# wind: wORKFLOW FOR PiRNAs AnD BEYONd

Computational workflow for Data Exploration resulted from smallRNA-seq

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#### 1. Load libraries

```
suppressPackageStartupMessages({
  library('tidyverse')
  library('data.table')
  library('plyranges')
  library('tximport')
  library('edgeR')
  library('NOISeq')
  library('rafalib')
  library('pheatmap')
  library('RColorBrewer')
})
```

# 2. Directory generation for the resulted files

#### i. Add date

Used as an identifier for the folder

```
todate <- format(Sys.time(), "%d_%b_%Y")</pre>
```

ii. Make the directory for the results of the exploratory data analysis

```
my_basename <- "GSE68246" ## INPUT name of the folder
my_exp <- "br_cancer" ## INPUT name of the analysis
genome_input <- "hg38" ## INPUT genome version here
my_exp_sal <- "salmon"
my_exp_fc <- "featureCounts"
dat_path <- str_glue("{my_basename}/ExpDatAnalysis_{my_exp}_{genome_input}_{todate}/{c(my_exp_sal,my_exp_sal)})
dat_path %>% map(~dir.create(str_glue("./{.x}"), recursive = TRUE))
```

# 3. Import the salmon files

```
# load salmon files----
files_salm <- list.files(path = my_basename,pattern = ".sf",
    recursive = TRUE, full.names = TRUE)

names(files_salm) <- files_salm %>%
    str_remove("/quant.sf") %>% basename %>% str_remove(".trimmed.+")

# tximport------
txi <- tximport::tximport(files_salm, type = "salmon",
    txOut = TRUE, countsFromAbundance = "lengthScaledTPM")</pre>
```

# 4. Make or import the targets file.

If you use a public dataset you can download it from SRA RUN selector Metadata or you can create it. It has to have at least three columns with the column names: "sample\_name", "group", "batch" Here we show both ways although we need to manually add some columns in the metadata of SRA run selector.

```
# make the targets object ----
# INPUT targets file
                          ## INPUT targets file here
targets file <- tibble(</pre>
 file_name = files_salm,
  sample_name = names(files_salm),
  group = as_factor(c(rep("MCF_7_Monolayer",3),rep("MCF_7_Spheroid",3))),
  batch = as factor(c(1:3,1:3))
)
targets_file <- list.files(pattern = "SraRun",</pre>
                            path = my_basename, full.names = T) %>%
 read_csv() %>%
  select(sample_name = Run, Genotype) %>%
  mutate(group = Genotype %>%
           str_remove(" .+") %>%
           str_remove("pri-") %>%
           str_replace("-","_")%>% as_factor(),
         batch = as factor(1:9))
```

# 5. Make a DGElist object for salmon

```
# DGElist
\# from https://bioconductor.org/packages/release/bioc/vignettes/tximport/inst/doc/tximport.html
# we follow the instructions to import for edgeR
cts <- txi$counts
normMat <- txi$length
# Obtaining per-observation scaling factors for length, adjusted to avoid
# changing the magnitude of the counts
normMat <- normMat/exp(rowMeans(log(normMat)))</pre>
normCts <- cts/normMat</pre>
# Computing effective library sizes from scaled counts, to account for
# composition biases between samples
eff.lib <- calcNormFactors(normCts) * colSums(normCts)</pre>
# Combining effective library sizes with the length factors, and calculating
# offsets for a log-link GLM
normMat <- sweep(normMat, 2, eff.lib, "*")</pre>
normMat <- log(normMat)</pre>
# Creating a DGEList object for use in edgeR.
dgl_salmon <- DGEList(cts, samples = targets_file) %>%
  scaleOffset( normMat) %>%
 write_rds(str_glue("{dat_path[1]}/dgl_edgeR_salmon.rds"))
```

```
# remove objects.
rm(cts, normCts, normMat, txi)
```

# 6. Import the FeatureCounts object and make a DGElist object

```
# load the rds from featureCOunts----
#INPUT rds featureCOunts
fc <- list.files(path = my_basename, recursive = TRUE,</pre>
 pattern = ".+counts.+.rds",
 full.names = TRUE) %>%
 read rds()
# write the matrix for the analysis, annotation stats----
colnames(fc$counts) <- targets_file$sample_name</pre>
fc$counts %>% as_tibble(rownames = "sRNA") %>%
  write_tsv(str_glue("{dat_path[2]}/raw_reads_fc.txt"))
fc$annotation %>%
  as_tibble() %>%
  write_tsv(str_glue("{dat_path[2]}/annotation_fc.txt"))
fc$stat %>%
  as_tibble() %>%
  write_tsv(str_glue("{dat_path[2]}/stats_fc.txt"))
dgl_fc <- DGEList(counts = fc$counts,</pre>
               samples = targets_file,
               lib.size = colSums(fc$counts),
               norm.factors = rep(1,ncol(fc$counts)))
# give colours to samples ----
pal1 <- tibble(value = c("#1C1718","#B6854D", "#79402E","#FF0000",</pre>
  "#D8B70A","#02401B","#046C9A","#5B1A18")) %>%
  dplyr::slice(1:length(levels(as_factor(targets_file$group)))) %>%
 mutate(
    group = as_factor(levels(as_factor(targets_file$group))))
dgl_fc$colours <- as_factor(inner_join(dgl_fc$samples, pal1,by= "group")$value)
# remove objects ----
rm(pal1)
```

# 7. Create biodetection plot with NOISeq

```
mybiotypes <- fc$annotation %>%
  mutate(gene_type = gene_type %>% str_remove(";.+")) %>%
  select(GeneID,gene_type) %>%
  column_to_rownames("GeneID")

function_Noiseq_plots <- function(exp_data, plot_path){
  mydata <- NOISeq::readData(data = exp_data,
  factors = as.data.frame(targets_file),</pre>
```

```
biotype = mybiotypes)
mybiodetection <- dat(mydata, k = 0, type = "biodetection")
pdf(str_glue("{plot_path}/NOISeq_biodetection_exprs_{todate}_{str_remove(plot_path,'.+/')}.pdf"))
seq(ncol(exp_data)) %>% map(~explo.plot(mybiodetection, samples = .x),plottype = "boxplot")
dev.off()
mycountsbio <- dat(mydata, factor = NULL, type = "countsbio")
pdf(str_glue("{plot_path}/NOISeq_countsbio_{todate}_{str_remove(plot_path,'.+/')}.pdf"))
seq(ncol(exp_data)) %>% map(~explo.plot(mycountsbio,
    samples = .x ,plottype = "boxplot"))
dev.off()
}
list( "salmon" = dgl_salmon$counts, "fc" = fc$counts) %>%
map2(.y = dat_path, ~function_Noiseq_plots(.x,.y))
```

#### 8. Create the design matrix

```
design <- model.matrix(~0 + targets_file$group)
colnames(design) <- colnames(design) %>%
    str_remove("targets_file\\$group")
rownames(design) <- targets_file$sample_name
design_2 <- model.matrix(~0 + targets_file$group + targets_file$batch)
colnames(design_2) <- colnames(design_2) %>%
    str_remove("targets_file\\$group") %>%
    str_remove("targets_file\\$batch")
rownames(design_2) <- targets_file$sample_name</pre>
```

#### 9. Perform various Filtering Methods: EdgeR, NOIseq

```
function_filtering <- function(dgl_data, data_path){</pre>
  # filtering with NOISEg -----
  noifil <- list("cpm" = 1L, "Prop" = 3L) %>%
    map(~NOISeq::filtered.data(dgl data$counts,
      factor = targets_file$group,
      norm = FALSE,
      method = .x, cv.cutoff = 100, cpm = 1)
  noifil %>%
    names %>%
    map( ~ dgl_data[rownames(dgl_data$counts) %in%
      rownames(noifil[.x]),,keep.lib.sizes = FALSE] %>%
        write_rds(str_glue("{data_path}/dgl_{.x}_filt.rds"))
  # filterwith EdgeR ----
  keep.exprs <- filterByExpr.DGEList(dgl_data, design = design)</pre>
  keep.exprs_2 <- filterByExpr.DGEList(dgl_data, design = design_2)</pre>
  dgl filt <- dgl data[keep.exprs,,keep.lib.sizes=FALSE] %>%
    write_rds(str_glue("{data_path}/dgl_filt_nobatch.rds"))
```

```
dgl_filt_2 <- dgl_data[keep.exprs_2,,keep.lib.sizes=FALSE] %>%
    write_rds(str_glue("{data_path}/dgl_filt_batch.rds"))
  features_NOIS <- map(noifil, ~ .x %>%
      rownames() %>%
      enframe(name = NULL))
  features_edgeR <- map(list(dgl_filt, dgl_filt_2) , ~ .x %>%
      rownames() %>%
      enframe(name = NULL)) %>%
    set names("no batch", "batch")
  common_edgeR_nobatch <- map2(features_edgeR[1], features_NOIS, ~ .x %>%
      inner join(.y))
  common_edgeR_batch <- map2(features_edgeR[2], features_NOIS, ~ .x %>%
      inner_join(.y))
  filter_info <- tibble(</pre>
    "features" = c("Starting_features:", "edgeR_nobatch_filter:",
      "edgeR_batch_filter:",
      "NOISeq_1cpm_filter:",
      "common_with_edgeR_nobatch:", "common_with_edgeR_batch:",
      "NOISeq_Proportion_filter:",
      "common_with_edgeR_nobatch:", "common_with_edgeR_batch:"
    "number of features" = c(nrow(dgl data$counts), nrow(dgl filt$counts),
      nrow(dgl filt 2$counts),
      nrow(noifil[[1]]),
      nrow(common_edgeR_nobatch[[1]]),nrow(common_edgeR_batch[[1]]),
      nrow(noifil[[2]]),
     nrow(common_edgeR_nobatch[[2]]),nrow(common_edgeR_batch[[2]])
   )
  ) %>%
    write_tsv(str_glue("{data_path}/filtering_info.txt"))
  dgl_filt
filtered_dgls <- list("salmon" = dgl_salmon, "fc" = dgl_fc) %>%
 map2(.y = dat_path, ~function_filtering(.x,.y))
```

#### 10. Histogram before and after filtering of data

```
function_hist <- function(dgl_data, dgl_fil_data, plot_path){
  AveLogCpm_Raw_Data <- aveLogCPM(dgl_data)
  AveLogCpm_Filtered_Data <-aveLogCPM(dgl_fil_data)
  pdf(str_glue("{plot_path}/histogram_plot_{todate}_{str_remove(plot_path,'.+/')}.pdf"))
  hist(AveLogCpm_Raw_Data)
  hist(AveLogCpm_Filtered_Data)
dev.off()
}
list(list("salmon" = dgl_salmon, "fc" = dgl_fc),
  filtered_dgls, dat_path) %>%
```

#### 11. Normalization

```
function EDA RLE <- function(data,name){EDASeq::plotRLE(data,</pre>
        col = as.character(dgl_fc$colours),
        outline=FALSE, las=3,
       ylab="Relative Log Expression",
        cex.axis=1, cex.lab=1, main = str_glue("{name}"))
      legend("topright",
       legend= levels(as_factor(dgl_fc$samples$group)),
       fill = levels(as_factor(dgl_fc$colours)),
       bty="n",
       cex = 0.5, inset = c(.01,.01))
   }
function_norm <- function(dgl_fil_data, data_path){</pre>
  # edgeR ----
 norm_method <- list("none", "TMM", "TMMwsp", "RLE") %>%
    set names(.)
  edger_norm <- map(norm_method, ~calcNormFactors(dgl_fil_data, method = .x))</pre>
  # voom ----
  pdf(str_glue("{data_path}/voom_plots_{str_remove(data_path,'.+/')}.pdf"))
  voom_norm <- edger_norm[1:3] %>%
   map2(.y = c("quantile", rep("none",2)),
      ~voom(.x, design = design,
       plot = TRUE, normalize.method = .y)) %>%
    set_names("voom_Quantile","voom_TMM","voom_TMMwsp")
  dev.off()
  # voom with quality weights ----
  pdf(str_glue("{data_path}/voom_quality_weights_plots_{str_remove(data_path,'.+/')}.pdf"))
  voom_norm_QW <- edger_norm[1:3] %>%
   map2(.y = c("quantile", rep("none",2)),
      ~voomWithQualityWeights(.x, design = design,
        plot = TRUE, normalize.method = .y)) %>%
   set_names("voomQW_Quantile","voomQW_TMM","voomQW_TMMwsp")
  dev.off()
  # list of normalized data ----
  norm_list <- c(edger_norm %>% map(~cpm(.x, normalized.lib.sizes = TRUE)),
    list(
    "voom_Quantile" = 2^voom_norm[[1]]$E,
    "voom_TMM" = 2^voom_norm[[2]]$E,
    "voom_TMMwsp" = 2^voom_norm[[3]]$E,
    "voomQW_Quantile" = 2^voom_norm_QW[[1]]$E,
    "voomQW_TMM" = 2^voom_norm_QW[[2]]$E,
    "voomQW_TMMwsp" = 2^voom_norm_QW[[3]]$E))
  pdf(str_glue("{data_path}/RLE_plots_{str_remove(data_path,'.+/')}.pdf"))
  norm_list %>%
    imap(~function_EDA_RLE(.x,.y))
  dev.off()
  norm_list[2:4] %>% imap(~.x %>%
```

# 12. Make h-clustering

```
function_clust <- function(dgl_norm_data, plot_path){</pre>
  hc_methods <- c("ward.D2",</pre>
                "complete",
                "average")
  list_distc <- c(dgl_norm_data[1:4] %>%
      map(~cpm(.x, normalized.lib.sizes = TRUE, log=TRUE, prior.count=5)),
      list("voom_Quantile" = dgl_norm_data[[5]]$E,
      "voom_TMM"= dgl_norm_data[[6]]$E,
      "voom_TMMwsp" = dgl_norm_data[[7]]$E,
      "voomQW_Quantile" = dgl_norm_data[[8]]$E,
      "voomQW_TMM"= dgl_norm_data[[9]]$E,
      "voomQW_TMMwsp" = dgl_norm_data[[10]]$E)) %>% map(~dist(t(.x)))
  #pheatmap start
  list_distc_mat <- list_distc %>% map(~as.matrix(.x))
  colours_pheat <- colorRampPalette(rev(brewer.pal(9, "Blues")))(255)</pre>
  pdf(str_glue("{plot_path}/distance_matrix_hclust_{str_remove(plot_path,'.+/')}.pdf"))
  list_distc_mat %>% imap(~pheatmap(.x,
           clustering_distance_rows = "euclidean",
           clustering_distance_cols = "euclidean",
           col = colours_pheat,
           main = str_glue({.y})))
  dev.off()
  #pheatmap end
  #list_distc <- log_cpm %>% map(~dist(t(.x)))
  list_hc <- sapply(hc_methods, function(x) map(list_distc, ~hclust(.x,method = x)))</pre>
  names(list_hc) <- rep(rownames(list_hc), times = ncol(list_hc))</pre>
  pdf(str_glue("{plot_path}/hierarchic_clust_{str_remove(plot_path, '.+/')}.pdf"))
  for (i in seq_along(list_hc)) {
       rafalib::myplclust(list_hc[[i]],
       lab.col = as.character(dgl_fc$colours),
       xlab = NULL,
       main = str_glue("{enframe(list_hc[[i]])$value[7]} - {enframe(list_hc[[i]])$value[5]} - {names(li
       legend("topright",
       legend = levels(dgl_fc$samples$group),
       fill = levels(dgl_fc$colours),
```

```
bty="n",
    cex = 0.9)
    }
    dev.off()
}
map2(norm_dgls, dat_path, ~function_clust(.x,.y))
```

# 13. Make MDS plot

```
function_MDS <- function(dgl_norm_data, plot_path){</pre>
  par(mar=c(6,5,2,1)+0.1)
  pdf(str_glue("{plot_path}/MDS_plot_{str_remove(plot_path,'.+/')}.pdf"))
  plotMDS(dgl_norm_data$TMM,
          labels = dgl fc$samples$sample name,
          pch = 10,
          cex = 0.7,
    col = as.character(dgl_fc$colours), dim.plot = c(1,2))
  legend("topright",
       legend = levels(dgl_fc$samples$group),
       fill = levels(dgl_fc$colours),
       bty="n",
       cex = 1.5, inset = c(.01,.09))
  map2(c(3,1,2,2),c(4,3,3,4),
  rplotMDS(dgl_norm_data$TMM, labels = dgl_fc$samples$sample_name, pch = 10,
   cex = 0.7,
   col = as.character(dgl_fc$colours),
   dim.plot = c(.x,.y),
   main = str_glue("MDS plot {names(dgl_norm_data[2])}"))
 )
 dev.off()
map2(norm_dgls, dat_path, ~function_MDS(.x,.y))
```

#### 14. Make PCA plot

```
## modified from DESeq2::plotPCA
## https://github.com/mikelove/DESeq2/blob/master/R/plots.R

# MAKE IF STATEMENT FOR LIMMA!!!!!!

function_PCA <- function(dgl_norm_data, plot_path, norm_method="TMM", ntop=500){
    library(DESeq2)
    # calculate the variance for each gene
    rv <- rowVars(dgl_norm_data[[norm_method]]$counts)
    # select the ntop genes by variance
    select <- order(rv, decreasing=TRUE)[seq_len(min(ntop, length(rv)))]

# perform a PCA on the data in assay(x) for the selected genes</pre>
```

```
pca <- prcomp(t(dgl_norm_data[[norm_method]]$counts[select,]))</pre>
  # the contribution to the total variance for each component
  percentVar <- pca$sdev^2 / sum( pca$sdev^2 )</pre>
  # create a new grouping factor
  group <- dgl_norm_data[[norm_method]]$samples$group</pre>
  # assembly the data for the plot
  d <- data.frame(PC1=pca$x[,1], PC2=pca$x[,2], group=group,</pre>
                  name=colnames(dgl_norm_data[[norm_method]]$counts))
  # create batch if exist factor
  if(dgl_norm_data[[norm_method]]$samples$batch){
    d$batch <- dgl_norm_data[[norm_method]]$samples$batch %>% as_factor
  p <- ggplot(data=d, aes_string(x="PC1", y="PC2", color="group", shape="batch"))</pre>
  }else{
  p <- ggplot(data=d, aes_string(x="PC1", y="PC2", color="group"))}</pre>
  p <- p +
    geom_point(size=3) +
    xlab(paste0("PC1: ",round(percentVar[1] * 100),"% variance")) +
    ylab(paste0("PC2: ",round(percentVar[2] * 100),"% variance")) +
    coord_fixed()+
    theme minimal()+
    labs(title=str_glue("PCA plot {norm_method}"))+
    theme(plot.title = element text(hjust = 0.5))
  # pdf
  par(mar=c(6,5,2,1)+0.1)
 pdf(str_glue("{plot_path}/PCA_plot_{norm_method}_{str_remove(plot_path,'.+/')}.pdf"))
 print(p)
 dev.off()
}
map2(norm_dgls, dat_path, ~function_PCA(.x,.y))
```

#### 15. Compare groups between FeatureCounts and salmon results

```
function_comp_groups <- function(dgl_norm_data, tool){
  grouped_cpm <- dgl_norm_data$TMM %>%
     cpmByGroup.DGEList

# keep <- rowSums(grouped_cpm > 1) > 0

# grouped_cpm[keep,] %>%
     grouped_cpm[keep,] %>%
     as_tibble(rownames = "sncRNA") %>%
     rename_at(vars(-matches("sncRNA")), list(~str_c(.,tool)))
}

comp_FC_sal <- map2(norm_dgls, list("_salmon", "_fc"), ~function_comp_groups(.x,.y))

#comp_FC_sal <- map2(list("salmon" = norm_dgls_sal, "FC" = norm_dgls_FC),
# list("_salmon", "_FC"), ~function_comp_groups(.x,.y))</pre>
```

```
#comp_FC_sal$salmon %>%
# inner_join(comp_FC_sal$fc, by = "sncRNA") %>%
\# write_tsv(str_glue("{str_remove(dat_path[1],'/.+')}/salmon_FC_1cpm_common_grouped.txt"))
salmon_FC_cpm_union_grouped <- comp_FC_sal$salmon %>%
  full_join(comp_FC_sal$fc, by = "sncRNA") %>%
  write_tsv(str_glue("{str_remove(dat_path[1],'/salmon|/featureCounts')}/salmon_FC_cpm_union_grouped.tx
# piCK the 100 top expressed molecules between FC salmon and all groups
all_exprs_cpm_TMM <- list.files(recursive = T ,</pre>
           pattern = "norm_cpm_TMM_",
           full.names = T) %>%
  vroom::vroom(id = "method") %>%
  mutate(method = method %>% basename() %>% str_remove("norm_cpm_TMM_") %>% str_remove(".txt"))
salmon_FC_cpm_union_grouped_top <- salmon_FC_cpm_union_grouped %>%
  filter(gene_type == "piRNA") %>%
  arrange(across(.fns = ~.x %>% desc ,
                 .cols = ends_with(c("salmon", "fc")))) %>%
  slice_head(n = 100) %>%
  all_exprs_cpm_TMM %>%
  mutate(method = if else(method == "featureCounts",
                          true = "fc",
                          false = "salmon")) %>%
  pivot_longer(cols = c(CM1:CM3)) %>% unite(col = "sample",c(name, method)) %>%
  pivot_wider(names_from = "sample",
              values_from = "value") %>%
  column_to_rownames("GeneIDs") %>%
  .[salmon_FC_cpm_union_grouped_top$sncRNA,] %>%
  as_tibble(rownames = "smallRNA") %>% write_tsv(str_glue("{str_remove(dat_path[1], '/salmon|/featureCou
```

#### 16. Make histogram of length

```
annot_tbl <- read_gff2("../genome_transc_human/ncRNA_transcripts_100bp_RNA_Central_piRNAbank_hg38.gtf")
as_tibble %>%
select(-c(type, score, phase)) %>%
dplyr::rename("length_w" = width)

function_prep_hist <- function(dgl_norm_data, annot_gtf){
   prep_hist <- annot_gtf %>%
        distinct(gene_id, .keep_all = T) %>%
        filter(gene_id %in% rownames(dgl_norm_data$TMM))
}

smallRNA_seqs <- map2(norm_dgls, list(annot_tbl, annot_tbl), ~function_prep_hist(.x,.y))

smallRNA_seqs[["fc"]] %>% # create one also per group
   dplyr::count(gene_type, sort = T) %>%
   dplyr::rename("fc_n" = n) %>%
   full_join( smallRNA_seqs[["salmon"]] %>%
```

```
dplyr::count(gene_type, sort = T) %>%
  dplyr::rename("salmon_n" = n)) %>%
  full_join(
    smallRNA_seqs[["fc"]] %>%
    inner_join(smallRNA_seqs[["salmon"]] ) %>%
    dplyr::count(gene_type, sort = T) %>%
    dplyr::rename("common_n" = n)) %>%
  full join(
smallRNA seqs[["fc"]] %>%
    anti_join(smallRNA_seqs[["salmon"]] ) %>%
    dplyr::count(gene_type, sort = T) %>%
    dplyr::rename("unique_FC_n" = n)) %>%
   full_join(
smallRNA_seqs[["salmon"]] %>%
    anti_join(smallRNA_seqs[["fc"]]) %>%
   dplyr::count(gene_type, sort = T) %>%
    dplyr::rename("unique_salmon_n" = n)) %>%
  write_tsv(str_glue("{str_remove(dat_path[1], '/salmon|/featureCounts')}/stats_gene_types_ids.txt"))
# per group summary ----
groups <- targets_file$group %>% levels() %>% set_names()
salmon groups <- groups %>% map(~comp FC sal$salmon %>%
  filter_at(vars(matches(!!str_c(.x,"_salmon"))),
    any vars(. > 0)) %>%
    select(sncRNA) %>%
   mutate(!!str_c("group_",.x) := .x)
) %>% purrr::reduce(full_join) %>%
  mutate(salmon = "salmon") %>%
  left_join(smallRNA_seqs$salmon, by = c("sncRNA" = "gene_id")) %>%
  select(-c(seqnames:end, strand, source, sRNA_id, sRNA_id2)) %>%
  pivot_longer(cols = starts_with("group_"), values_to = "rank_salmon"
   ) %>% select(-name)
FC_groups <- groups %>% map(~comp_FC_sal$fc %>%
  filter_at(vars(matches(!!str_c(.x,"_fc"))),
    any_vars(. > 0)) %>%
    select(sncRNA) %>%
   mutate(!!str_c("group_",.x) := .x)
) %>%
 purrr::reduce(full_join) %>%
  mutate(FeatureCount = "FeatureCount") %>%
  left_join(smallRNA_seqs$fc, by = c("sncRNA" = "gene_id")) %>%
  select(-c(seqnames:end, strand, source, sRNA_id, sRNA_id2)) %>%
  pivot_longer(cols = starts_with("group_"), values_to = "rank_FC"
   ) %>% select(-name)
## histograms ----
pdf(str_glue("{str_remove(dat_path[1], '/salmon|/featureCounts')}/length_histogram.pdf"))
map(FC_groups$gene_type %>% as_factor() %>% levels(),
  ~salmon_groups %>%
  full_join(FC_groups,
   by = c("sncRNA", "seq_RNA",
```

```
"gene_type", "length_w")) %>%
filter(gene_type == .x) %>%
pivot_longer(c(salmon, FeatureCount),
   "tool", values_to = "Quantification") %>%
pivot_longer(c(rank_salmon, rank_FC), "rank", values_to = "groups") %>%
filter(!is.na(Quantification),!is.na(groups)) %>%
ggplot() +
geom_bar(mapping = aes(x = factor(length_w), fill = Quantification),
   position = "dodge") +
facet_wrap(~ groups,ncol = 2)+
   scale_x_discrete(name = 'length')+
   scale_y_continuous(labels = scales::comma)+
   ggtitle(.x) +
   coord_flip()
)
dev.off()
```

#### 17. Sequence logos

```
# sequences logos -----
library(ggseqlogo)
pdf(str_glue("{str_remove(dat_path[1],'/salmon|/featureCounts')}/piRNA_logos_FC_salmon.pdf"))
groups %>% map2(list(salmon_groups),
  ~.y %>%
   filter(gene_type == "piRNA") %>%
   filter_at(vars(starts_with("rank")), any_vars(. == .x)) %>%
  .$seq_RNA %>%
  str_sub(1,15) %>%
  ggseqlogo(method = 'prob', font="roboto_regular") +
  ggtitle(str_glue("Salmon_{.x}")))
#featurecounts
groups %>% map2(list(FC_groups),
  ~.y %>%
   filter(gene_type == "piRNA") %>%
   filter_at(vars(starts_with("rank")), any_vars(. == .x)) %>%
  .$seq_RNA %>%
  str sub(1,15) %>%
  ggseqlogo(method = 'prob', font="roboto_regular") +
  ggtitle(str_glue("FeatureCounts_{.x}")))
dev.off()
```

#### 18. Reads info and histograms

i. extract the information of length for reads

```
## filter for reads of 15-49 bases
for file in my_data/star_results/*/*_sorted.bam;
```

```
where_to_save=`dirname ${file}`;
regex=`basename ${file}`;
samp="${regex%%.trimmed_sorted.bam}";
echo "Processing sample ${samp} start: $(date)";
echo "end:$(date)";
done
## find length of reads from STAR - FeatureCounts
\#cut - f1 \ my\_data/piRNAs\_hist.txt > my\_data/piRNA\_ids.txt \ | \ grep \ -F \ -w \ -f \ my\_data/piRNA\_ids.txt \ -
for file in my_data/spike_ins/star_results/*.featureCounts.bam;
where_to_save=`dirname ${file}`;
regex=`basename ${file}`;
samp="${regex%%.trimmed_sorted.bam.featureCounts.bam}";
echo "Processing sample ${samp} start: $(date) and saving in: ${where_to_save}/${samp}_hist.txt";
samtools view -@ 6 ${file} | awk 'BEGIN{FS=OFS="\t"}{print length($10),$18}'| sed 's/XT:Z://g' | sort
echo "end:$(date)";
done
## find length of reads from salmon
for file in my_data/quants/*.fastq.gz.bam;
where_to_save=`dirname ${file}`;
regex=`basename ${file}`;
samp="${regex%%.trimmed*}";
echo "Processing sample ${samp} start: $(date) and saving in: ${where_to_save}/${samp}_hist_allRNA.txt"
samtools view -@ 2 ${file} | awk 'BEGIN{FS=OFS="\t"}{print $1,length($10),$3}'| sort -k1,1 | bedtools
echo "end:$(date)";
done
## if was to make only for piRNA
samtools view -@ 6 ${file} | awk 'BEGIN{FS=OFS="\t"}{print length($10),$3}' | grep -F -w -f my_data/piRN.
```

#### ii. make a txt file with the info

#### iii. ggplot for histograms

```
library(tidyverse)
library(plyranges)
```

```
#piRNAs_hist <- read_tsv("piRNAs_hist.txt")</pre>
# qtf_piB_RCentr <- read_qff2("mouse_data/ncRNA_transcripts_100bp_RNA_Central_piRNAbank_mm10.qtf") %>%
  as_tibble() %>%
  select(gene_id,gene_type) %>%
  distinct(gene_id, .keep_all = T) %>%
    mutate(gene_type = as_factor(gene_type))
# FC and salmon files -----
hist_files_fc <- list.files(path ="mouse_datasets/star_results",
                         pattern = "_hist.txt",
                         recursive = TRUE, full.names = TRUE) %>%
    set names(. %>%
                basename() %>%
                str remove(" hist.txt")
hist_files_salmon <- list.files(path ="mouse_datasets/quants",
                         pattern = "_hist_allRNA.txt",
                         recursive = TRUE, full.names = TRUE) %>%
    set_names(. %>%
                basename() %>%
                str_remove("_hist_allRNA.txt")
#FC reads info----
## add the regex of smallRNA names
smallRNA_Categ_regex <- levels(gtf_piB_RCentr$gene_type) %>%
  set names(.) %>%
  map(~gtf_piB_RCentr %>%
  filter(gene_type == .x) %>%
  .$gene_id %>%
  str_c(collapse = "|")) %>%
  c(.,"Not_assigned" = "Not_assigned")
## function to filter a df of mappeds reads per small RNA category using regex
reads_info_fc <- function(df, Categ_regex_list){</pre>
  Categ_regex_list %% #regex with rna_names "mmu_piR_039595|mmu_piR_039148"
    imap(~df %>% # for each category summarise the reads
           filter(str_detect(sncRNA,.x)) %>%
           summarise(!!str_c(.y,"_Reads"):= sum(Reads)) %>%
           add_column(Category = "Reads")
         ) %>%
   purrr::reduce(left_join) %>%
   left join(df %>%
                filter(Alignment!="Not_assigned") %>%
                group_by(Alignment) %>%
                summarise(S=sum(Reads)) %>%
                mutate( Category = "Reads") %>%
                pivot_wider(names_from = Alignment, values_from = S)) %>%
   mutate(Reads_in_analysis = Not_assigned_Reads + Multimapped + Unique,
      Mapped_Reads = Multimapped + Unique,
     Mapped_Reads_perc =
        as.character(round((Mapped_Reads/Reads_in_analysis)*100, digits = 2)) %>%
        str_c(.,"%")) %>%
```

```
select(Category, Reads_in_analysis, Mapped_Reads,
                Mapped_Reads_perc,
                Unique,
                Multimapped,
                Not_assigned_Reads, everything())
}
## read the hist files and mutate them
df_list <- hist_files_fc %>%
  map(~read_tsv(.x, col_names = c("Reads", "Length", "sncRNA")) %>%
  mutate(Length = as_factor(Length),
         sncRNA = if else(is.na(sncRNA), "Not assigned", sncRNA),
         Alignment = case_when(
           sncRNA == "Not_assigned" ~ "Not_assigned",
           str_detect(sncRNA, ",") ~"Multimapped" ,
           TRUE ~ "Unique" )
         )
  )
## create the data
FC_reads_info <- df_list %>%
  imap(~reads_info_fc(.x, smallRNA_Categ_regex) %>%
         mutate(Category = .y)) %>%
  purrr::reduce(bind rows)
dgl_norm$featureCounts$TMM$counts %>%
  as_tibble(rownames = "smallRNA") %>%
  filter(smallRNA %in% annot_tbl$gene_id) %>% # annot_tbl <- gtf_piB_RCentr %>% filter(gene_type =="piR
  pivot_longer(-smallRNA) %>%
  mutate(piRNA = if_else(value > 0,1,0)) %>%
  group_by(name) %>%
  summarise(piRNAs_in_sample =sum(piRNA)) %>%
  dplyr::rename(Category = name) %>%
  left_join(FC_reads_info) %>% # save them on a txt
write_tsv("mouse_datasets/FC_reads_info.txt")
# FeatureCounts histograms of piRNAs-----
piRNA_DF_hist<- df_list %>%
  bind_rows(.id = "Sample") %>%
  filter(sncRNA != "Not_assigned") %>%
  filter(str_detect(sncRNA, smallRNA_Categ_regex[["piRNA"]]))
#FC reads hist piRNA----
piRNA_DF_hist %>%
   group_by(Sample, Length, Alignment) %>%
   summarise(Reads = sum(Reads)) %>%
   ggplot() +
   geom_bar(mapping =
              aes(x = Length,
                  y = Reads,
                  fill = Alignment),
```

```
stat = "identity") +
   scale_y_continuous(labels = scales::comma) +
   coord_cartesian(ylim = c(0,2000000))+
   theme bw() +
   facet_wrap(~Sample, ncol = 3) +
   theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
pdf(str_glue("mouse_datasets/histograms_piRNA_reads_facets_fc.pdf"))
df_list %>% imap(~.x %>%
           filter(str_detect(sncRNA, smallRNA_Categ_regex[["piRNA"]])) %>%
           group_by(Length,Alignment) %>%
           summarise(Reads = sum(Reads)) %>%
           ggplot() +
             geom_bar(mapping = aes(x = Length, y = Reads),
                      stat = "identity") +
             scale_y_continuous(labels = scales::comma) +
             theme_bw() +
             facet_grid(Alignment ~ .) +
             ggtitle(str_glue(" Histogram of {.y} sample"))+
             theme(plot.title = element_text(hjust = 0.5),
        axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
dev.off()
## pick only spike ins----
spike_reg_ex <- piRNAs_hist %>%
 filter(gene_type == "spike_in") %>%
  .$gene_id %>%
  str_c(collapse = "|") %>%
  set_names("spike_ins")
piRNA_reg_ex <- piRNAs_hist %>%
  filter(gene_type == "piRNA") %>%
  .$gene_id %>%
  str_c(collapse = "|") %>%
  set_names("piRNA")
miRNA_reg_ex <- gtf_piB_RCentr %>%
  as tibble() %>%
  filter(gene_type == "miRNA") %>%
  distinct(gene_id) %>%
  .$gene id %>%
  str_c(collapse = "|") %>%
  set_names("miRNA")
pdf(str_glue("histograms_spike_ins_reads_Salmon.pdf"))
map(hist_files_salmon,~read_tsv(.x, col_names = c("Reads", "Length", "sncRNA")) %>%
  mutate(Length = as_factor(Length)) %>%
  filter(str_detect(sncRNA,spike_reg_ex)) %>%
  group_by(Length) %>%
  summarise(Reads = sum(Reads)) %>%
```

```
ggplot() +
  geom_bar(mapping = aes(x = Length, y = Reads), stat = "identity")+
  scale_y_continuous(labels = scales::comma)+
  theme minimal()+
  ggtitle(.x %>% basename %>% str_replace("allRNA.txt","spike_ins"))
  dev.off()
pdf(str glue("histograms piRNA reads FC filtered.pdf"))
map(hist_files_fc, ~read_tsv(.x, col_names = c("Reads", "Length", "sncRNA")) %>%
 mutate(Length = as_factor(Length)) %>%
  filter(str_detect(sncRNA,piRNA_reg_ex)) %>%
  group_by(Length) %>%
  summarise(Reads = sum(Reads)) %>%
 ggplot() +
  geom_bar(mapping = aes(x = Length, y = Reads), stat = "identity")+
  scale_y_continuous(labels = scales::comma)+
  theme_minimal()+
  ggtitle(.x %>% basename %>% str_replace("allRNA.txt","piRNA"))
  dev.off()
pdf(str_glue("histograms_miRNA_reads_Salmon.pdf"))
map(hist files salmon,~read tsv(.x, col names = c("Reads", "Length", "sncRNA")) %>%
 mutate(Length = as factor(Length)) %>%
  filter(str_detect(sncRNA,miRNA_reg_ex)) %>%
  group_by(Length) %>%
  summarise(Reads = sum(Reads)) %>%
  ggplot() +
  geom_bar(mapping = aes(x = Length, y = Reads), stat = "identity")+
  scale_y_continuous(labels = scales::comma)+
  theme_minimal()+
  ggtitle(.x %>% basename %>% str_replace("allRNA.txt","miRNA"))
 dev.off()
pdf(str glue("histograms all RNA reads Salmon.pdf"))
map(hist_files_salmon,~read_tsv(.x, col_names = c("Reads", "Length", "sncRNA")) %>%
  mutate(Length = as factor(Length)) %>%
  #filter(str_detect(sncRNA,miRNA_reg_ex)) %>%
  group_by(Length) %>%
  summarise(Reads = sum(Reads)) %>%
  ggplot() +
  geom_bar(mapping = aes(x = Length, y = Reads), stat = "identity")+
  scale_y_continuous(labels = scales::comma)+
  theme_minimal()+
  ggtitle(.x %>% basename %>% str_replace("allRNA.txt","all_RNA"))+
  coord_flip()
)
```

```
dev.off()
## piRNA reads -----
reads_piRNA <- read_tsv("reads_piRNA.txt",</pre>
                        col_names = c("Read","Length","sRNAs","read_sequence", "Sigar"))
reads piRNA %>% count(Sigar)
reads_piRNA %>% count(Length)
piRNA_reads <- reads_piRNA %>%
  select(Read, sRNAs, read_sequence, Length) %>%
  separate(sRNAs, str_c("V",1:2),
           extra = "merge", fill = "right", sep = ",") %>%
  filter(is.na(V2)) %>%
  select(-V2) %>%
  filter(str_detect(V1, piRNA_reg_ex))
piRNA_reads %>%
  mutate(Length = as_factor(Length)) %>%
  group_by(Length) %>%
  #summarise(Read) %>%
  ggplot() +
  geom_bar(mapping = aes(x = Length))+
  scale_y_continuous(labels = scales::comma)+
  theme minimal()+
  ggtitle("COLO205_dil_A_NT_1_piRNA_reads" %>% basename %>% str_replace("allRNA.txt", "all_RNA"))+
  coord_flip()
key_mIrna_pIrna <- gtf_piB_RCentr %>%
  as_tibble() %>%
  distinct(gene_id, .keep_all = T) %>%
  select(gene_id,seq_RNA,gene_type)
test_mut_reads <- reads_piRNA %>%
  #head(1000) %>%
  filter(str_detect(sRNAs, miRNA_reg_ex)) %>%
  select(Read, sRNAs, read_sequence) %>%
  separate(sRNAs, str_c("V",1:12),
           extra = "merge", fill = "right", sep = ",") %>%
  pivot_longer(cols = starts_with("V"),
               names_to = "alignment", values_to = "sRNAs", values_drop_na = T) %>%
  left_join(key_mIrna_pIrna, by = c("sRNAs" = "gene_id"))
test_mut_reads %>% filter(gene_type %in% c("miRNA", "piRNA"))
test_mut_reads %>%
  group_by(Read) %>%
# different way -----
mutate_all(~replace(., is.na(.), "SS_22"))
get_gene_type <- function(x){key_mIrna_pIrna %>%
```

```
filter(gene_id== x) %>%
.$seq_RNA}

test_mut_reads %>% head %>%
  mutate_at(vars(starts_with("V")), ~map_chr(.,get_gene_type))
```

# 19. piRNA Targets

```
library(plyranges)
library(tidyverse)
#load the gtf of genes mm10
ensembl_mm10 <- read_gff2("mouse_data/Mus_musculus.GRCm38.100.gtf.gz") %>%
  as_tibble() %>%
  select(type, gene_id, gene_name, gene_biotype, transcript_id, transcript_biotype, protein_id) %>%
  arrange(protein_id) %>%
  distinct(gene_id, transcript_id, .keep_all = T) %>%
 filter(!is.na(transcript_id))
# Load the targets
targets <- list.files(path = "mouse_data/targets_piRNA_mm10", pattern = ".UTR|CDS",full.names = T) %>%
  set_names(. %>% basename() %>% str_remove(".txt")) %>%
  map(~read_tsv(.x,col_names = F)
  ) %>%
 bind_rows(.id = "id") %>%
  mutate(
    condition = case_when(
                          is.na(X8) ~ "Complete",
                          str_detect(X8,",",negate = T) ~"MissMatch_1",
                          TRUE ~ "multiple_MissMatch")
   ) %>%
  select(id, piRNA = X1, X3, condition) %>%
  separate(X3, c("gene_id", "gene_id_v", "transcript_id", "transcript_id_v") , sep = "\\|") %>%
  select(-c("gene_id_v","transcript_id_v")) %>%
  distinct(piRNA, transcript_id, .keep_all = T) %>%
  left_join(ensembl_mm10) %>%
  select(piRNA, condition, gene_name, transcript_id, everything()) %>%
  arrange(piRNA, transcript id, gene name)
targets all %>% write tsv("mouse data/gene targets piRNA mm10.txt")
```