# DE analysis - Day1

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#### Overview

- Principal Investigator: Beth Overmoyer
- Experiment: RNAseq\_analysis\_of\_inflammatory\_breast\_cancer\_hbc04141
- study 6 was excluded because if low read depth in 3373-3
- $\verb| https://www.bioconductor.org/packages/release/bioc/vignettes/DEGreport/inst/doc/DEGreport. \\ \verb| html|$
- AnnotationHub. We use ensembl version matching bebio pipeline v94.
- HBC materials
- HBC materials functional analysis
- $\bullet \ \ http://bioconductor.org/packages/release/bioc/vignettes/DESeq2/inst/doc/DESeq2.html$
- this is DE for Day1 samples

#### Checking to see that the transcript to gene mapping is correct

When you have annotations that are from a different source from your reference you can run into problems (i.e lose genes). Some checks you can do before proceeding:

- 1. Look at the dimensions of your count matrix. Do you have ~20k genes present? dim(txi\$counts)
- 2. When running tximport() you will get a message in your console. If you see something like transcripts missing from tx2gene start troubleshooting.

```
dim(txi$counts)
## [1] 58735 20
```

## Sanity check that metadata matches your expression

It is always a good idea to check if:

- 1. Do you have expression data for all samples listed in your metadata?
- 2. Are the samples in your expression data in the same order as your metadata?

```
### Check that sample names match in both files
all(colnames(txi$counts) %in% rownames(meta))

### [1] TRUE

### Check that sample names match in both files
all(colnames(txi$counts) %in% rownames(meta))

## [1] TRUE

### Check that all samples are in the same order

meta <- meta[colnames(txi$counts),]
all(colnames(txi$counts) == rownames(meta))</pre>
```

#### Run DESeq2

```
estimating size factors
estimating dispersions
gene-wise dispersion estimates
mean-dispersion relationship
final dispersion estimates
fitting model and testing
```

- Estimating size factors and count normalization
- Gene-wise dispersions
- Mean-dispersion(variance) relationship and the Negative Binomial Model
- Model fitting and hypothesis testing

#### Wald test

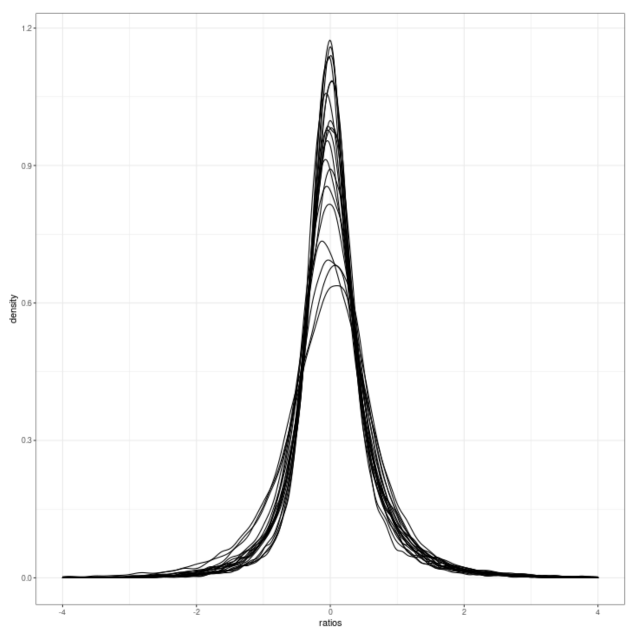
Here we subset protein coding genes.

```
## Create DESeq2Dataset object
dds_file <- "data/dds.day1.RDS"</pre>
meta$treatment <- as.factor(meta$treatment)</pre>
meta$response <- as.factor(meta$response)</pre>
meta$er <- as.factor(meta$er)</pre>
meta$date_of <- as.factor(meta$date_of)</pre>
meta$tumor_percentage <- as.factor(meta$tumor_percentage)</pre>
meta$tumor_percentage_high <- as.factor(meta$tumor_percentage_high)</pre>
if (remove_cases_2_19){
  non_responders <- meta %>% dplyr::filter(study_id %in% c(2, 19)) %>% row.names()
}
if (!rebuild_rds & file.exists(dds_file)){
    dds <- readRDS(dds_file)</pre>
}else{
    dds <- DESeqDataSetFromTximport(txi,</pre>
                                   colData = meta,
                                   design = ~response)
    if (remove_cases_2_19){
        dds <- dds[,!colnames(dds) %in% non_responders]</pre>
    design(dds) <- formula(~response + er + tumor_percentage_high + date_of)</pre>
    # subset protein-coding genes
    pc_genes <- intersect(protein_coding_genes$ensembl_gene_id, row.names(dds))</pre>
    dds <- dds[pc genes,]</pre>
    # 100 reads / 20 samples
    keep <- rowSums(counts(dds)) >= 100
    dds <- dds[keep,]</pre>
    # Run DESeq2
    dds <- DESeq(dds)
    saveRDS(dds, dds_file)
```

# DEGreport QC

## Size factor QC - samples 1-20

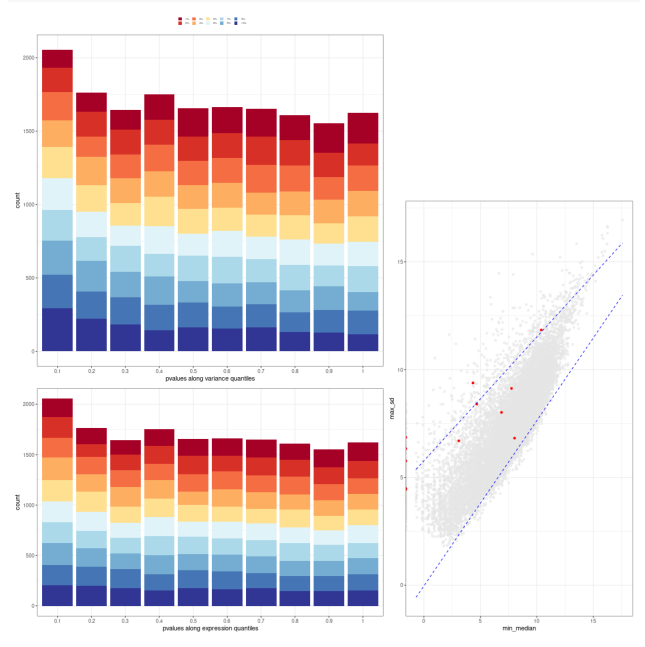
```
counts <- counts(dds, normalized = TRUE)
design <- as.data.frame(colData(dds))
degCheckFactors(counts[, 1:20])</pre>
```



## Mean-Variance QC plots

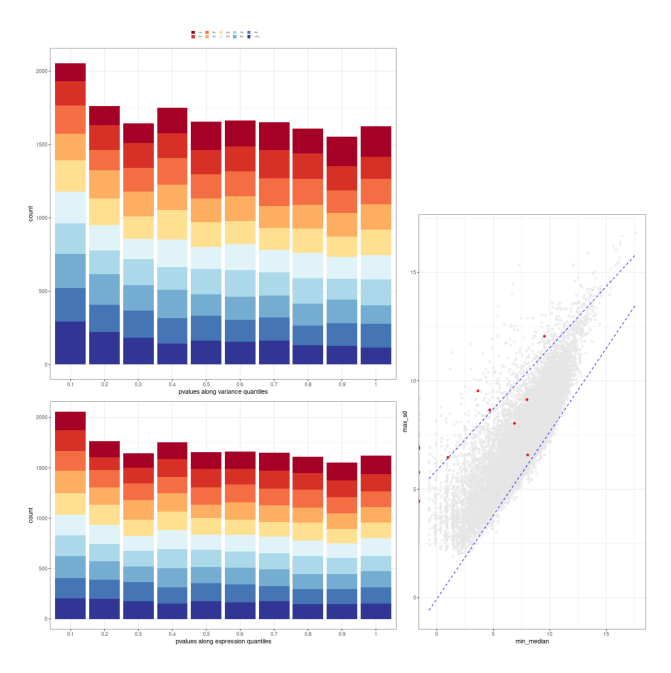
### response

```
res <- results(dds)
degQC(counts, design[["response"]], pvalue = res[["pvalue"]])</pre>
```



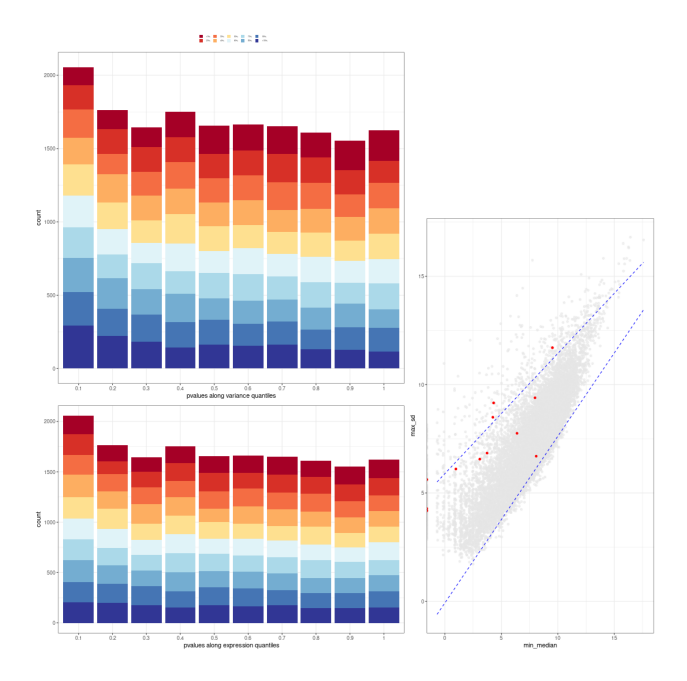
#### $\mathbf{E}\mathbf{R}$

```
degQC(counts, design[["er"]], pvalue = res[["pvalue"]])
```



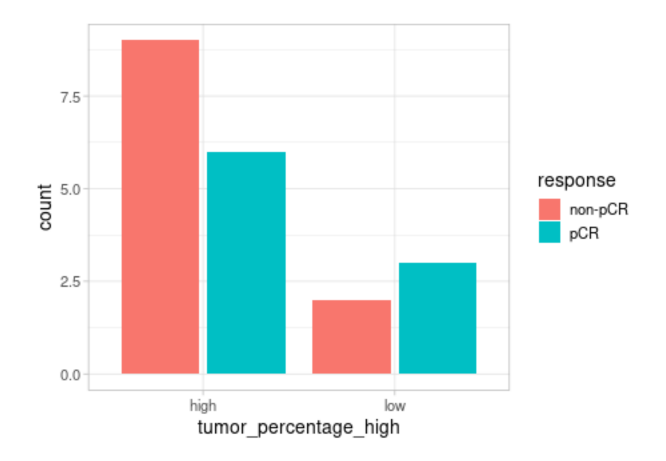
## $tumor\_percentage\_high$

degQC(counts, design[["tumor\_percentage\_high"]], pvalue = res[["pvalue"]])



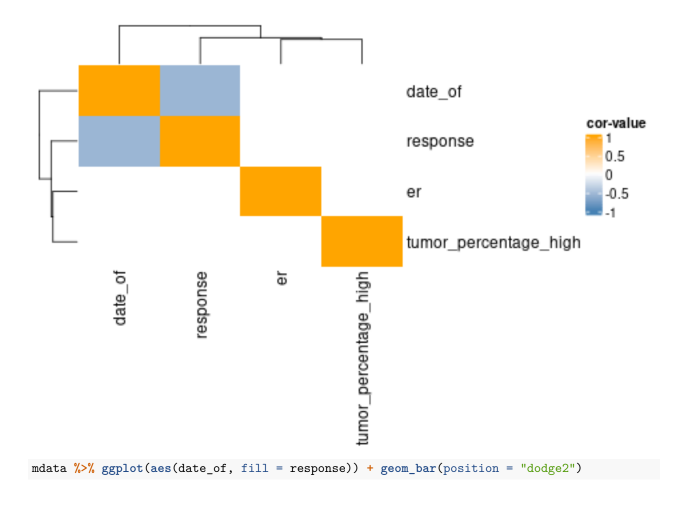
#### Covariates effect on count data

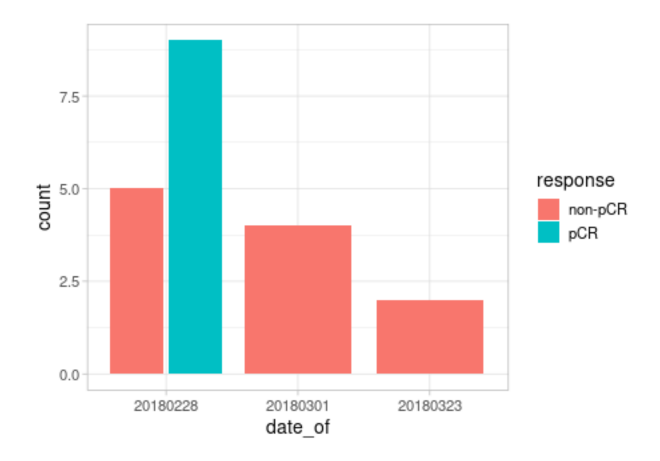
```
mdata <- colData(dds) %>% as.data.frame() %>%
   dplyr::select(response, er, date_of, tumor_percentage_high)
#resCov <- degCovariates(log2(counts(dds)+0.5), mdata)
mdata %>% ggplot(aes(tumor_percentage_high, fill = response)) + geom_bar(position = "dodge2")
```



## Covariates correlation with metrics

cor <- degCorCov(mdata)</pre>





### Sample-level QC analysis

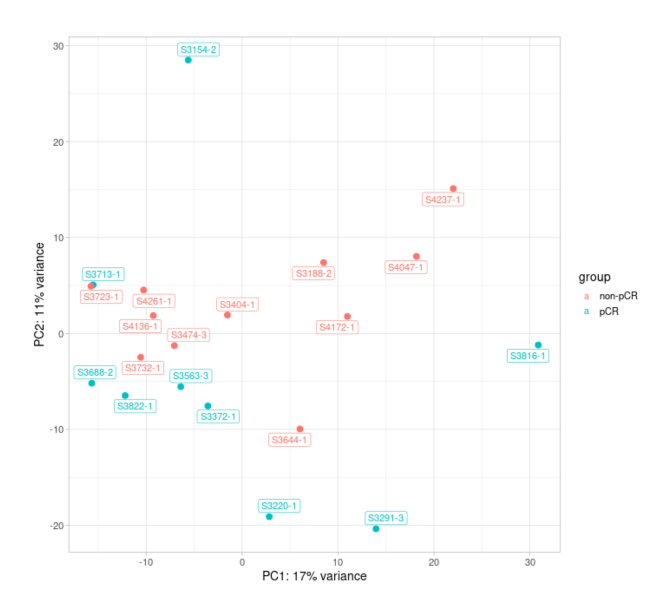
```
### Transform counts for data visualization (unsupervised analysis)
rld_file <- "data/rld.day1.RDS"
if (!rebuild_rds &file.exists(rld_file)){
    rld <- readRDS(rld_file)
}else{
    rld <- rlog(dds, blind = TRUE)
        saveRDS(rld, rld_file)
}
class(rld) # what type of object is this

## [1] "DESeqTransform"
## attr(,"package")
## [1] "DESeq2"

# we also need just a matrix of transformed counts
rld_mat <- assay(rld)</pre>
```

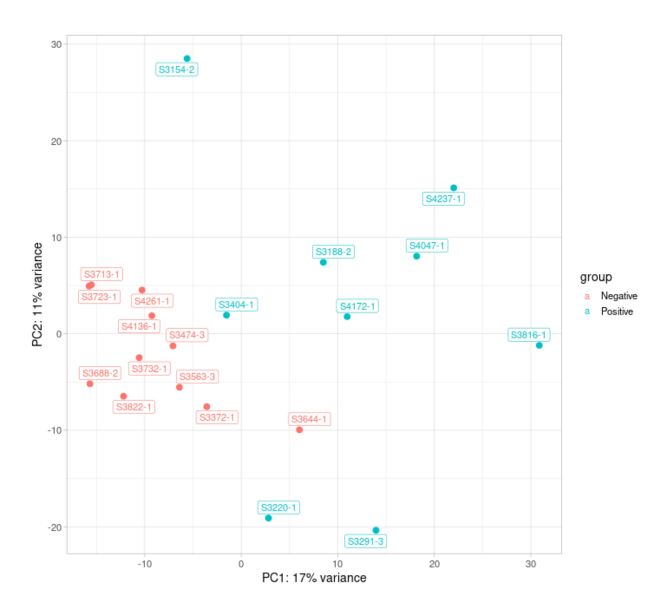
#### PCA - response

```
# Use the DESeq2 function
plotPCA(rld, intgroup = c("response")) + geom_label_repel(aes(label = name))
```



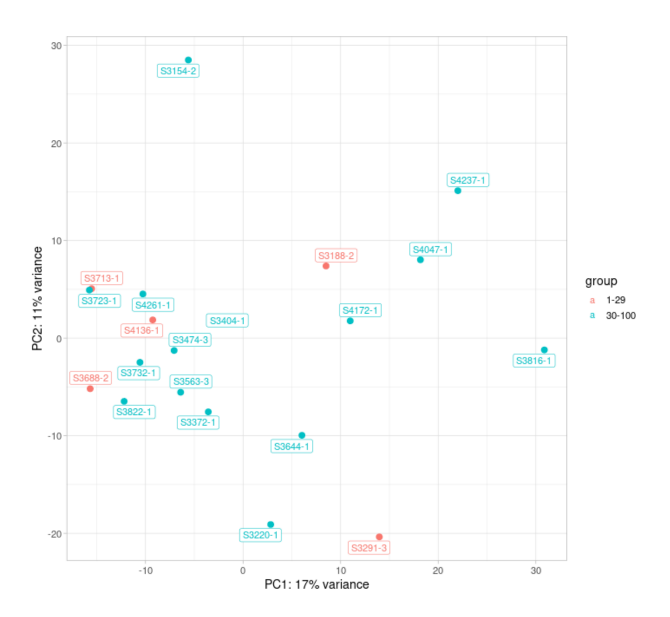
#### PCA - ER

```
# Use the DESeq2 function
plotPCA(rld, intgroup = c("er")) + geom_label_repel(aes(label = name))
```



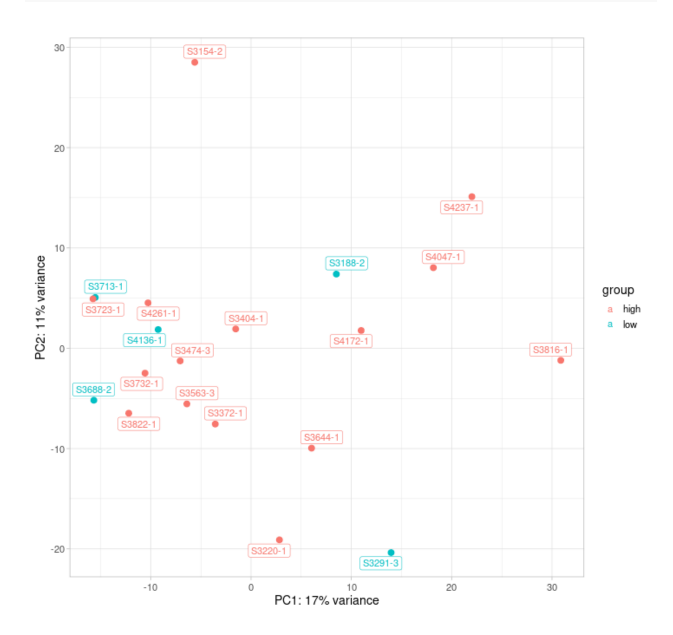
#### PCA - tumor\_percentage

```
# Use the DESeq2 function
plotPCA(rld, intgroup = c("tumor_percentage")) + geom_label_repel(aes(label = name))
```



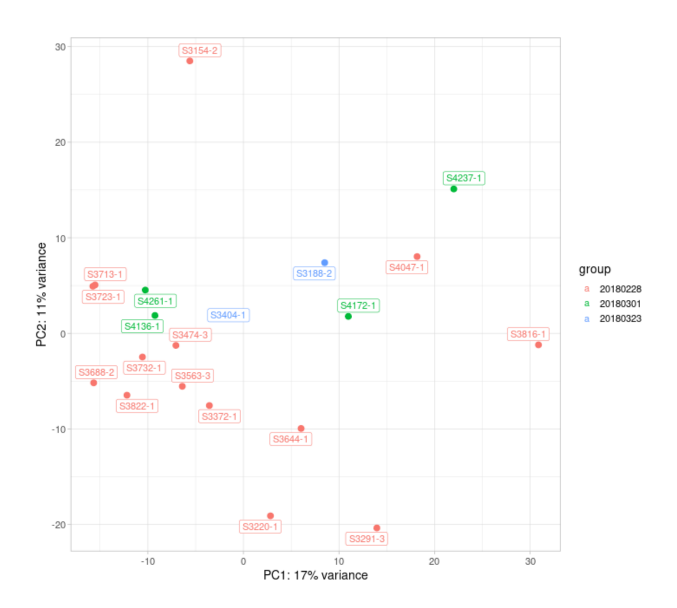
### $PCA - tumor\_percentage\_high$

```
# Use the DESeq2 function
plotPCA(rld, intgroup = c("tumor_percentage_high")) + geom_label_repel(aes(label = name))
```



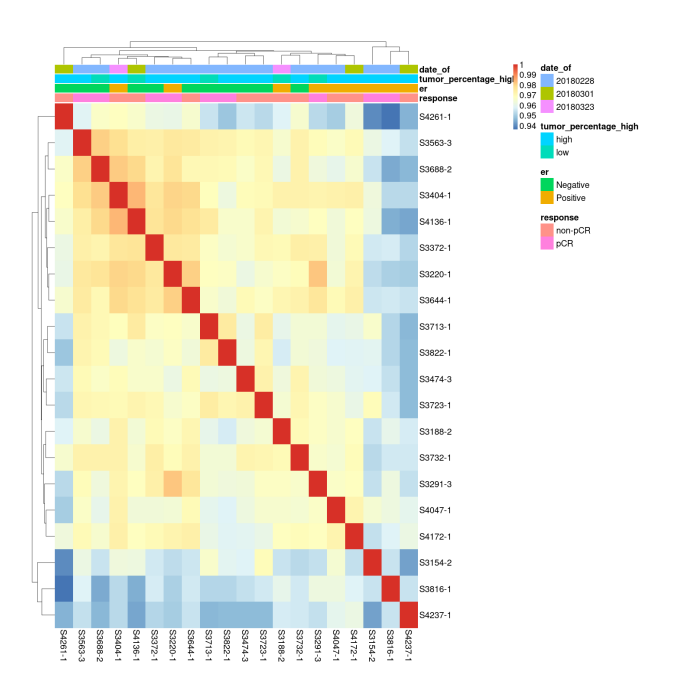
#### PCA - date of

```
# Use the DESeq2 function
plotPCA(rld, intgroup = c("date_of")) + geom_label_repel(aes(label = name))
```

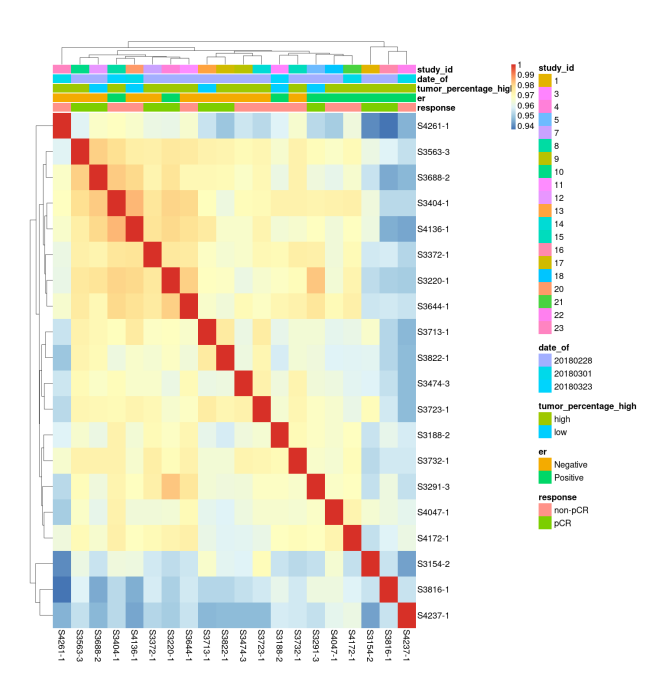


## Inter-correlation analysis

#### Without study\_id



### With study\_id



Response pCR vs non-pCR for Day 1- see Table9

ER: Positive vs Negative for Day1 - Table 10

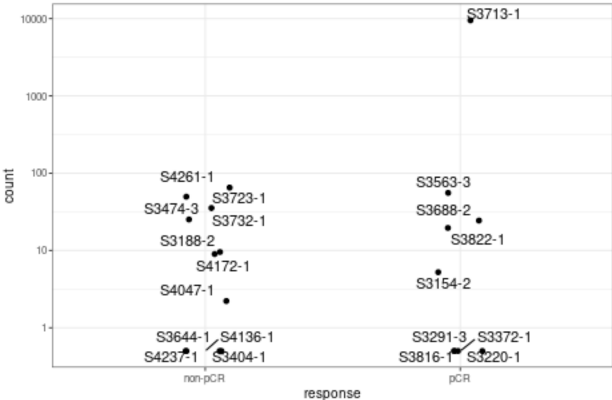
tumor\_percentage\_high: High vs Low for Day1- Table 11

date of: 20180323 vs 20180228 - for Day1: Table 12

#### Visualization

 $Gene\ example$ 





```
# Add a column for significant genes
resResponse_tb_vis <- resResponse_tb %>% mutate(threshold = padj < 0.01)
resResponse_tb_vis$symbol <- ifelse((abs(resResponse_tb_vis$log2FoldChange) > 1.5),
                                resResponse_tb_vis$symbol, NA)
resResponse_tb_vis$symbol <- ifelse(resResponse_tb_vis$threshold,</pre>
                                    resResponse_tb_vis$symbol, NA)
ggplot(resResponse_tb_vis,
       aes(log2FoldChange, -log10(padj), label = symbol)) +
  geom_point(aes(x = log2FoldChange, y = -log10(padj), colour = threshold)) +
  ggtitle("Response pCR vs non-pCR") +
  xlab("log2 fold change") +
  ylab("-log10 adjusted p-value") +
  scale_x_continuous(limits = c(-10,10)) +
  scale_y_continuous(limits = c(0, 5))+
  theme(legend.position = "none",
        plot.title = element_text(size = rel(1.5), hjust = 0.5),
        axis.title = element_text(size = rel(1.25)),
        panel.grid.major = element_blank(),
        panel.grid.minor = element_blank(),
        panel.border = element_blank(),
        panel.background = element_blank()) +
  geom_text_repel(aes(label = symbol))
```

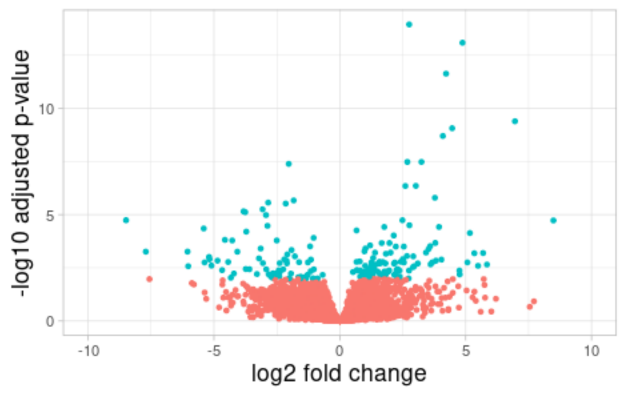
# Response pCR vs non-pCR



```
# Add a column for significant genes
resER_tb <- resER_tb %>% mutate(threshold = padj < 0.01)

ggplot(resER_tb) +
    geom_point(aes(x = log2FoldChange, y = -log10(padj), colour = threshold)) +
    ggtitle("ER: Positive vs Negative") +
    xlab("log2 fold change") +
    ylab("-log10 adjusted p-value") +
    scale_x_continuous(limits = c(-10,10)) +
    theme(legend.position = "none",
        plot.title = element_text(size = rel(1.5), hjust = 0.5),
        axis.title = element_text(size = rel(1.25)))</pre>
```

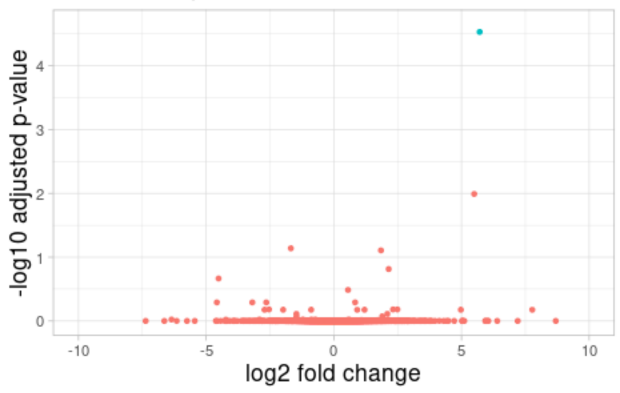
# ER: Positive vs Negative



```
# Add a column for significant genes
resTP_tb <- resTP_tb %>% mutate(threshold = padj < 0.01)

ggplot(resTP_tb) +
   geom_point(aes(x = log2FoldChange, y = -log10(padj), colour = threshold)) +
   ggtitle("Tumor_percentage_high: High vs Low") +
   xlab("log2 fold change") +
   ylab("-log10 adjusted p-value") +
   scale_x_continuous(limits = c(-10,10)) +
   theme(legend.position = "none",
        plot.title = element_text(size = rel(1.5), hjust = 0.5),
        axis.title = element_text(size = rel(1.25)))</pre>
```

# Tumor\_percentage\_high: High vs Low



```
# Add a column for significant genes
resDO_tb <- resDO_tb %>% mutate(threshold = padj < 0.01)

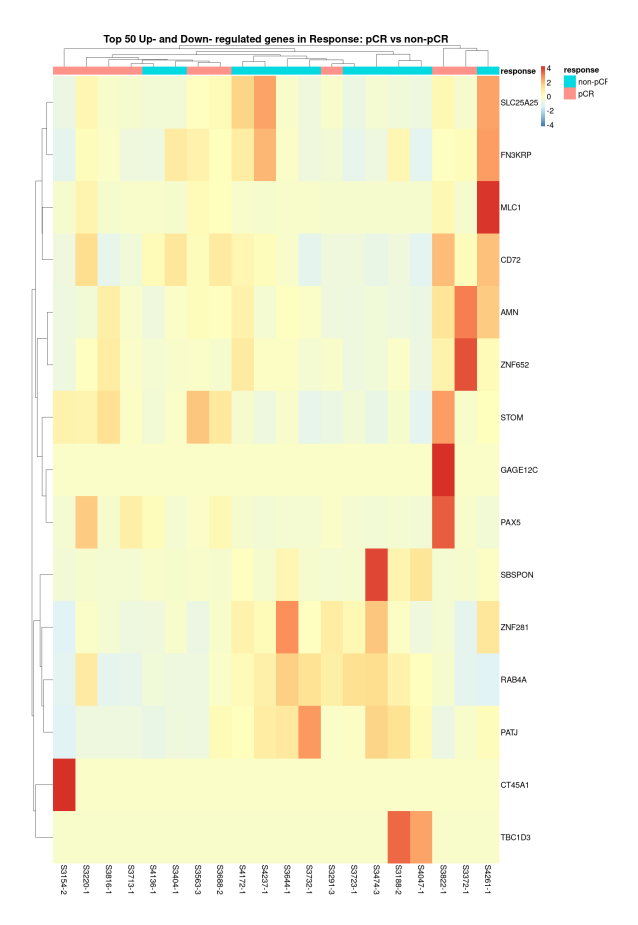
ggplot(resDO_tb) +
   geom_point(aes(x = log2FoldChange, y = -log10(padj), colour = threshold)) +
   ggtitle("Dafe of: 20180323 vs 20180228") +
   xlab("log2 fold change") +
   ylab("-log10 adjusted p-value") +
   scale_x_continuous(limits = c(-10,10)) +
   theme(legend.position = "none",
        plot.title = element_text(size = rel(1.5), hjust = 0.5),
        axis.title = element_text(size = rel(1.25)))</pre>
```

# Dafe of: 20180323 vs 20180228

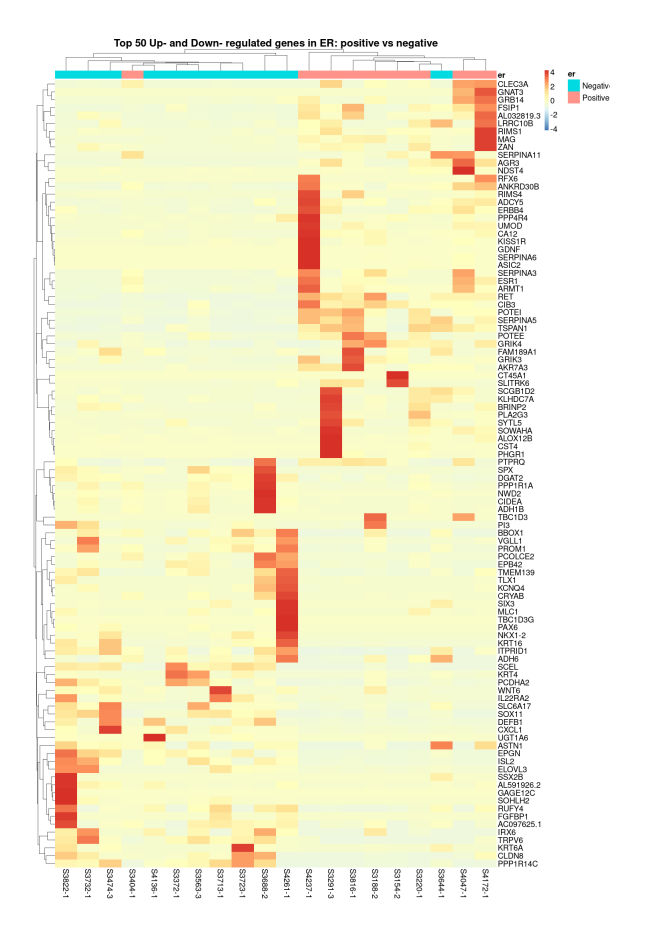


#### Heatmaps

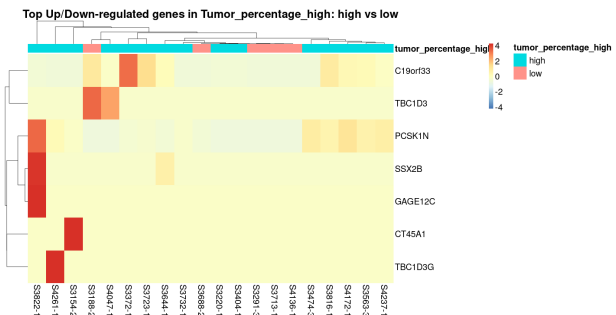
```
# Create a matrix of normalized expression
sig_up <- resResponse_tb_significant %>% arrange(-log2FoldChange) %>% head(50) %>% pull(gene)
sig_down <- resResponse_tb_significant %>% arrange(log2FoldChange) %>% head(50) %>% pull(gene)
sig <- c(sig_up, sig_down)</pre>
row_annotation <- gene_symbol %>%
                    as_tibble() %>%
                    dplyr::filter(gene_id %in% sig)
plotmat <- txi$abundance[c(sig_up, sig_down),] %>% as.data.frame() %>%
          rownames_to_column(var = "ensembl_gene_id") %>%
          left_join(gene_symbol, by = c("ensembl_gene_id" = "gene_id")) %>%
          drop_na(symbol)
plotmat$ensembl_gene_id <- NULL</pre>
plotmat <- plotmat %>% column_to_rownames(var = "symbol") %>% as.matrix()
# Color palette
heat.colors <- brewer.pal(6, "YlOrRd")</pre>
# Plot heatmap
pheatmap(plotmat,
         scale = "row",
         show_rownames = TRUE,
         border = FALSE,
         annotation = meta[, c("response"), drop = FALSE],
         main = "Top 50 Up- and Down- regulated genes in Response: pCR vs non-pCR",
         fontsize = 20)
```



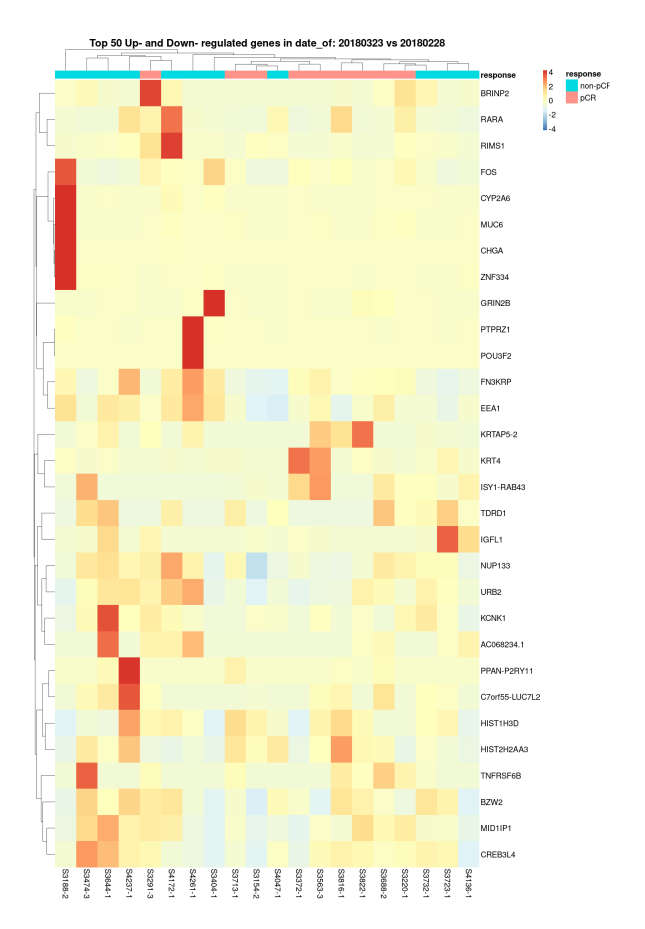
```
# Create a matrix of normalized expression
sig_up <- resER_tb_significant %>% arrange(-log2FoldChange) %>% head(50) %>% pull(gene)
sig_down <- resER_tb_significant %>% arrange(log2FoldChange) %>% head(50) %>% pull(gene)
sig <- c(sig_up, sig_down)</pre>
row_annotation <- gene_symbol %>%
                    as_tibble() %>%
                    dplyr::filter(gene_id %in% sig)
plotmat <- txi$abundance[c(sig_up, sig_down),] %>% as.data.frame() %>%
          rownames_to_column(var = "ensembl_gene_id") %>%
          left_join(gene_symbol, by = c("ensembl_gene_id" = "gene_id")) %>%
          drop_na(symbol)
plotmat$ensembl_gene_id <- NULL</pre>
plotmat <- plotmat %>% column_to_rownames(var = "symbol") %>% as.matrix()
# Color palette
heat.colors <- brewer.pal(6, "YlOrRd")</pre>
# Plot heatmap
pheatmap(plotmat,
         scale = "row",
         show_rownames = TRUE,
         border = FALSE,
         annotation = meta[, c("er"), drop = FALSE],
         main = "Top 50 Up- and Down- regulated genes in ER: positive vs negative",
         fontsize = 20)
```



```
# Create a matrix of normalized expression
sig_up <- resTP_tb_significant %>% arrange(-log2FoldChange) %>% head(50) %>% pull(gene)
sig_down <- resTP_tb_significant %>% arrange(log2FoldChange) %>% head(50) %>% pull(gene)
sig <- c(sig_up, sig_down)</pre>
row_annotation <- gene_symbol %>%
                    as_tibble() %>%
                    dplyr::filter(gene_id %in% sig)
plotmat <- txi$abundance[c(sig_up, sig_down),] %>% as.data.frame() %>%
          rownames_to_column(var = "ensembl_gene_id") %>%
          left_join(gene_symbol, by = c("ensembl_gene_id" = "gene_id")) %>%
          drop_na(symbol)
plotmat$ensembl_gene_id <- NULL</pre>
plotmat <- plotmat %>% column_to_rownames(var = "symbol") %>% as.matrix()
# Color palette
heat.colors <- brewer.pal(6, "YlOrRd")
# Plot heatmap
pheatmap(plotmat,
         scale = "row",
         show_rownames = TRUE,
         border = FALSE,
         annotation = meta[, c("tumor_percentage_high"), drop = FALSE],
         main = "Top Up/Down-regulated genes in Tumor_percentage_high: high vs low",
         fontsize = 20)
```



```
# Create a matrix of normalized expression
sig_up <- resD0_tb_significant %>% arrange(-log2FoldChange) %>% head(50) %>% pull(gene)
sig_down <- resDO_tb_significant %>% arrange(log2FoldChange) %>% head(50) %>% pull(gene)
sig <- c(sig_up, sig_down)</pre>
row_annotation <- gene_symbol %>%
                    as_tibble() %>%
                    dplyr::filter(gene_id %in% sig)
plotmat <- txi$abundance[c(sig_up, sig_down),] %>% as.data.frame() %>%
          rownames_to_column(var = "ensembl_gene_id") %>%
          left_join(gene_symbol, by = c("ensembl_gene_id" = "gene_id")) %>%
          drop_na(symbol)
plotmat$ensembl_gene_id <- NULL</pre>
plotmat <- plotmat %>% column_to_rownames(var = "symbol") %>% as.matrix()
# Color palette
heat.colors <- brewer.pal(6, "YlOrRd")</pre>
# Plot heatmap
pheatmap(plotmat,
         scale = "row",
         show_rownames = TRUE,
         border = FALSE,
         annotation = meta[, c("response"), drop = FALSE],
         main = "Top 50 Up- and Down- regulated genes in date_of: 20180323 vs 20180228",
         fontsize = 20)
```



#### Generate input files for GSEA

```
# prepares an expression profile for GSEA
# http://software.broadinstitute.org/cancer/software/gsea/wiki/index.php/Data_formats#Expression_Data_
# for GSEA it is important to report all genes - genome wide
# hopefully cpms are better than logcpms
counts <- counts[rowSums(counts)>0,]
result_file <- paste0("tables/1day.4gsea.txt")</pre>
counts_gsea <- counts %>% as.data.frame() %>%
    rownames_to_column(var = "ensembl_gene_id") %>%
    left_join(gene_symbol, by = c("ensembl_gene_id" = "gene_id")) %>%
    dplyr::relocate(symbol)
#%>%
     dplyr::relocate(ensembl_gene_id)
colnames(counts_gsea)[1:2] <- c("NAME", "DESCRIPTION")</pre>
d <-duplicated(counts_gsea$NAME)</pre>
o <- order(rowSums(counts_gsea[,rownames(meta)]),decreasing = T)</pre>
counts_gsea <- counts_gsea[o, ]</pre>
counts_gsea <- counts_gsea[!d, ]</pre>
samples_yes <- meta %>% dplyr::filter(response == "Yes") %>% row.names()
samples_no <- meta %>% dplyr::filter(response == "No") %>% row.names()
counts_gsea <- counts_gsea[,c("NAME", "DESCRIPTION", samples_yes, samples_no)]</pre>
# qsea now supports ENSEMBL IDs
write_tsv(counts_gsea, result_file)
```

#### Functional analysis

#### Biological Process (BP)

```
bg_genes <- rownames(resResponse)</pre>
## Run GO enrichment analysis
compGO <- enrichGO(gene = sigResponse_up,</pre>
                   universe = bg_genes,
                   keyType = "ENSEMBL",
                   OrgDb = "org.Hs.eg.db",
                   ont = "BP",
                   qvalueCutoff = 0.05,
                   pAdjustMethod = "BH",
                   readable = TRUE)
## Error in enrichGO(gene = sigResponse_up, universe = bg_genes, keyType = "ENSEMBL", : could not find
#dotplot(compGO,
         showCategory = 20,
         title = "GO (Biological Process) Enrichment \n Analysis for UP in Responders)",
#
         label\_format = 20,
         font.size = 10)
# image pdf 12 x 12
## Output results from GO analysis to a table
print("UP")
## [1] "UP"
results_up <- data.frame(compGO@result) %>% dplyr::filter(p.adjust < 0.05)
## Error in data.frame(compGO@result): object 'compGO' not found
nrow(results_up)
## Error in h(simpleError(msg, call)): error in evaluating the argument 'x' in selecting a method for f
write_csv(results_up, "tables/T21.day8.G0_BP_UP.csv")
## Error in is.data.frame(x): object 'results_up' not found
compGO <- enrichGO(gene = sigResponse_down,</pre>
                   universe = bg_genes,
                   keyType = "ENSEMBL",
                   OrgDb = "org.Hs.eg.db",
                   ont = "BP",
                   qvalueCutoff = 0.05,
                   pAdjustMethod = "BH",
                   readable = TRUE)
## Error in enrichGO(gene = sigResponse_down, universe = bg_genes, keyType = "ENSEMBL", : could not fin
results_down <- data.frame(compGO@result) %>% dplyr::filter(p.adjust < 0.05)
## Error in data.frame(compGO@result): object 'compGO' not found
print("Down")
```

```
## [1] "Down"
nrow(results_down)
```

## Error in h(simpleError(msg, call)): error in evaluating the argument 'x' in selecting a method for f

#### R session

#### sessionInfo()

```
## R version 4.0.3 (2020-10-10)
## Platform: x86_64-redhat-linux-gnu (64-bit)
## Running under: Fedora 32 (Workstation Edition)
##
## Matrix products: default
## BLAS/LAPACK: /usr/lib64/libopenblas-r0.3.12.so
## locale:
  [1] LC_CTYPE=en_CA.UTF-8
                                   LC_NUMERIC=C
   [3] LC_TIME=en_CA.UTF-8
                                   LC_COLLATE=en_CA.UTF-8
## [5] LC_MONETARY=en_CA.UTF-8
                                   LC_MESSAGES=en_CA.UTF-8
  [7] LC_PAPER=en_CA.UTF-8
                                   LC_NAME=C
## [9] LC_ADDRESS=C
                                   LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_CA.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] parallel stats4
                           stats
                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
##
## other attached packages:
## [1] ensembldb_2.14.1
                                     AnnotationFilter_1.14.0
## [3] GenomicFeatures_1.42.3
                                     AnnotationDbi_1.52.0
## [5] AnnotationHub_2.22.1
                                    BiocFileCache_1.14.0
## [7] dbplyr_2.1.1
                                    knitr_1.30
## [9] ggrepel_0.9.1
                                    tximport_1.18.0
## [11] DEGreport_1.26.0
                                    pheatmap_1.0.12
## [13] RColorBrewer_1.1-2
                                     forcats_0.5.1
## [15] stringr_1.4.0
                                    dplyr_1.0.5
                                    readr_1.4.0
## [17] purrr_0.3.4
                                    tibble_3.1.1
## [19] tidyr_1.1.3
## [21] ggplot2_3.3.3
                                     tidyverse_1.3.1
## [23] DESeq2_1.30.1
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## [25] Biobase_2.50.0
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## [29] GenomeInfoDb_1.26.7
                                     IRanges_2.24.1
## [31] S4Vectors_0.28.1
                                    BiocGenerics_0.36.1
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     [1] readxl_1.3.1
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     [3] circlize_0.4.12
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##
     [5] lazyeval_0.2.2
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##
     [7] splines_4.0.3
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##
     [9] digest_0.6.27
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  [11] fansi_0.4.2
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[13] memoise_2.0.0
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## [15] limma_3.46.0
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## [17] Biostrings 2.58.0
                                       annotate 1.68.0
## [19] Nozzle.R1_1.1-1
                                       modelr_0.1.8
##
   [21] askpass_1.1
                                       prettyunits_1.1.1
## [23] colorspace_2.0-0
                                       blob 1.2.1
## [25] rvest 1.0.0
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## [27] haven 2.4.1
                                       xfun 0.19
## [29] crayon_1.4.1
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## [31] jsonlite_1.7.2
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## [33] survival_3.2-7
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## [37] XVector_0.30.0
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## [39] GetoptLong_1.0.5
## [41] shape_1.4.5
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## [45] edgeR_3.32.1
                                       Rcpp_1.0.6
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## [53] httr_1.4.2
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## [59] locfit 1.5-9.4
                                       utf8 1.2.1
## [61] labeling_0.4.2
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## [63] rlang_0.4.10
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## [65] munsell_0.5.0
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## [67] cellranger_1.1.0
                                       tools_4.0.3
## [69] cachem_1.0.4
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                                       RSQLite_2.2.7
## [73] broom_0.7.6
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## [83] xml2_1.3.2
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## [85] compiler 4.0.3
                                       rstudioapi 0.13
## [87] curl_4.3
                                       png_0.1-7
   [89] interactiveDisplayBase_1.28.0 reprex_2.0.0
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## [91] geneplotter_1.68.0
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## [95] Matrix 1.2-18
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## [97] vctrs 0.3.7
## [99] lifecycle_1.0.0
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## [101] GlobalOptions_0.1.2
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## [103] cowplot_1.1.1
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## [105] rtracklayer_1.50.0
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## [107] R6_2.5.0
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## [113] withr_2.4.2
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## [115] Rsamtools_2.6.0
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## [117] GenomeInfoDbData_1.2.4
                                       hms_1.0.0
## [119] grid_4.0.3
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lubridate\_1.7.10