Differential Expression Analysis for bulk RNA-seq data

In 22q patient, Vehicle condition vs TTX condition

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Ses	Session information	
lib lib	prary(tibble) prary(tidyr) prary(dplyr) prary(rtracklayer)	
	<pre>load function from local files urce(here::here("source", "DEG_functions.R"))</pre>	

1. Read the count data

In this section, we will read the clean count data from the synaptosomes_bulkRNA folder. We will read the data and merge them into a single table. The final table will be stored in results/02-DEG-Vehicle/synaptosomes_bulkRNA_counts

```
input_count <- read.csv(here::here("data", "synaptosomes_bulkRNA",</pre>
                                         "synaptosomes_bulkRNA_counts_cleaned.csv"))
counts <- as.data.frame(input_count) %>%
  column_to_rownames(var = "gene")
colnames(counts) <- gsub("_", "-", colnames(counts))</pre>
# raw sample list
sample_list_raw <- read.csv(here::here("data", "synaptosomes_bulkRNA",</pre>
                                         "sample_info_22q.csv")) %>%
                     mutate(condition = paste0(Diagosis, "_", Treatment))
# Ensure the column names of counts exist in Sample.name
new_colnames <- sample_list_raw$Label[match(colnames(counts), sample_list_raw$Sample.name)]</pre>
# Assign new column names
colnames(counts) <- new_colnames</pre>
# sort the columns by the colname
condition_list <- data.frame(</pre>
  group =sample_list_raw$condition
)
row.names(condition_list) <- sample_list_raw$Label</pre>
counts<- counts[, rownames(condition_list)]</pre>
# load the 22q gene_list
target_22q_gene <- read.csv(here::here("data","ref" ,"22q_gene_2024_10_17.csv"))</pre>
target_gene <- target_22q_gene$gene</pre>
target_gene <- target_gene[1:(length(target_gene) - 4)]</pre>
gene_name_mapping<- readRDS(here::here("data", "ref" , "gene_name_mapping.rds"))</pre>
```

2. Differential expression analysis

In this section, we will perform differential expression analysis using DESeq2. We will compare the 22q vs Control in the vehicle condition. The results will be stored in results/02-DEG-Vehicle/DESeq2_results.csv.

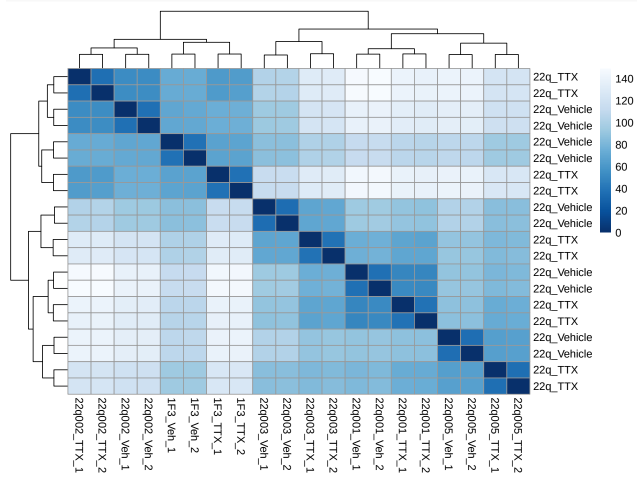
```
# Init the result folder structure for the result
result_folder = file.path("results", "02-DEG-22q")
Result_folder_structure(result_folder)

# load the comparison group information
reference_group <- "22q_Vehicle"
compare_group <- "22q_TTX"</pre>
```

```
filter_sample_info <- condition_list %>%
  filter(group %in% c(reference_group, compare_group))
filter_counts <- counts[, rownames(filter_sample_info)]</pre>
# Run the DESeq2 analysis
dds_obj <- DEAnalysis(counts =filter_counts,</pre>
                       reference_group = reference_group,
                       compare group = compare group,
                        condition_list = filter_sample_info,
                       target_gene = target_gene,
                       result_folder = result_folder)
res <- results(dds_obj)</pre>
resOrdered <- res[order(res$padj), ]
# omit the NA values
resOrdered <- resOrdered[!is.na(resOrdered$padj),]</pre>
dds_obj <- dds_obj[rownames(resOrdered),]</pre>
write.csv(resOrdered, file.path(result_folder, "02-DEG", "01_all_gene_results.csv"))
# DEG with log2fc > 1 and padj < 0.05
deg_1 <- resOrdered %>% as.data.frame() %>% rownames_to_column(var = "gene") %>%
 filter(padj < 0.05 & abs(log2FoldChange) > 1) %>% arrange(padj)
deg 1 <- deg 1[!is.na(deg 1$padj),]
write.csv(deg_1, file.path(result_folder, "02-DEG", "02_DEG_log2fc_1.csv"), row.names = FALSE)
# DEG with log2fc > 1.5 and padj < 0.05
deg_1.5 <- resOrdered %>% as.data.frame() %>% rownames_to_column(var = "gene") %>%
 filter(padj < 0.05 & abs(log2FoldChange) > 1.5) %>% arrange(padj)
deg_1.5 <- deg_1.5 [!is.na(deg_1.5 $padj),]
write.csv(deg_1.5 , file.path(result_folder, "02-DEG", "03_DEG_log2fc_1_5.csv"), row.names = FALSE)
print("DEG analysis is done")
## [1] "DEG analysis is done"
# Save the normalized counts
normalized_counts <- counts(dds_obj, normalized = TRUE)</pre>
write.csv(normalized_counts, file.path(result_folder, "02-DEG", "DESeq2_normalized_counts.csv"))
```

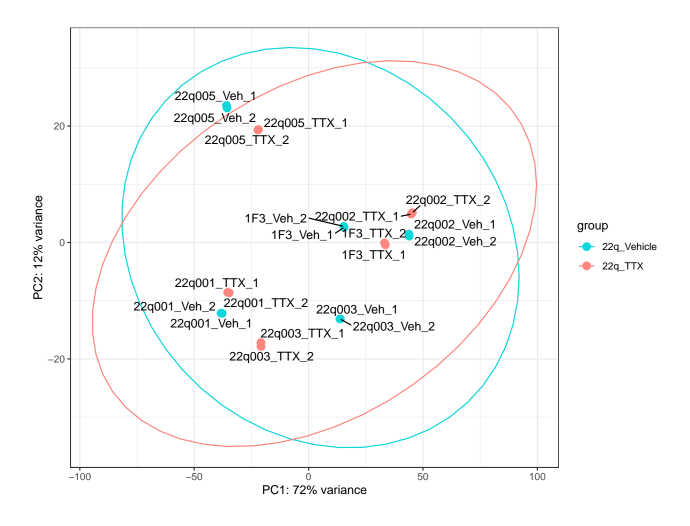
3. Visualization for reuslt

(1) Sample information



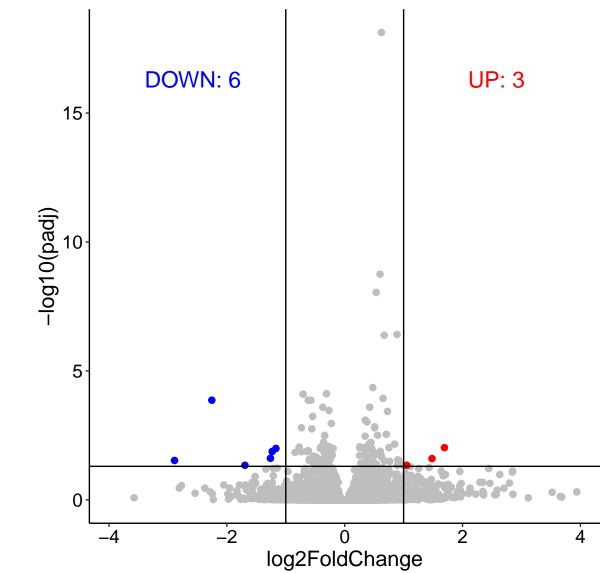
[1] "Sample distance heatmap is done"

[1] "PCA plot is done"



(2) DEG visualization - Volcano plot and Heatmap

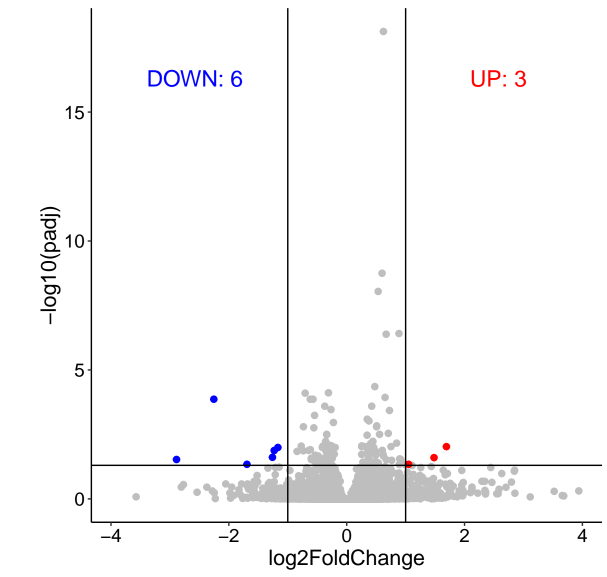
[1] "Volcano plot for 02_volcano_plot_log2fc_1"



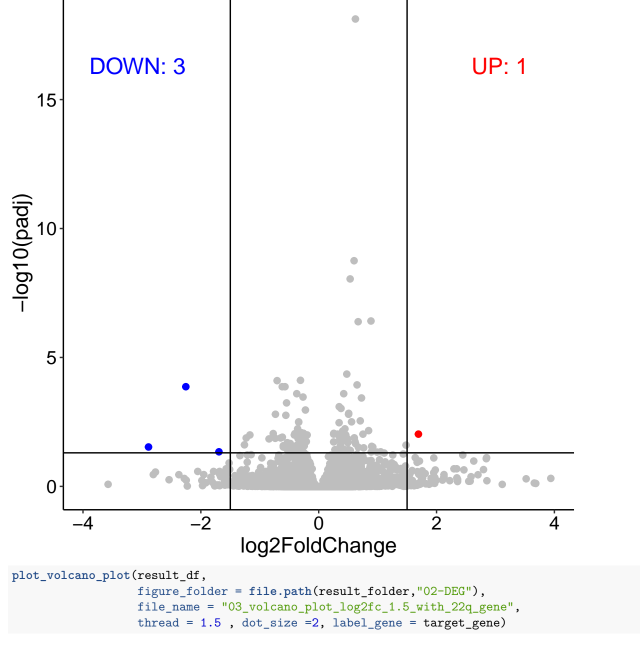
plot_volcano_plot(result_df,

```
figure_folder = file.path(result_folder,"02-DEG"),
file_name = "02_volcano_plot_log2fc_1_with_22q_gene",
thread = 1 , dot_size =2, label_gene = target_gene)
```

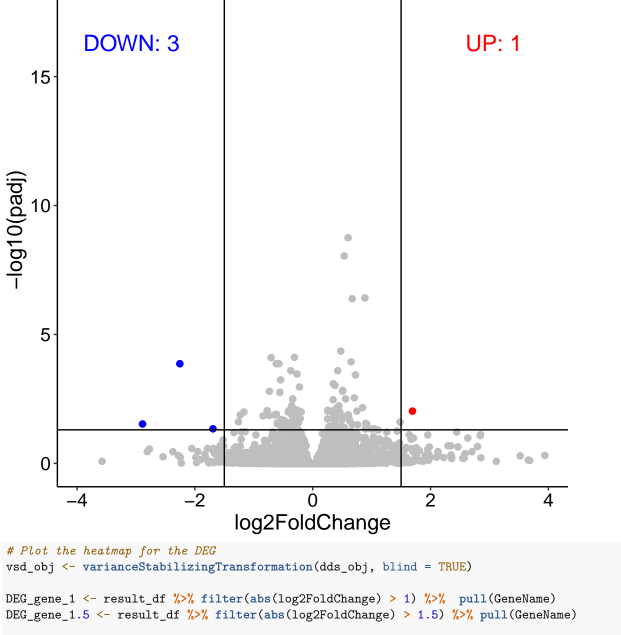
[1] "Volcano plot for 02_volcano_plot_log2fc_1_with_22q_gene"



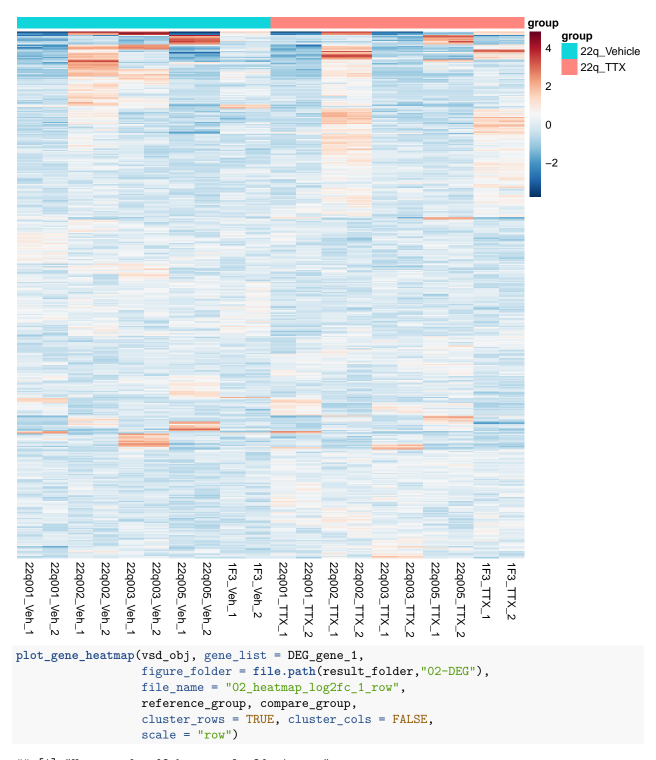
[1] "Volcano plot for 03_volcano_plot_log2fc_1.5"



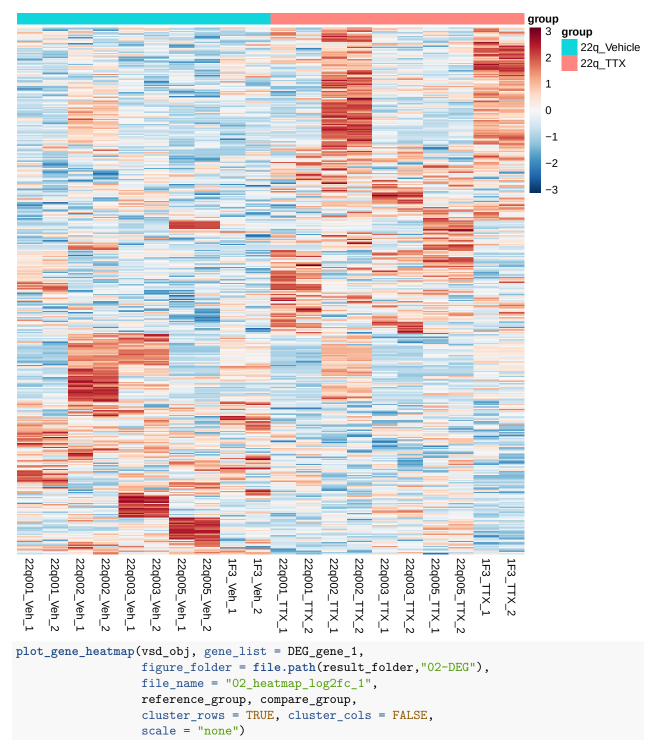
[1] "Volcano plot for 03_volcano_plot_log2fc_1.5_with_22q_gene"



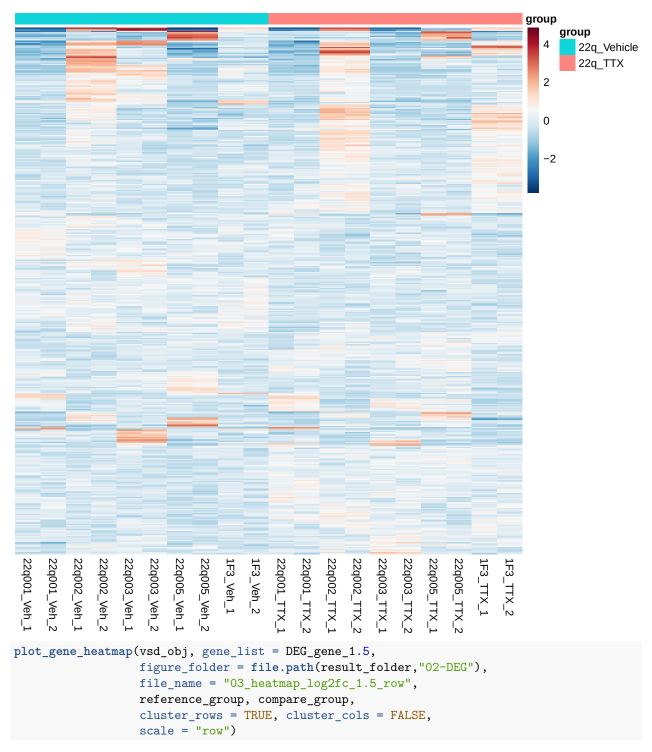
[1] "Heatmap for 02_heatmap_log2fc_1 "



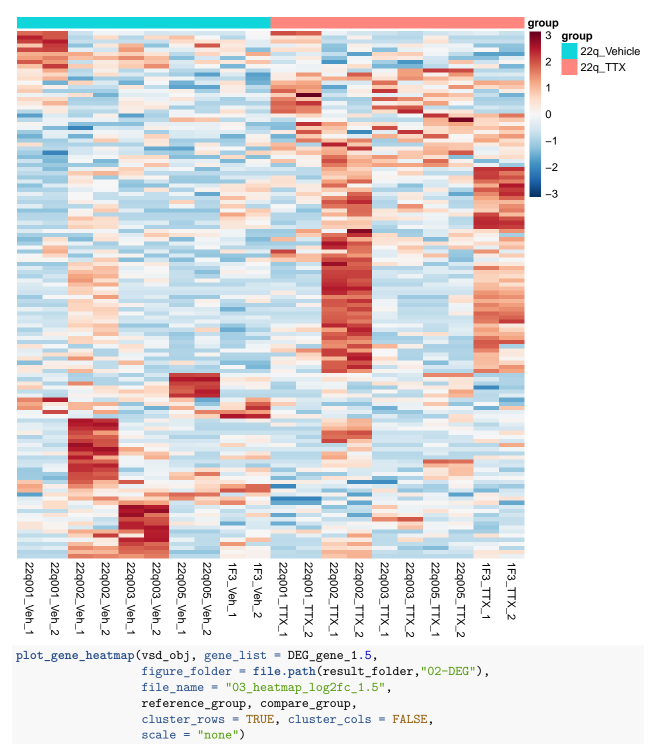
[1] "Heatmap for 02_heatmap_log2fc_1_row "



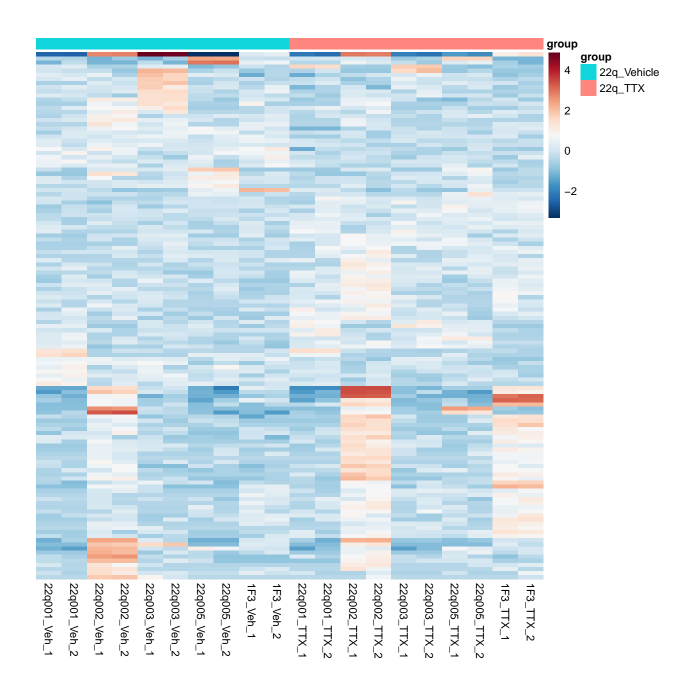
[1] "Heatmap for 02_heatmap_log2fc_1 "



[1] "Heatmap for 03_heatmap_log2fc_1.5_row "

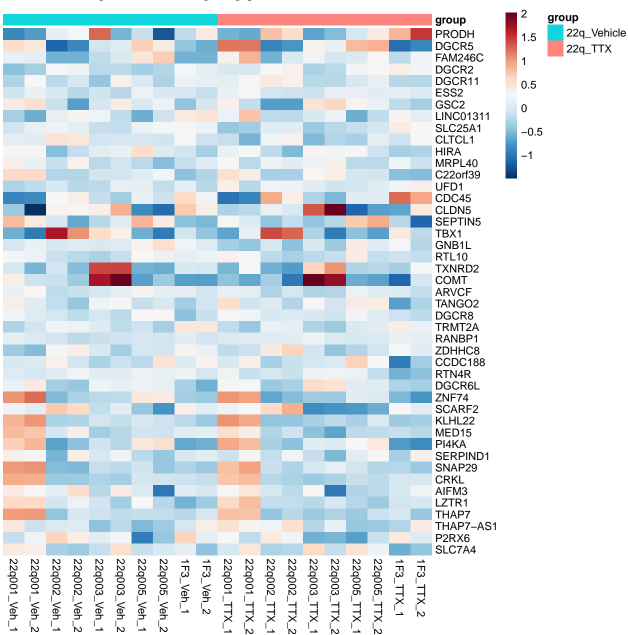


[1] "Heatmap for 03_heatmap_log2fc_1.5 "

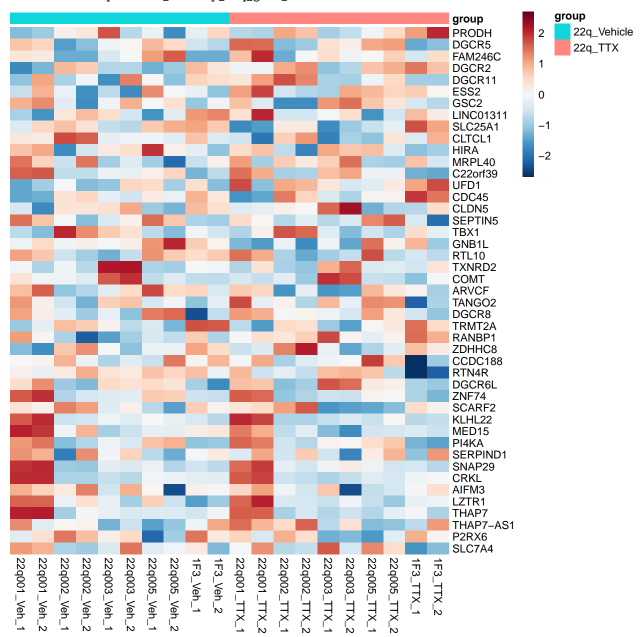


(3) 22q gene visualization - Heatmap

[1] "Heatmap for O1_heatmap_22q_gene "



[1] "Heatmap for O1_heatmap_22q_gene_row "

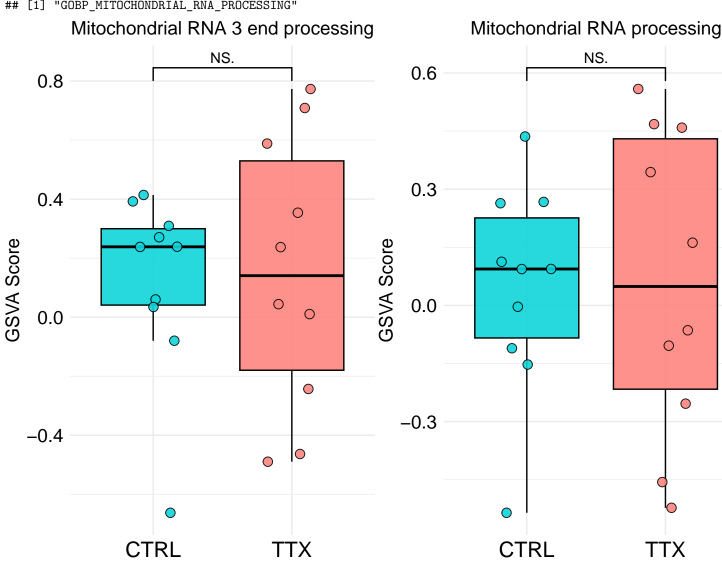


4. GSVA analysis

```
# # The following code is used to generate the GSVA matrix , only need to run once
# gmxFile <- here::here("data", "ref", "c5.go.v2023.1.Hs.symbols.gmt")</pre>
# go_list <- getGmt(gmxFile)</pre>
# geneset <- go_list</pre>
# dat <- as.matrix(counts)</pre>
# gsvapar <- gsvaParam(dat, geneset, maxDiff=TRUE)</pre>
# qsva es <- qsva(qsvapar)</pre>
# gsva_matrix <- as.data.frame(gsva_es)</pre>
# # save the result
 \begin{tabular}{ll} \# \ write. \ csv(gsva\_matrix, \ file.path(result\_folder,"04-GSVA", "01\_GSVA\_matrix.csv")) \\ \end{tabular} 
gsva_matrix <- read.csv(file.path(result_folder, "04-GSVA", "01_GSVA_matrix.csv"),</pre>
                          row.names = 1)
colnames(gsva_matrix) <- sub("^X", "", colnames(gsva_matrix))</pre>
condition_list_label <- condition_list %>%
                          filter(group %in% c(reference_group, compare_group)) %>%
  mutate(group = recode(group,
                          "22q Vehicle" = "CTRL",
                          "22q TTX" = "TTX"))
reference_group_short = "CTRL"
compare_group_short = "TTX"
# plot the heatmap for the GSVA result
pathway_list <- read.csv(here::here("data", "ref", "focus-pathway_2024_10_03.csv"))</pre>
# # plot for all pathway
# for (i in 1:nrow(pathway_list)){
# if (i %% 10 == 0) print(i)
# pathway name <- pathway list$pathway[i]</pre>
   plot_gsva_boxplot(gsva_matrix,
#
                        condition_list_label =condition_list_label,
#
                        pathway_name = pathway_name,
#
                        figure_folder = file.path(result_folder, "04-GSVA", "Boxplot"),
#
                        file_name = pasteO("GSVA_", pathway_name),
#
                        fig.height = 6, fig.width = 4,
#
                        reference_group =reference_group_short ,
#
                        compare_group = compare_group_short)
#
# }
# plot for the focus pathway
for (i in 1:2){
```

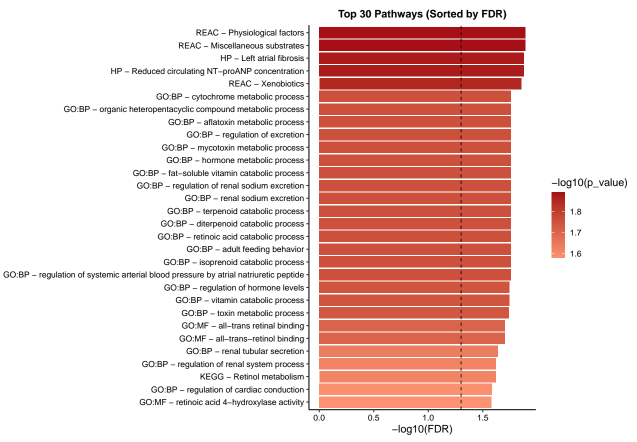
```
pathway_name <- pathway_list$pathway[i]</pre>
  print(pathway_name)
  p<-plot_gsva_boxplot(gsva_matrix,</pre>
                      condition_list_label =condition_list_label,
                      pathway_name = pathway_name,
figure_folder = file.path(result_folder,"04-GSVA","Boxplot"),
                      file_name = pasteO("GSVA_", pathway_name),
                      fig.height = 6, fig.width = 4,
                      reference_group =reference_group_short ,
                      compare_group = compare_group_short)
  print(p)
}
```

- ## [1] "GOBP_MITOCHONDRIAL_RNA_3_END_PROCESSING"
- ## [1] "GOBP_MITOCHONDRIAL_RNA_PROCESSING"

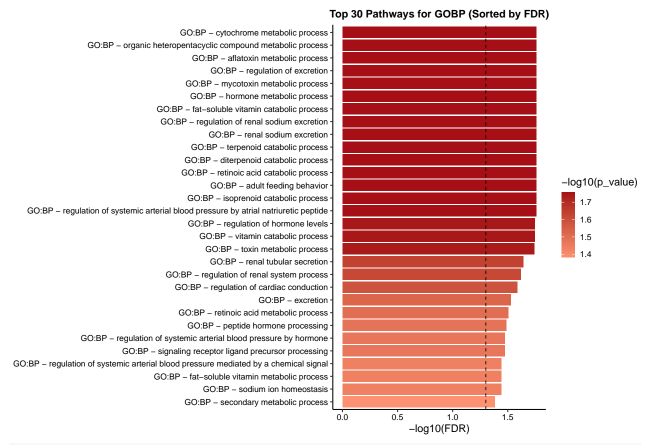


5. Pathway Enrichment Analysis

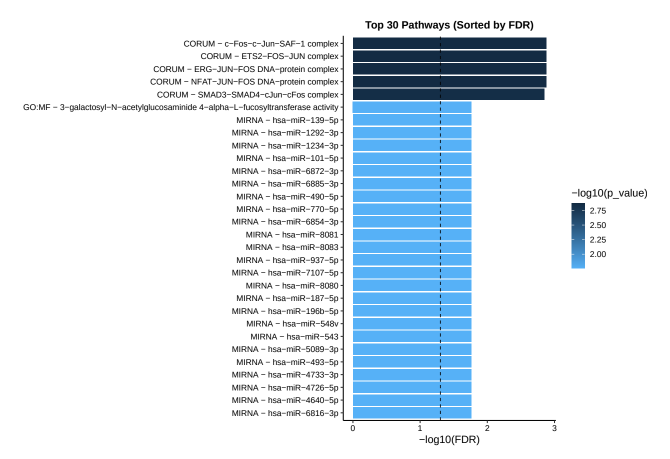
[1] "Enrichment analysis for 01-DEG_1.0_up "



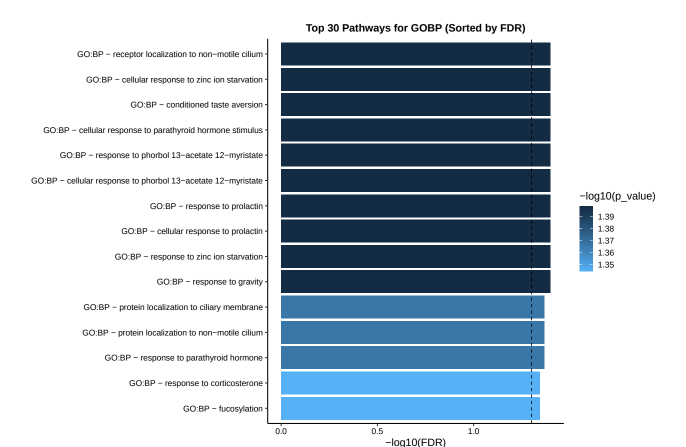
[1] "Enrichment analysis for GOBP 01-DEG_1.0_up "



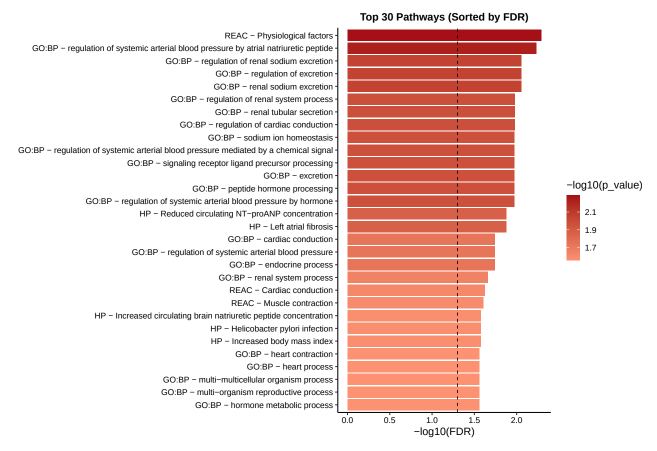
[1] "Enrichment analysis for 01-DEG_1.0_down "



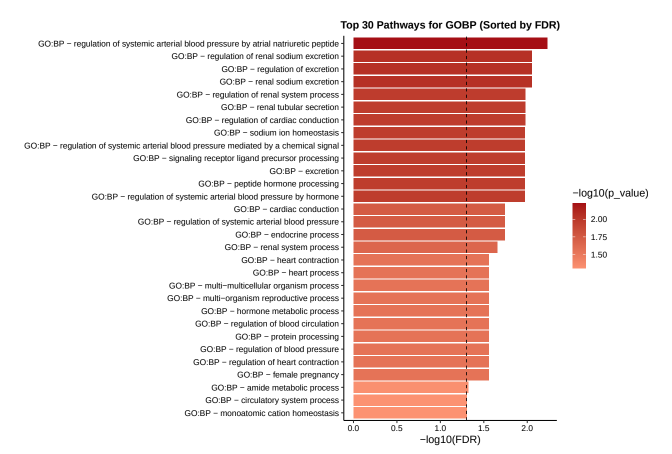
[1] "Enrichment analysis for GOBP 01-DEG_1.0_down "



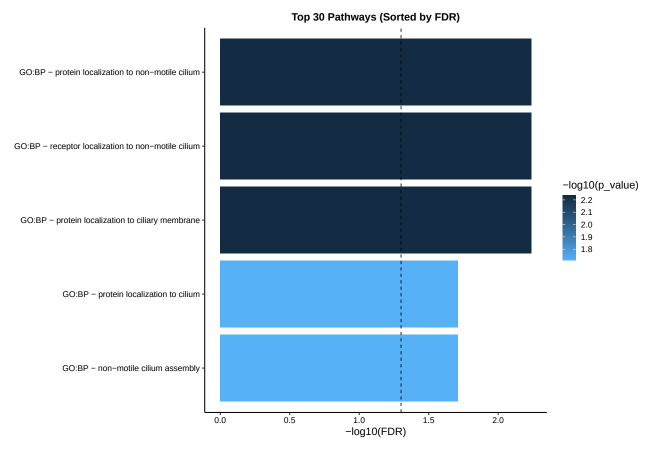
[1] "Enrichment analysis for 02-DEG_1.5_up "



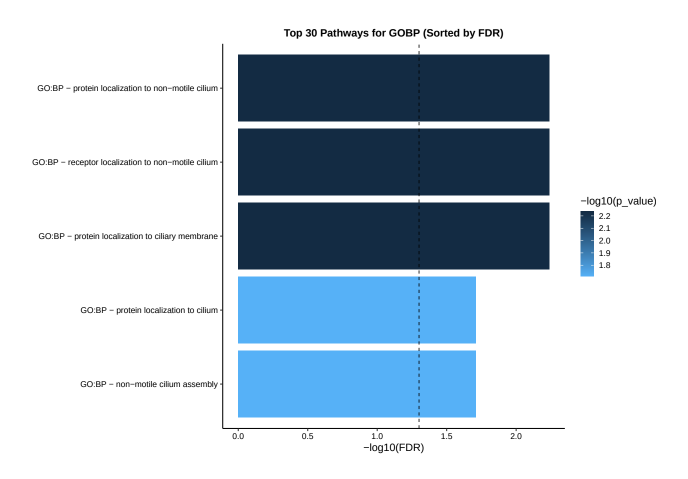
[1] "Enrichment analysis for GOBP 02-DEG_1.5_up "



[1] "Enrichment analysis for 02-DEG_1.5_down "



[1] "Enrichment analysis for GOBP 02-DEG_1.5_down "



Session information

##

[1] SpatialExperiment_1.14.0

[3] progress_1.2.3

```
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] parallel stats4
                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
##
## other attached packages:
## [1] GSEABase_1.66.0
                                    graph_1.82.0
## [3] annotate_1.82.0
                                    XML_3.99-0.18
## [5] extrafont_0.19
                                    ggsignif_0.6.4
## [7] patchwork_1.3.0
                                    decoupleR_2.10.0
## [9] GSVA_1.52.3
                                    BiocParallel_1.38.0
## [11] edgeR_4.2.2
                                    limma_3.60.6
## [13] GenomicFeatures_1.56.0
                                    biomaRt_2.60.1
## [15] gprofiler2_0.2.3
                                    RColorBrewer_1.1-3
## [17] data.table_1.16.4
                                    org.Hs.eg.db_3.19.1
## [19] AnnotationDbi_1.66.0
                                    clusterProfiler_4.12.6
## [21] ggfortify_0.4.17
                                    pheatmap_1.0.12
## [23] EnhancedVolcano_1.22.0
                                    ggrepel_0.9.6
## [25] apeglm_1.26.1
                                    DESeq2_1.44.0
## [27] SummarizedExperiment_1.34.0 Biobase_2.64.0
## [29] MatrixGenerics_1.16.0
                                    matrixStats_1.5.0
## [31] reshape2_1.4.4
                                    Matrix_1.7-2
## [33] Signac_1.14.0
                                    Seurat_5.2.1
## [35] SeuratObject_5.0.2
                                    sp_2.2-0
## [37] rtracklayer_1.64.0
                                    GenomicRanges_1.56.2
## [39] GenomeInfoDb_1.40.1
                                    IRanges_2.38.1
## [41] S4Vectors_0.42.1
                                    BiocGenerics_0.50.0
## [43] knitr_1.49
                                    lubridate_1.9.4
## [45] forcats_1.0.0
                                    stringr_1.5.1
## [47] dplyr_1.1.4
                                    purrr_1.0.4
## [49] readr_2.1.5
                                    tidyr_1.3.1
## [51] tibble_3.2.1
                                    ggplot2_3.5.1
## [53] tidyverse_2.0.0
## loaded via a namespace (and not attached):
```

LAPACK v

R.methodsS3_1.8.2 goftest_1.2-3

```
[5] HDF5Array_1.32.1
##
                                     Biostrings_2.72.1
##
     [7] vctrs_0.6.5
                                     spatstat.random_3.3-2
##
     [9] digest_0.6.37
                                     png 0.1-8
   [11] deldir_2.0-4
                                     parallelly_1.42.0
##
##
    [13] magick_2.8.5
                                     MASS_7.3-64
##
   [15] httpuv 1.6.15
                                     qvalue 2.36.0
   [17] withr 3.0.2
                                     xfun 0.51
##
   [19] ggfun_0.1.8
                                     survival_3.8-3
##
    [21] memoise_2.0.1
                                     gson_0.1.0
##
   [23] systemfonts_1.2.1
                                     ragg_1.3.3
   [25] tidytree_0.4.6
                                     zoo_1.8-12
##
   [27] pbapply_1.7-2
                                     R.oo_1.27.0
##
  [29] prettyunits_1.2.0
                                     KEGGREST_1.44.1
##
  [31] promises_1.3.2
                                     httr_1.4.7
##
  [33] restfulr_0.0.15
                                     rhdf5filters_1.16.0
##
   [35] globals_0.16.3
                                     fitdistrplus_1.2-2
##
   [37] rhdf5_2.48.0
                                     rstudioapi_0.17.1
   [39] UCSC.utils 1.0.0
                                     miniUI 0.1.1.1
##
   [41] generics_0.1.3
                                     DOSE_3.30.5
##
   [43] curl 6.2.1
                                     zlibbioc_1.50.0
##
   [45] ScaledMatrix_1.12.0
                                     ggraph_2.2.1
                                     GenomeInfoDbData_1.2.12
  [47] polyclip_1.10-7
##
  [49] SparseArray_1.4.8
                                     xtable_1.8-4
##
   [51] evaluate 1.0.3
                                     S4Arrays_1.4.1
##
  [53] BiocFileCache 2.12.0
                                     hms 1.1.3
  [55] irlba_2.3.5.1
                                     colorspace_2.1-1
##
  [57] filelock_1.0.3
                                     ROCR_1.0-11
##
  [59] reticulate_1.40.0
                                     spatstat.data_3.1-4
##
  [61] magrittr_2.0.3
                                     lmtest_0.9-40
## [63] later_1.4.1
                                     viridis_0.6.5
##
   [65] ggtree_3.12.0
                                     lattice_0.22-6
##
   [67] spatstat.geom_3.3-5
                                     future.apply_1.11.3
##
   [69] scattermore_1.2
                                     shadowtext_0.1.4
##
   [71] cowplot_1.1.3
                                     RcppAnnoy_0.0.22
##
    [73] pillar_1.10.1
                                     nlme_3.1-167
##
  [75] compiler_4.4.0
                                     beachmat_2.20.0
  [77] RSpectra 0.16-2
                                     stringi 1.8.4
## [79] tensor_1.5
                                     GenomicAlignments_1.40.0
##
   [81] plyr_1.8.9
                                     crayon_1.5.3
##
  [83] abind_1.4-8
                                     BiocIO_1.14.0
## [85] gridGraphics_0.5-1
                                     emdbook 1.3.13
## [87] locfit 1.5-9.11
                                     graphlayouts_1.2.2
## [89] bit_4.5.0.1
                                     fastmatch 1.1-6
## [91] textshaping_1.0.0
                                     codetools_0.2-20
## [93] BiocSingular_1.20.0
                                     plotly_4.10.4
## [95] mime_0.12
                                     splines_4.4.0
## [97] Rcpp_1.0.14
                                     fastDummies_1.7.5
  [99] sparseMatrixStats_1.16.0
                                     dbplyr_2.5.0
## [101] Rttf2pt1_1.3.12
                                     blob_1.2.4
## [103] here_1.0.1
                                     fs_1.6.5
## [105] listenv_0.9.1
                                     ggplotify_0.1.2
## [107] statmod 1.5.0
                                     tzdb_0.4.0
## [109] tweenr_2.0.3
                                     pkgconfig_2.0.3
## [111] tools 4.4.0
                                     cachem 1.1.0
```

```
## [113] RSQLite 2.3.9
                                     viridisLite_0.4.2
## [115] DBI_1.2.3
                                     numDeriv_2016.8-1.1
## [117] fastmap 1.2.0
                                     rmarkdown 2.29
## [119] scales_1.3.0
                                     grid_4.4.0
                                     Rsamtools_2.20.0
## [121] ica_1.0-3
## [123] coda 0.19-4.1
                                     dotCall64 1.2
## [125] RANN 2.6.2
                                     farver 2.1.2
## [127] tidygraph_1.3.1
                                     scatterpie_0.2.4
## [129] yaml_2.3.10
                                     cli_3.6.4
## [131] lifecycle_1.0.4
                                     uwot_0.2.2
## [133] mvtnorm_1.3-3
                                     timechange_0.3.0
## [135] gtable_0.3.6
                                     rjson_0.2.23
## [137] ggridges_0.5.6
                                     progressr_0.15.1
## [139] ape_5.8-1
                                     jsonlite_1.9.0
## [141] RcppHNSW_0.6.0
                                     bitops_1.0-9
## [143] bit64_4.6.0-1
                                     Rtsne_0.17
## [145] yulab.utils_0.2.0
                                     spatstat.utils_3.1-2
## [147] bdsmatrix 1.3-7
                                     GOSemSim 2.30.2
## [149] spatstat.univar_3.1-1
                                     R.utils_2.12.3
## [151] lazyeval 0.2.2
                                     shiny 1.10.0
## [153] htmltools_0.5.8.1
                                     enrichplot_1.24.4
## [155] GO.db 3.19.1
                                     sctransform_0.4.1
## [157] rappdirs_0.3.3
                                     tinytex_0.55
## [159] glue_1.8.0
                                     spam 2.11-1
## [161] httr2 1.1.0
                                     XVector 0.44.0
## [163] RCurl_1.98-1.16
                                     rprojroot_2.0.4
## [165] treeio_1.28.0
                                     gridExtra_2.3
                                     igraph_2.1.4
## [167] extrafontdb_1.0
## [169] R6_2.6.1
                                     SingleCellExperiment_1.26.0
## [171] labeling_0.4.3
                                     RcppRoll_0.3.1
## [173] cluster_2.1.8
                                     bbmle_1.0.25.1
## [175] Rhdf5lib_1.26.0
                                     aplot_0.2.4
## [177] DelayedArray_0.30.1
                                     tidyselect_1.2.1
## [179] ggforce_0.4.2
                                     xml2_1.3.6
## [181] future 1.34.0
                                     rsvd 1.0.5
## [183] munsell_0.5.1
                                     KernSmooth_2.23-26
## [185] htmlwidgets 1.6.4
                                     fgsea 1.30.0
## [187] rlang_1.1.5
                                     spatstat.sparse_3.1-0
## [189] spatstat.explore_3.3-4
```