# Differential Expression Analysis for small RNA-seq data $$\operatorname{TTX}$$ contition: 22q vs Control

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#### 1. Read the count data

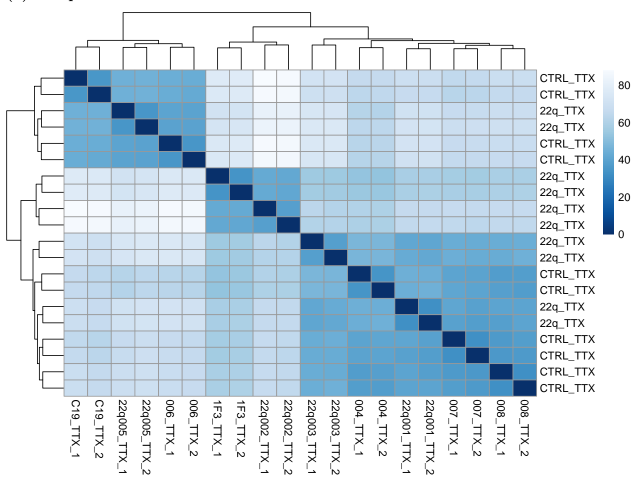
In this section, we will read the clean count data from the synaptosomes\_miRNARNA folder. The data is stored in the format of .csv. We will read the data and filter the columns based on the condition list. The final table will be stored in results/02-DEM-Vehicle/DESeq2\_results.csv.

# 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/01-DEM-TTX/DESeq2\_results.csv.

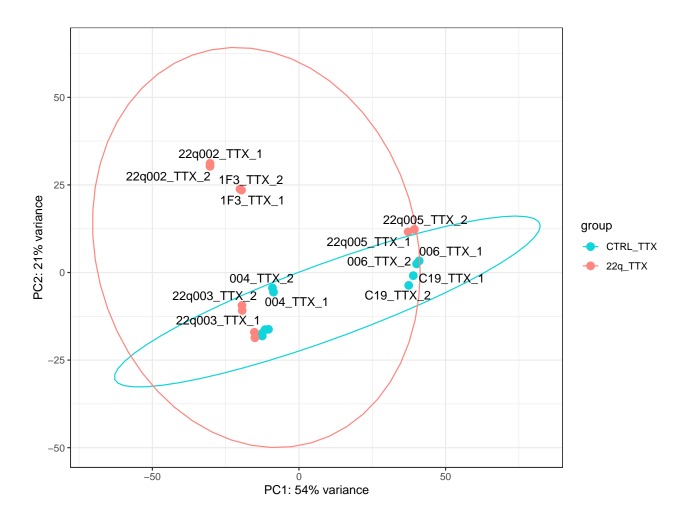
#### 3. Visualization for reuslt

#### (1) Sample information



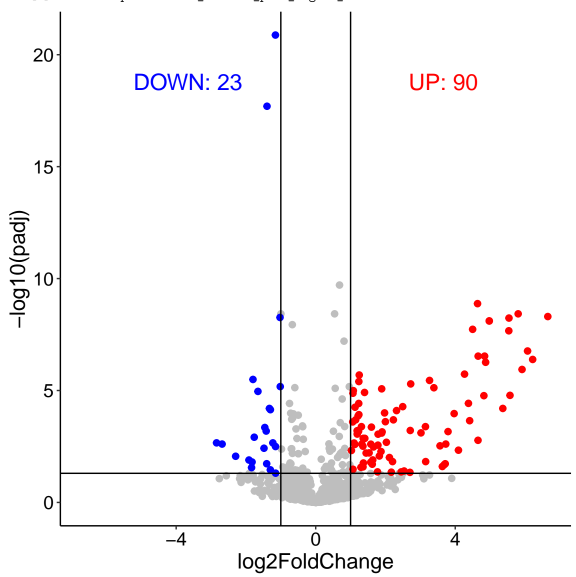
## [1] "Sample distance heatmap is done"

## [1] "PCA plot is done"

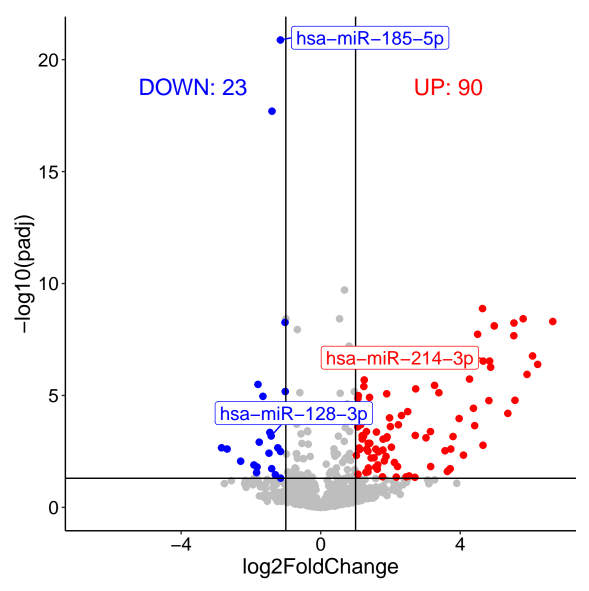


### (2) DEM visualization - Volcano plot and Heatmap

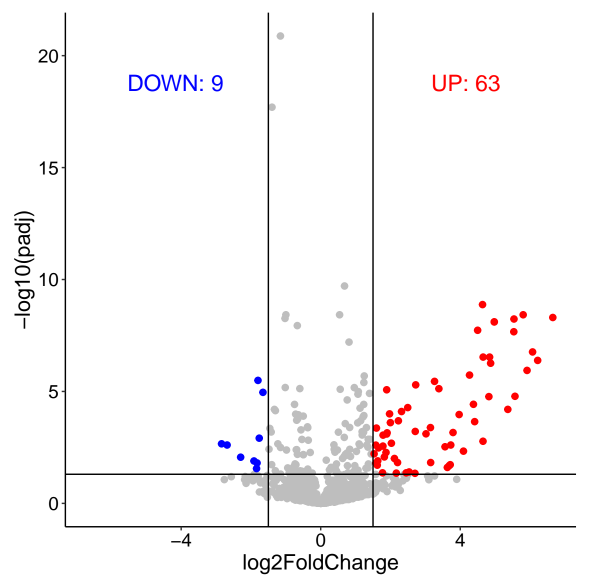
## [1] "Volcano plot for 02\_volcano\_plot\_log2fc\_1"



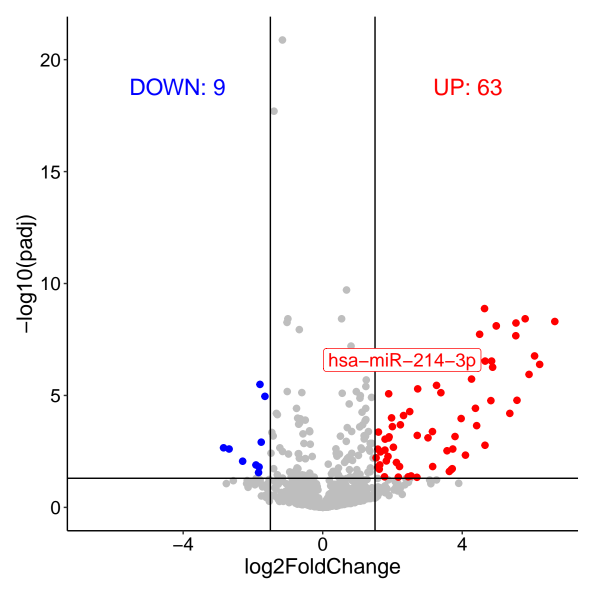
## [1] "Volcano plot for 02\_volcano\_plot\_log2fc\_1\_with\_label\_miRNA"



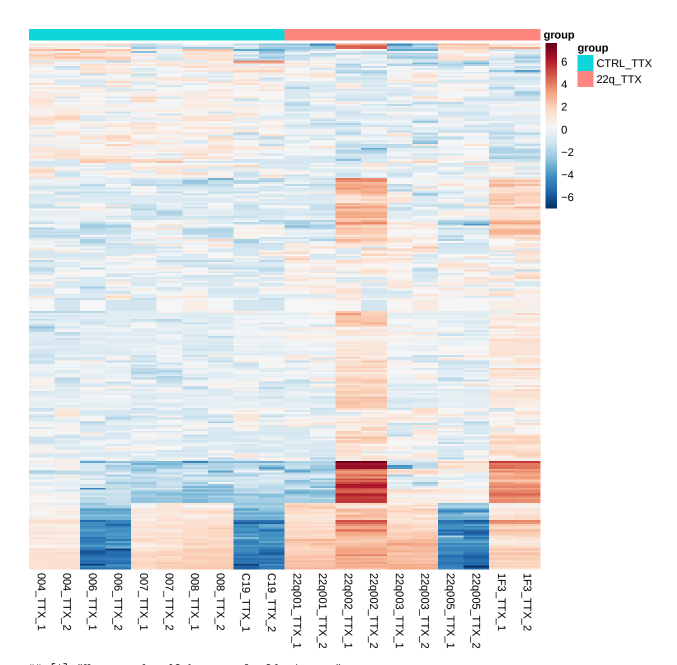
## [1] "Volcano plot for 03\_volcano\_plot\_log2fc\_1.5"



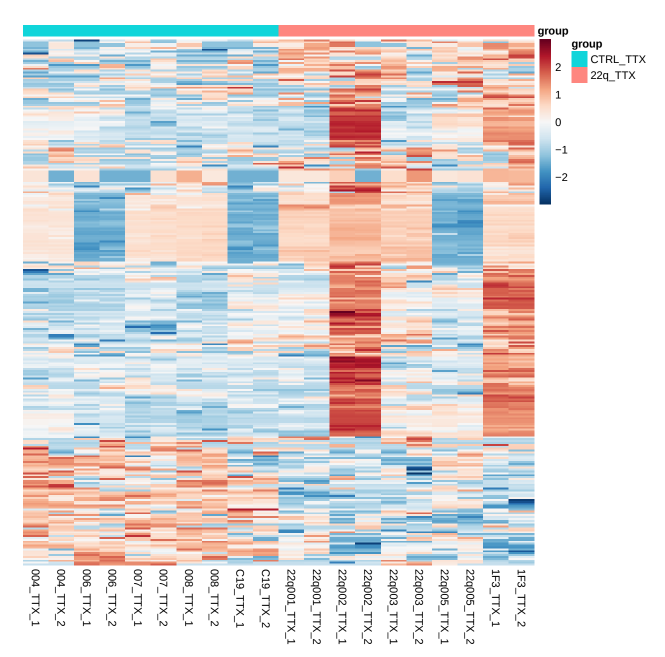
## [1] "Volcano plot for 03\_volcano\_plot\_log2fc\_1.5\_with\_label\_miRNA"



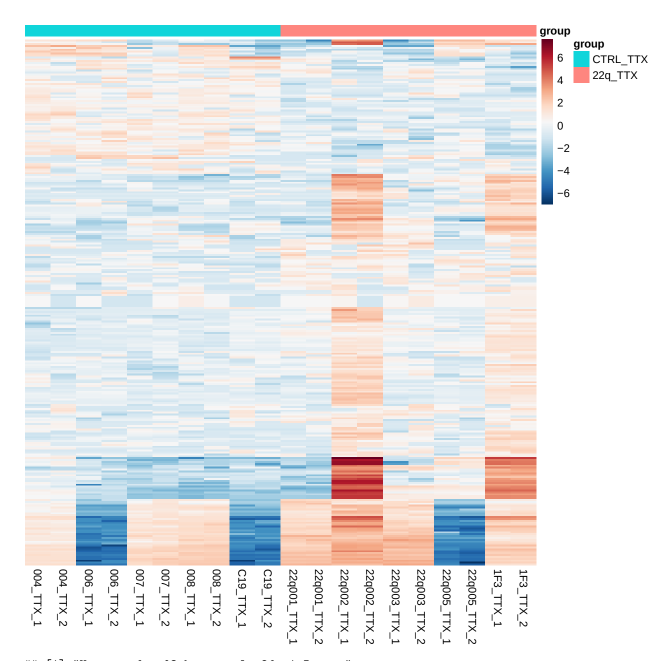
## [1] "Heatmap for 02\_heatmap\_log2fc\_1 "



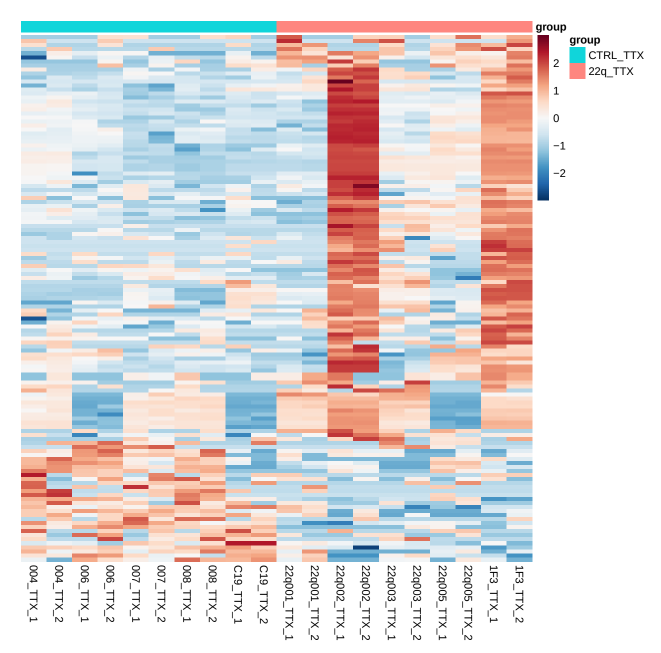
## [1] "Heatmap for 02\_heatmap\_log2fc\_1\_row "



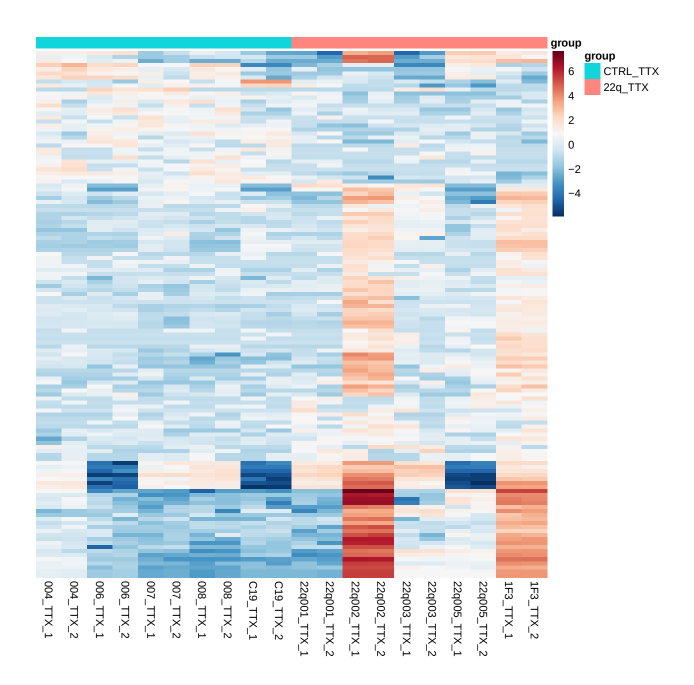
## [1] "Heatmap for 02\_heatmap\_log2fc\_1 "



## [1] "Heatmap for 03\_heatmap\_log2fc\_1.5\_row "

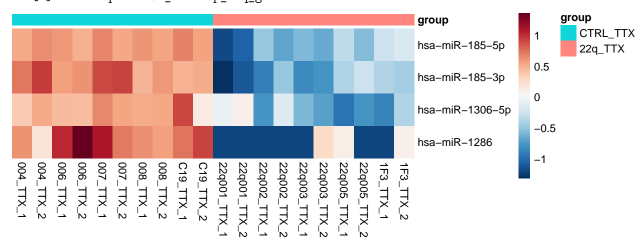


## [1] "Heatmap for 03\_heatmap\_log2fc\_1.5 "

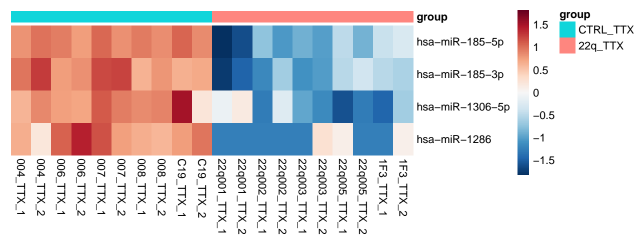


#### (3) 22q miRNA visualization - Heatmap

## [1] "Heatmap for 01\_heatmap\_22q\_gene "



## [1] "Heatmap for O1\_heatmap\_22q\_gene\_row "



#### Session information

```
## R version 4.4.0 (2024-04-24)
## Platform: aarch64-apple-darwin20
## Running under: macOS 15.4
## Matrix products: default
          /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; LAPACK v
## 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
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## [1] parallel stats4
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                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
##
## other attached packages:
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## [3] annotate_1.82.0
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## [5] extrafont_0.19
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## [13] GenomicFeatures_1.56.0
## [15] gprofiler2_0.2.3
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