Quality Control for bulk RNA-seq data

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<pre>library(tibble) library(tidyr) library(dplyr) library(rtracklayer)</pre>	
# load function from local files	

1. Read the count data

In this section, we will read the raw count data from the synaptosomes_bulkRNA folder. The data is stored in the format of V1-1_final_gene_with_names.csv and T1-1_final_gene_with_names.csv. We will read the data and merge them into a single table. The final table will be stored in results/01-QC/synaptosomes_bulkRNA_counts_raw.csv.

```
sample_list \leftarrow c(paste0("V", rep(1:10, each = 2), "-", 1:2),
                  paste0("T", rep(1:10, each = 2), "-", 1:2))
condition_list <- c(rep("control", 10), rep("treated", 10))</pre>
counts_ref <- read.csv(here::here("data", "synaptosomes_bulkRNA",</pre>
                                     "raw", "V1-1 final gene with names.csv"),
                         header = TRUE)
counts <- tibble()</pre>
for (i in 1:length(sample_list)) {
  sample <- sample_list[i]</pre>
  gene_file <- here::here("data", "synaptosomes_bulkRNA", "raw",</pre>
                            pasteO(sample, "_final_gene_with_names.csv"))
  gene <- read.csv(gene_file, header = TRUE)</pre>
  if (i == 1) {
    counts <- gene[, c("GeneID", "GeneName")]</pre>
    counts <- counts %>% mutate(!!sample := gene$Count)
  } else {
    temp_counts <- gene[, c("GeneID", "Count")]</pre>
    colnames(temp_counts)[2] <- sample</pre>
    counts <- merge(counts, temp_counts, by = "GeneID")</pre>
```

2. Map the gene name.

In this section, we will map the gene name to the gene ID using the GTF file. Since some gene may have different gene name, we will check if the gene name is unique. If not, we will find the gene that has more than one gene name. And merge their counts. The gene annotation comes from the the file Homo_sapiens.GRCh38.106.gtf. The final table will be stored in results/01-QC/synaptosomes bulkBNA counts cleaned.csv.

```
The final table will be stored in results/01-QC/synaptosomes_bulkRNA_counts_cleaned.csv.
# read the gtf file
gtf data <- import(here::here("data", "ref", "Homo sapiens.GRCh38.106.gtf"))
gtf_df <- as.data.frame(gtf_data)</pre>
gtf_genes <- gtf_df %>%
  filter(type == "gene") %>%
  select(gene_id, gene_name)
colnames(gtf_genes) <- c("GeneID", "GeneName")</pre>
# check the annotation format
head(gtf_genes)
##
              GeneID GeneName
## 1 ENSG00000186827 TNFRSF4
## 2 ENSG00000186891 TNFRSF18
## 3 ENSG00000160072
                        ATAD3B
## 4 ENSG00000260179
                          <NA>
## 5 ENSG00000234396
                          <NA>
## 6 ENSG00000225972 MTND1P23
gtf_gene <- na.omit(gtf_genes)</pre>
# check if the gene is unique
unique_genename <- length(unique(gtf_genes$GeneName)) == nrow(gtf_genes)
print(paste("GeneName is unique:", unique_genename))
## [1] "GeneName is unique: FALSE"
# Find the genes that appear more than once
duplicate_genes <- gtf_genes$GeneName[duplicated(gtf_genes$GeneName)]</pre>
duplicate_genes <- unique(duplicate_genes)</pre>
# Correct sprintf statement
print(sprintf("There are %d genes with duplicate gene names",
              length(duplicate_genes)))
```

[1] "There are 68 genes with duplicate gene names"

```
# Merge counts data with GTF information
merged_data <- counts %>%
 left join(gtf genes, by = "GeneID")%>%
  select(-GeneName.x,-GeneID)
names(merged_data) [names(merged_data) == "GeneName.y"] <- "gene"</pre>
# find the same gene
aggregated_data <- merged_data %>%
  group_by(gene) %>%
  summarise(across(everything(), \(x) sum(x, na.rm = TRUE)))
# Using sprintf (Recommended for better formatting)
print(sprintf("The shape of the count matrix is: %d x %d",
              dim(aggregated_data)[1], dim(aggregated_data)[2]))
## [1] "The shape of the count matrix is: 39853 \times 41"
# clean the NA in gene
aggregated_data <- aggregated_data[complete.cases(aggregated_data), ]</pre>
# check if here is any NA in gene
print(sprintf("There are %d genes with NA gene name",
              sum(is.na(aggregated_data$gene))))
## [1] "There are O genes with NA gene name"
write.csv(aggregated data, here::here("data", "synaptosomes bulkRNA",
                                       "synaptosomes_bulkRNA_counts_cleaned.csv"),
          row.names = FALSE)
```

3. make gene refernce

ALso, we will make a reference table for the gene length. The reference table will be stored in results/01-QC/gene_lengths.csv.

sessionInfo()

```
## R version 4.4.0 (2024-04-24)
## Platform: aarch64-apple-darwin20
## Running under: macOS Sonoma 14.3.1
## Matrix products: default
## BLAS:
         /Library/Frameworks/R.framework/Versions/4.4-arm64/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.4-arm64/Resources/lib/libRlapack.dylib;
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## time zone: America/New_York
## tzcode source: internal
##
## attached base packages:
## [1] stats4
                 stats
                           graphics grDevices utils
                                                          datasets methods
## [8] base
##
## other attached packages:
## [1] rtracklayer_1.64.0
                             GenomicRanges_1.56.2 GenomeInfoDb_1.40.1
## [4] IRanges_2.38.1
                             S4Vectors_0.42.1
                                                  BiocGenerics_0.50.0
## [7] knitr_1.49
                             lubridate_1.9.4
                                                   forcats_1.0.0
## [10] stringr_1.5.1
                             dplyr_1.1.4
                                                  purrr_1.0.4
                                                  tibble_3.2.1
## [13] readr_2.1.5
                             tidyr_1.3.1
## [16] ggplot2_3.5.1
                             tidyverse_2.0.0
##
## loaded via a namespace (and not attached):
## [1] SummarizedExperiment_1.34.0 gtable_0.3.6
## [3] rjson_0.2.23
                                    xfun_0.51
## [5] lattice_0.22-6
                                    Biobase_2.64.0
## [7] tzdb_0.4.0
                                    vctrs_0.6.5
## [9] tools_4.4.0
                                    bitops_1.0-9
                                    curl_6.2.1
## [11] generics_0.1.3
## [13] parallel_4.4.0
                                    pkgconfig_2.0.3
## [15] Matrix_1.7-2
                                    lifecycle_1.0.4
## [17] GenomeInfoDbData_1.2.12
                                    compiler_4.4.0
## [19] Rsamtools_2.20.0
                                    Biostrings_2.72.1
## [21] munsell_0.5.1
                                    codetools_0.2-20
## [23] htmltools_0.5.8.1
                                    RCurl_1.98-1.16
## [25] yaml_2.3.10
                                    pillar_1.10.1
## [27] crayon_1.5.3
                                    BiocParallel_1.38.0
## [29] DelayedArray_0.30.1
                                    abind_1.4-8
## [31] tidyselect_1.2.1
                                    digest 0.6.37
## [33] stringi_1.8.4
                                    restfulr_0.0.15
## [35] rprojroot_2.0.4
                                    fastmap_1.2.0
## [37] grid_4.4.0
                                    here_1.0.1
## [39] SparseArray_1.4.8
                                    colorspace_2.1-1
                                    magrittr_2.0.3
## [41] cli_3.6.4
## [43] S4Arrays_1.4.1
                                    XML_3.99-0.18
## [45] withr_3.0.2
                                    scales_1.3.0
## [47] UCSC.utils_1.0.0
                                    timechange_0.3.0
                                    XVector_0.44.0
## [49] rmarkdown_2.29
```

## [51]	httr_1.4.7	matrixStats_1.5.0
## [53]	hms_1.1.3	evaluate_1.0.3
## [55]	BiocIO_1.14.0	rlang_1.1.5
## [57]	glue_1.8.0	rstudioapi_0.17.1
## [59]	jsonlite_1.9.0	R6_2.6.1
## [61]	MatrixGenerics_1.16.0	GenomicAlignments_1.40.0
## [63]	zlibbioc_1.50.0	