset(top, adj.P.Val<=0.05 & abs(logFC)>0.50),
         points(logFC, -log10(P.Value), pch=20, col='green', cex=0.6))
    dev.off()
}
save_MAplot <- function(top, label, feature){</pre>
    pdf(file=paste0(feature, "/MAplot_", label, ".pdf"), 8, 6)
    with(top, plot(AveExpr, logFC, pch=20, cex=0.5))
    with(subset(top, adj.P.Val<0.05),</pre>
         points(AveExpr, logFC, col="red", pch=20, cex=0.5))
    dev.off()
}
extract_de <- function(contrast, label, efit, feature){</pre>
    top <- topTable(efit, coef=contrast, number=Inf, sort.by="P")</pre>
    top <- top[order(top$P.Value), ]</pre>
    top.fdr <- top %>% filter(adj.P.Val<=0.05)</pre>
    print(paste("Comparison for:", label))
    print(paste('There are:', dim(top.fdr)[1], 'DE features!'))
    fwrite(top,
           file=pasteO(feature, "/diffExpr_", label, "_full.txt"),
           sep='\t', row.names=TRUE)
    fwrite(top.fdr,
           file=pasteO(feature, "/diffExpr_", label, "_FDR05.txt"),
           sep='\t', row.names=TRUE)
    save volcanoPlot(top, label, feature)
    save_MAplot(top, label, feature)
}
```

### 1.1.2 Cached functions

```
[3]: prep_data <- function(feature){
    ancestry = "../../../input/ancestry_structure/structure.
    out_ancestry_proportion_raceDemo_compare"
    counts_lt = list("genes"="../../../input/counts/_m/
    ohippo_brainseq_phase2_hg38_rseGene_merged_n447.rda",
```

```
"transcripts"="../../input/counts/_m/

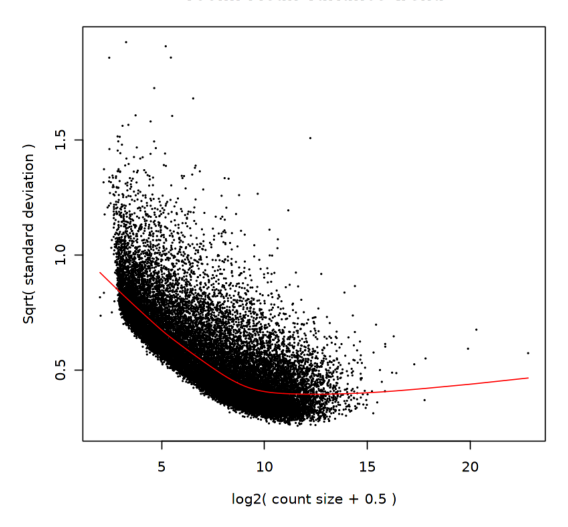
→hippo_brainseq_phase2_hg38_rseTx_merged_n447.rda",
                    "exons"="../../input/counts/ m/
→hippo_brainseq_phase2_hg38_rseExon_merged_n447.rda",
                    "junctions"="../../input/counts/_m/
→hippo_brainseq_phase2_hg38_rseJxn_merged_n447.rda")
   tx_file = "../../input/counts/_m/transcripts_counts/hippo_counts.txt"
  load(counts lt[[feature]])
  if(exists("rse_gene")){
       rse_df = rse_gene
  } else if (exists("rse tx")){
       counts <- fread(tx_file) %>% column_to_rownames("transcript_id") %>%
           select(colnames(rse_tx)) %>% as.matrix
       annot <- fread(tx_file) %>% column to rownames("transcript_id") %>%
           select(-starts_with("R"))
       rse_df <- SummarizedExperiment(assays=SimpleList(counts=counts),</pre>
                                       rowData=annot,
                                       colData=colData(rse_tx)) %>%
           as("RangedSummarizedExperiment")
  } else if (exists("rse_exon")){
       rse_df = rse_exon
  } else {
       rse_df = rse_jxn
  keepIndex = which((rse_df$Dx %in% c("Control")) &
                     rse_df$Age > 17 &
                     rse_df$Race %in% c("AA", "CAUC"))
  rse_df = rse_df[, keepIndex]
  rse df$Sex <- factor(rse df$Sex)</pre>
  rse_df <- merge_rse_metrics(rse_df)</pre>
   colData(rse_df)$RIN = sapply(colData(rse_df)$RIN,"[",1)
  rownames(colData(rse_df)) <- sapply(strsplit(rownames(colData(rse_df)),__
\rightarrow"_"), "[", 1)
  pheno = colData(rse_df) %>% as.data.frame %>%
       inner join(fread(ancestry), by=c("BrNum"="id", "Race"="group"))
   # Generate DGE list
  x <- DGEList(counts=assays(rse_df)$counts[, pheno$RNum],</pre>
                genes=rowData(rse df),
                samples=pheno)
   # Filter by expression
  design0 <- model.matrix(~Race, data=x$samples)</pre>
  keep.x <- filterByExpr(x, design=design0)</pre>
  x <- x[keep.x, , keep.lib.sizes=FALSE]</pre>
  print(paste('There are:', sum(keep.x), 'features left!', sep=' '))
   # Normalize library size
  x <- calcNormFactors(x, method="TMM")</pre>
  return(x)
```

```
}
memo_prepData <- memoise(prep_data)</pre>
qSV_model <- function(feature){</pre>
    x <- memo_prepData(feature)</pre>
    # Design matrix
    mod = model.matrix(~Eur + Sex + Age + mitoRate + rRNA_rate +
                        totalAssignedGene + overallMapRate, data = x$samples)
    colnames(mod) <- gsub("SexM", "Male", colnames(mod))</pre>
    colnames(mod) <- gsub("Eur", "EA", colnames(mod))</pre>
    colnames(mod) <- gsub("\\(Intercept\\)", "Intercept", colnames(mod))</pre>
    # Load qSV
    qsv_file = "../../input/phenotypes/_m/qSV_hippo.csv"
    modQsva <- mod %>% as.data.frame %>% rownames_to_column() %>%
        inner_join(fread(qsv_file), by=c("rowname"="V1")) %>%
        rename_all(list(~str_replace_all(., 'PC', 'qPC'))) %>%
        column_to_rownames("rowname") %>% as.matrix
    return(modQsva)
}
memo_qsvModel <- memoise(qSV_model)</pre>
get voom <- function(feature){</pre>
    ### Preform voom
    x <- memo_prepData(feature)</pre>
    modQsva <- memo_qsvModel(feature)</pre>
    v <- voom(x[, rownames(modQsva)], modQsva, plot=TRUE)</pre>
    return(v)
}
memo_voom <- memoise(get_voom)</pre>
cal_res <- function(feature){</pre>
    ### Calculate residuals
    v <- memo_voom(feature)</pre>
    null_model <- v$design %>% as.data.frame %>% select(-c("EA")) %>% as.matrix
    fit_res <- lmFit(v, design=null_model)</pre>
    res = v$E - ( fit res$coefficients %*% t(null model) )
    res_sd = apply(res, 1, sd)
    res_mean = apply(res, 1, mean)
    res_norm = (res - res_mean) / res_sd
    write.table(res_norm, file=paste0(feature, '/residualized_expression.tsv'),
                 sep="\t", quote=FALSE)
}
memo_res <- memoise(cal_res)</pre>
```

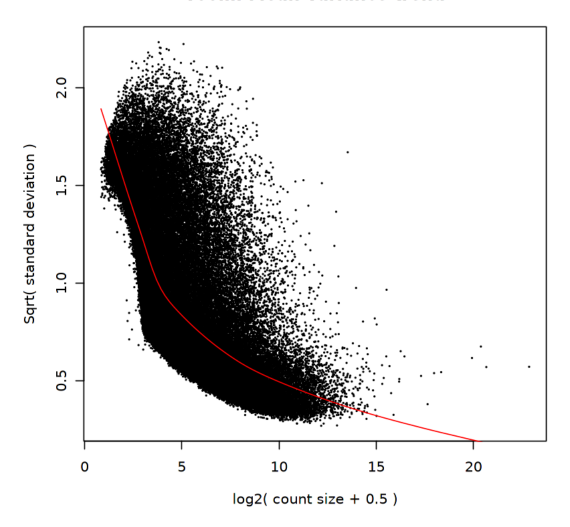
### 1.2 Differential Expression Analysis

```
[4]: for(feature in c('genes', 'transcripts', 'junctions', 'exons')){
    dir.create(feature)
    # Preform voom
    v <- memo_voom(feature)
    save(v, file=paste0(feature,'/voomSVA.RData'))
    # Fit model and apply eBayes
    efit = memo_efit(feature)
    # Save differential expression
    extract_de(1, "EAvsAA", efit, feature)
    # Calculate residuals
    memo_res(feature)
}</pre>
```

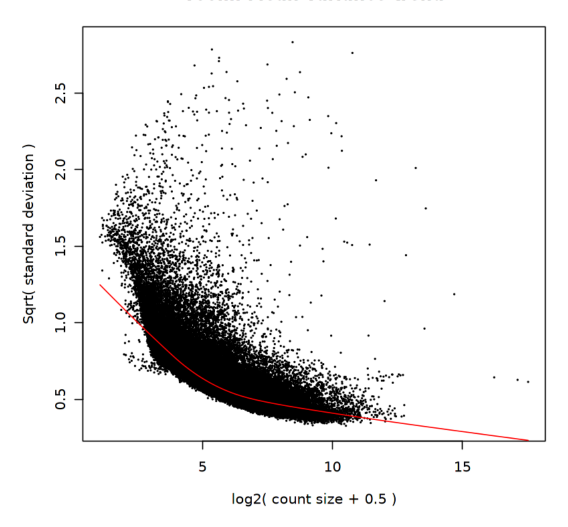
[1] "There are: 22269 features left!"
[1] "Comparison for: EAvsAA"
[1] "There are: 2956 DE features!"
[1] "There are: 75309 features left!"
[1] "Comparison for: EAvsAA"
[1] "There are: 3118 DE features!"



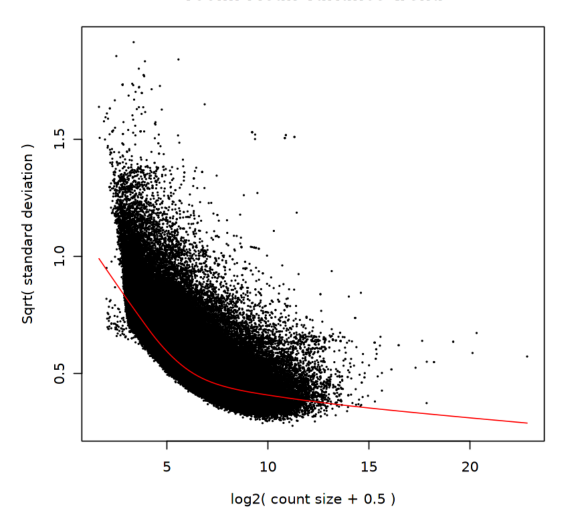
- [1] "There are: 136285 features left!"
- [1] "Comparison for: EAvsAA"
- [1] "There are: 4052 DE features!"



[1] "There are: 327876 features left!"



- [1] "Comparison for: EAvsAA"
- [1] "There are: 15448 DE features!"



## 1.3 Reproducibility Information

```
[5]: Sys.time()
    proc.time()
    options(width = 120)
    sessioninfo::session_info()

[1] "2021-07-10 12:17:13 EDT"

    user system elapsed
    1611.023 90.392 1304.695

    Session info
    setting value
```

version R version 4.0.3 (2020-10-10)

os Arch Linux

system x86\_64, linux-gnu

ui X11 language (EN)

collate en\_US.UTF-8
ctype en\_US.UTF-8
tz America/New\_York

date 2021-07-10

### Packages package

package	*	version	date	lib	source
annotate		1.68.0	2020-10-27	[1]	Bioconductor
AnnotationDbi		1.52.0	2020-10-27	[1]	Bioconductor
assertthat		0.2.1	2019-03-21	[1]	CRAN (R 4.0.2)
backports		1.2.1	2020-12-09	[1]	CRAN (R 4.0.2)
base64enc		0.1-3	2015-07-28	[1]	CRAN (R 4.0.2)
Biobase	*	2.50.0	2020-10-27	[1]	Bioconductor
BiocGenerics	*	0.36.1	2021-04-16	[1]	Bioconductor
BiocParallel	*	1.24.1	2020-11-06	[1]	Bioconductor
bit		4.0.4	2020-08-04	[1]	CRAN (R 4.0.2)
bit64		4.0.5	2020-08-30	[1]	CRAN (R 4.0.2)
bitops		1.0-7	2021-04-24	[1]	CRAN (R 4.0.3)
blob		1.2.1	2020-01-20	[1]	CRAN (R 4.0.2)
broom		0.7.8	2021-06-24	[1]	CRAN (R 4.0.3)
cachem		1.0.5	2021-05-15	[1]	CRAN (R 4.0.3)
Cairo		1.5-12.2	2020-07-07	[1]	CRAN (R 4.0.2)
cellranger		1.1.0	2016-07-27	[1]	CRAN (R 4.0.2)
cli		3.0.0	2021-06-30	[1]	CRAN (R 4.0.3)
colorspace		2.0-2	2021-06-24	[1]	CRAN (R 4.0.3)
crayon		1.4.1	2021-02-08	[1]	CRAN (R 4.0.3)
data.table	*	1.14.0	2021-02-21	[1]	CRAN (R 4.0.3)
DBI		1.1.1	2021-01-15	[1]	CRAN (R 4.0.2)
dbplyr		2.1.1	2021-04-06	[1]	CRAN (R 4.0.3)
DelayedArray		0.16.3	2021-03-24	[1]	Bioconductor
digest		0.6.27	2020-10-24	[1]	CRAN (R 4.0.2)
dplyr	*	1.0.7	2021-06-18	[1]	CRAN (R 4.0.3)
edgeR	*	3.32.1	2021-01-14	[1]	Bioconductor
ellipsis		0.3.2	2021-04-29	[1]	CRAN (R 4.0.3)
evaluate		0.14	2019-05-28	[1]	CRAN (R 4.0.2)
fansi		0.5.0	2021-05-25	[1]	CRAN (R 4.0.3)
fastmap		1.1.0	2021-01-25	[1]	CRAN (R 4.0.2)
forcats	*	0.5.1	2021-01-27	[1]	CRAN (R 4.0.2)
fs		1.5.0	2020-07-31	[1]	CRAN (R 4.0.2)
genefilter	*	1.72.1	2021-01-21	[1]	Bioconductor
generics		0.1.0	2020-10-31	[1]	CRAN (R 4.0.2)
GenomeInfoDb	*	1.26.7	2021-04-08	[1]	Bioconductor
${\tt GenomeInfoDbData}$		1.2.4	2021-02-02	[1]	Bioconductor

GenomicRanges	*	1.42.0	2020-10-27	[1]	Bioconductor
ggplot2		3.3.5	2021-06-25	[1]	CRAN (R 4.0.3)
glue	.,.	1.4.2	2020-08-27	[1]	CRAN (R 4.0.2)
gtable		0.3.0	2019-03-25	[1]	CRAN (R 4.0.2)
haven		2.4.1	2019 03 23	[1]	CRAN (R 4.0.3)
hms		1.1.0	2021 04 23		CRAN (R 4.0.3)
htmltools		0.5.1.1	2021-01-22		CRAN (R 4.0.2)
httr		1.4.2	2021-01-22		CRAN (R 4.0.2)
IRanges	Ψ.	2.24.1			Bioconductor
IRdisplay	71	1.0	2020 12 12 2021-01-20	[1]	CRAN (R 4.0.2)
IRkernel		1.2	2021 01 20	[1]	CRAN (R 4.0.3)
jsonlite		1.7.2	2021 03 11		CRAN (R 4.0.2)
lattice		0.20-41	2020-12-09		CRAN (R 4.0.2)
lifecycle		1.0.0	2020-04-02	[1]	CRAN (R 4.0.3)
limma	Ψ.	3.46.0	2021-02-13		Bioconductor
locfit	•	1.5-9.4	2020-10-27		CRAN (R 4.0.2)
lubridate		1.7.10	2020-03-25		CRAN (R 4.0.2) CRAN (R 4.0.3)
magrittr		2.0.1	2020-11-17		CRAN (R 4.0.2) CRAN (R 4.0.3)
Matrix MatrixGenerics	.1.	1.3-4 1.2.1	2021-06-01	[1] [1]	•
	*		2021-01-30	[1]	Bioconductor CRAN (R 4.0.3)
matrixStats		0.59.0	2021-06-01		•
memoise	*	2.0.0	2021-01-26	[1]	CRAN (R 4.0.2)
mgcv	*	1.8-33	2020-08-27		CRAN (R 4.0.3)
modelr		0.1.8	2020-05-19		
munsell		0.5.0	2018-06-12		CRAN (R 4.0.2)
nlme	*	3.1-152	2021-02-04	[1]	CRAN (R 4.0.3)
pbdZMQ		0.3-5	2021-02-10	[1]	CRAN (R 4.0.3)
pillar		1.6.1	2021-05-16	[1]	CRAN (R 4.0.3)
pkgconfig		2.0.3	2019-09-22	[1]	CRAN (R 4.0.2)
purrr	*	0.3.4	2020-04-17	[1]	CRAN (R 4.0.2)
R6		2.5.0	2020-10-28		CRAN (R 4.0.2)
Rcpp		1.0.7	2021-07-07		CRAN (R 4.0.3)
RCurl			2021-03-16	[1]	CRAN (R 4.0.3)
readr	*	1.4.0			CRAN (R 4.0.2)
readxl		1.3.1	2019-03-13		
repr		1.1.3			CRAN (R 4.0.2)
reprex		2.0.0	2021-04-02		
rlang		0.4.11	2021-04-30		
RSQLite		2.2.7	2021-04-22		
rstudioapi		0.13	2020-11-12		
rvest		1.0.0	2021-03-09		
S4Vectors	*	0.28.1	2020-12-09		
scales		1.1.1			CRAN (R 4.0.2)
sessioninfo		1.1.1			CRAN (R 4.0.2)
stringi		1.6.2	2021-05-17		
stringr		1.4.0	2019-02-10	[1]	
SummarizedExperiment	*				Bioconductor
survival		3.2-7	2020-09-28	[2]	CRAN (R 4.0.3)

```
* 3.38.0
                                2020-10-27 [1] Bioconductor
sva
tibble
                     * 3.1.2
                                2021-05-16 [1] CRAN (R 4.0.3)
                     * 1.1.3
                                2021-03-03 [1] CRAN (R 4.0.3)
tidyr
tidyselect
                       1.1.1
                                2021-04-30 [1] CRAN (R 4.0.3)
tidyverse
                     * 1.3.1
                                2021-04-15 [1] CRAN (R 4.0.3)
                                2021-03-12 [1] CRAN (R 4.0.3)
utf8
                       1.2.1
uuid
                       0.1-4
                                2020-02-26 [1] CRAN (R 4.0.2)
                                2021-04-29 [1] CRAN (R 4.0.3)
vctrs
                       0.3.8
withr
                       2.4.2
                                2021-04-18 [1] CRAN (R 4.0.3)
XML
                       3.99-0.6 2021-03-16 [1] CRAN (R 4.0.3)
                                2020-04-23 [1] CRAN (R 4.0.2)
xm12
                       1.3.2
xtable
                       1.8-4
                                2019-04-21 [1] CRAN (R 4.0.2)
                                2020-10-27 [1] Bioconductor
XVector
                       0.30.0
                       1.36.0
                                2020-10-27 [1] Bioconductor
zlibbioc
```

- [1] /home/jbenja13/R/x86\_64-pc-linux-gnu-library/4.0
- [2] /usr/lib/R/library