Package Downloads

```
# Run this chunk top-to-bottom in a fresh R session to ensure all required
# packages are available. It safely skips anything already installed.
cran_repo <- "https://cloud.r-project.org"</pre>
options(repos = c(CRAN = cran_repo))
if (!requireNamespace("BiocManager", quietly = TRUE)) {
  install.packages("BiocManager", repos = cran_repo)
}
BiocManager::install(version = "3.21", ask = FALSE, update = FALSE)
install_if_missing <- function(pkgs, installer) {</pre>
 missing_pkgs <- pkgs[!vapply(pkgs, requireNamespace,</pre>
                               FUN.VALUE = logical(1), quietly = TRUE)]
 if (length(missing_pkgs) > 0) {
    installer(missing_pkgs)
 }
  invisible(NULL)
install_if_missing(
 pkgs = c("devtools", "GenomeInfoDb", "tidyverse"),
  installer = function(pkgs) install.packages(pkgs, repos = cran_repo)
install_if_missing(
 pkgs = c("DESeq2", "org.Hs.eg.db"),
  installer = function(pkgs) BiocManager::install(pkgs, ask = FALSE, update = FALSE)
```

Section 1

Section 1a

```
# Attach the library
library(org.Hs.eg.db)

## Loading required package: AnnotationDbi

## Loading required package: stats4

## Loading required package: BiocGenerics

## Loading required package: generics
```

```
##
## Attaching package: 'generics'
## The following objects are masked from 'package:base':
##
##
       as.difftime, as.factor, as.ordered, intersect, is.element, setdiff,
##
       setequal, union
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, sd, var, xtabs
## The following objects are masked from 'package:base':
##
##
       anyDuplicated, aperm, append, as.data.frame, basename, cbind,
##
       colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find,
##
       get, grep, grepl, is.unsorted, lapply, Map, mapply, match, mget,
##
       order, paste, pmax, pmax.int, pmin, pmin.int, Position, rank,
##
       rbind, Reduce, rownames, sapply, saveRDS, table, tapply, unique,
##
       unsplit, which.max, which.min
## Loading required package: Biobase
## Welcome to Bioconductor
##
##
       Vignettes contain introductory material; view with
##
       'browseVignettes()'. To cite Bioconductor, see
       'citation("Biobase")', and for packages 'citation("pkgname")'.
##
## Loading required package: IRanges
## Loading required package: S4Vectors
##
## Attaching package: 'S4Vectors'
## The following object is masked from 'package:utils':
##
##
       findMatches
## The following objects are masked from 'package:base':
##
##
       expand.grid, I, unname
##
## Attaching package: 'IRanges'
```

```
## The following object is masked from 'package:grDevices':
##
##
       windows
##
# We will need this so we can use the pipe: %>%
library(magrittr)
\# Create the data folder if it doesn't exist
if (!dir.exists("data")) {
  dir.create("data")
# Define the file path to the plots directory
plots_dir <- "plots"</pre>
# Create the plots folder if it doesn't exist
if (!dir.exists(plots_dir)) {
  dir.create(plots_dir)
}
# Define the file path to the results directory
results_dir <- "results"</pre>
# Create the results folder if it doesn't exist
if (!dir.exists(results_dir)) {
  dir.create(results_dir)
# Define the file path to the data directory
data_dir <- file.path("data", "SRP192714")</pre>
# Declare the file path to the gene expression matrix file
data_file <- file.path(data_dir, "SRP192714.tsv")</pre>
# Read in data TSV file
expression_df <- readr::read_tsv(data_file) %>%
tibble::column_to_rownames("Gene")
## Rows: 43363 Columns: 1022
## -- Column specification ----
## Delimiter: "\t"
## chr
         (1): Gene
## dbl (1021): SRR8907879, SRR8907880, SRR8907881, SRR8907882, SRR8907883, SRR8...
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
# Load refine.bio metadata
refinebio_meta <- readr::read_tsv(file.path(data_dir, "metadata_SRP192714.tsv"))
```

```
## Rows: 1021 Columns: 25
## -- Column specification ------
## Delimiter: "\t"
## chr (11): refinebio_accession_code, experiment_accession, refinebio_organism...
## dbl (3): refinebio_age, refinebio_processor_id, MetaSRA_age
## lgl (11): refinebio_cell_line, refinebio_compound, refinebio_developmental_s...
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
rownames(refinebio_meta) <- refinebio_meta$refinebio_accession_code</pre>
## Warning: Setting row names on a tibble is deprecated.
# Load GEO metadata (CSV)
geo_meta <- readr::read_csv("data/GSE129882_PhenoData.transcript.csv")</pre>
## New names:
## Rows: 261 Columns: 21
## -- Column specification
                                         ----- Delimiter: "," chr
## (15): ...1, Sample, Time, Sex, DENV.at.Inception, DENV.Exposure.at.Time.... dbl
## (6): Patient, Visit, Age, DENV.Infections, lib.size, norm.factors
## i Use 'spec()' to retrieve the full column specification for this data. i
## Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## * ' ' -> ' . . . 1 '
# Merge on 'refinebio_title' (refine.bio) and 'Sample' (GEO), keep 'refinebio_title' as column name
if ("Sample" %in% colnames(geo_meta) && "refinebio_title" %in% colnames(refinebio_meta)) {
 merged meta <- dplyr::left join(refinebio meta, geo meta, by = c("refinebio title" = "Sample"))
} else {
 merged_meta <- refinebio_meta</pre>
# Choose how many samples to keep for this run (set to Inf to keep all samples)
sample_limit <- 25</pre>
all_samples <- colnames(expression_df)</pre>
selected_samples <- all_samples</pre>
if (is.finite(sample_limit)) {
  selected_samples <- all_samples[seq_len(min(sample_limit, length(all_samples)))]</pre>
}
is_trimmed_run <- length(selected_samples) < length(all_samples)</pre>
run_label <- if (is_trimmed_run) paste0("trimmed_", length(selected_samples)) else "full"</pre>
expression_df <- expression_df[, selected_samples, drop = FALSE]</pre>
merged_meta <- merged_meta %>%
  dplyr::filter(refinebio_accession_code %in% selected_samples)
rownames(merged_meta) <- merged_meta$refinebio_accession_code</pre>
```

Warning: Setting row names on a tibble is deprecated.

```
merged_meta_path <- file.path(data_dir, "metadata_SRP192714_merged.tsv")
readr::write_tsv(merged_meta, merged_meta_path)
```

Section 1b

```
# Bring back the "Gene" column in preparation for mapping
expression_df <- expression_df %>%
 tibble::rownames_to_column("Gene")
# Map Ensembl IDs to their first mapped Symbol
gene_symbols <- mapIds(</pre>
 org.Hs.eg.db,
 keys = expression_df$Gene,
 keytype = "ENSEMBL",
 column = "SYMBOL",
 multiVals = "first"
```

'select()' returned 1:many mapping between keys and columns

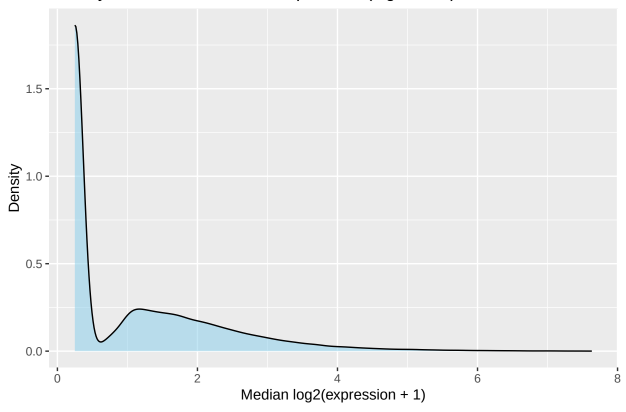
```
# Add the mapped symbols as a new column
expression_df$Symbol <- gene_symbols[expression_df$Gene]</pre>
# Reorder columns to have Gene, Symbol, then the rest
expression_df <- expression_df %>%
  dplyr::select(Gene, Symbol, dplyr::everything(), -Gene)
# Write mapped data frame to output file
readr::write_tsv(expression_df, file.path(
  results dir,
  "SRP192714_Symbols.tsv"
))
```

Section 1c

```
# Get matrix size
matrix_dim <- dim(expression_df)</pre>
cat("Expression matrix dimensions (genes x samples):", matrix_dim[1], "x", matrix_dim[2], "\n")
## Expression matrix dimensions (genes x samples): 43363 x 26
# Number of genes
cat("Number of genes:", nrow(expression_df), "\n")
## Number of genes: 43363
```

```
# Select only numeric columns for log transformation
expr_numeric <- expression_df %>% dplyr::select(where(is.numeric))
# Log-scale the data (add pseudocount to avoid log(0))
log_expr <- log2(expr_numeric + 1)</pre>
# Calculate per-gene median expression
gene_medians <- apply(log_expr, 1, median, na.rm = TRUE)</pre>
# Show summary statistics for gene medians
summary(gene_medians)
                              Mean 3rd Qu.
##
      Min. 1st Qu. Median
                                               Max.
## 0.2486 0.2486 0.2625 1.0755 1.7040 7.6328
# Density plot of per-gene median expression
library(ggplot2)
plot_obj <- ggplot(data.frame(median=gene_medians), aes(x=median)) +</pre>
  geom_density(fill="skyblue", alpha=0.5) +
  labs(title="Density of Per-Gene Median Expression (log2 scale)",
       x="Median log2(expression + 1)",
       y="Density")
# Save plot to the plots directory
plot_path <- file.path(plots_dir, "per_gene_median_density.png")</pre>
ggsave(plot_path, plot=plot_obj, width=6, height=4, dpi=300)
# Also print the plot in the notebook
plot_obj
```

Density of Per-Gene Median Expression (log2 scale)



The dataset contains 43,363 genes measured across 26 samples. The summary statistics show that most genes have low median expression (median ≈ 0.26 on the log2 scale), with a long tail of higher expression values (max ≈ 7.63). From our research, this is typical for transcriptome data, where a small number of genes are highly expressed while the majority have low expression. The density plot visualizes this distribution, showing a peak at lower expression values and a gradual decline towards higher values. Log transformation helps to reduce the impact of extreme values and makes the distribution more interpretable. # Section 2 ## Section 2a-c

```
suppressPackageStartupMessages({
  library(DESeq2)
  library(dplyr)
  library(magrittr)
  library(readr)
  library(tibble)
  library(stringr)
})
if (!exists("results_dir")) {
  results_dir <- "results"</pre>
  if (!dir.exists(results_dir)) dir.create(results_dir, recursive = TRUE)
if (!exists("plots_dir")) {
  plots_dir <- "plots"</pre>
  if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
}
if (!exists("data_dir")) {
```

```
}
if (!exists("expression_df") || !exists("merged_meta")) {
  sample_limit_local <- if (exists("sample_limit")) sample_limit else 25</pre>
  expression_df <- readr::read_tsv(file.path(data_dir, "SRP192714.tsv"), show_col_types = FALSE) %>%
    tibble::column_to_rownames("Gene")
  refinebio_meta <- readr::read_tsv(file.path(data_dir, "metadata_SRP192714.tsv"), show_col_types = FAL
  rownames(refinebio_meta) <- refinebio_meta$refinebio_accession_code</pre>
  all_samples <- colnames(expression_df)</pre>
  selected samples <- all samples
  if (is.finite(sample_limit_local)) {
    selected_samples <- all_samples[seq_len(min(sample_limit_local, length(all_samples)))]</pre>
  }
  expression_df <- expression_df[, selected_samples, drop = FALSE]</pre>
  merged_meta <- refinebio_meta %>%
    dplyr::filter(refinebio_accession_code %in% selected_samples)
  rownames(merged_meta) <- merged_meta$refinebio_accession_code</pre>
  is_trimmed_run <- length(selected_samples) < length(all_samples)</pre>
 run_label <- if (is_trimmed_run) paste0("trimmed_", length(selected_samples)) else "full"</pre>
expr_mat <- expression_df %>% dplyr::select(where(is.numeric))
meta <- merged_meta
common_samples <- intersect(colnames(expr_mat), meta$refinebio_accession_code)</pre>
expr_mat <- expr_mat %>% dplyr::select(all_of(common_samples))
meta <- meta %>%
  dplyr::filter(refinebio_accession_code %in% common_samples)
rownames(meta) <- meta$refinebio_accession_code</pre>
## Warning: Setting row names on a tibble is deprecated.
cts <- as.matrix(expr_mat)</pre>
cts <- cts[, rownames(meta), drop = FALSE]</pre>
cts[cts < 0] <- 0
cts <- round(cts)
stopifnot(ncol(cts) == nrow(meta))
plot_title <- sprintf("PCA of %d Samples (colored by Exposure)", length(common_samples))
plot_file <- paste0(run_label, "_pca_exposure.png")</pre>
```

converting counts to integer mode

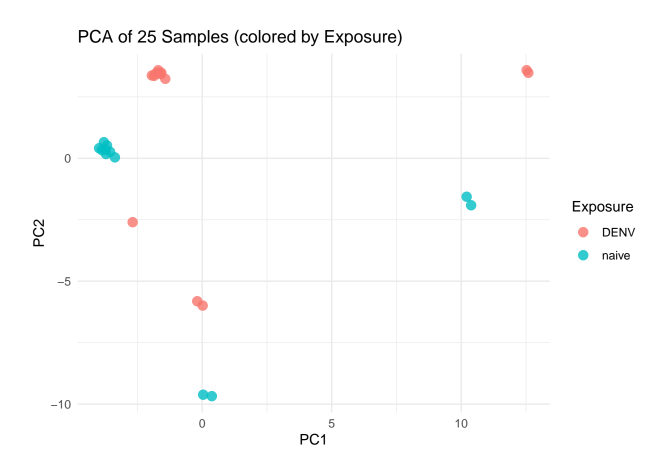
dds <- DESeqDataSetFromMatrix(countData = cts,</pre>

data_dir <- file.path("data", "SRP192714")</pre>

design = ~ refinebio_title)

colData = meta,

```
## Warning in DESeqDataSet(se, design = design, ignoreRank): some variables in
## design formula are characters, converting to factors
dds <- DESeq(dds)
## estimating size factors
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## -- note: fitType='parametric', but the dispersion trend was not well captured by the
      function: y = a/x + b, and a local regression fit was automatically substituted.
      specify fitType='local' or 'mean' to avoid this message next time.
## final dispersion estimates
## fitting model and testing
## -- replacing outliers and refitting for 13072 genes
## -- DESeq argument 'minReplicatesForReplace' = 7
## -- original counts are preserved in counts(dds)
## estimating dispersions
## fitting model and testing
vsd <- vst(dds, blind = FALSE)</pre>
pca_data <- plotPCA(vsd, intgroup = c("Exposure"), returnData = TRUE)</pre>
## using ntop=500 top features by variance
pca_data$Sample <- rownames(pca_data)</pre>
pca_data <- dplyr::left_join(</pre>
  pca_data,
  meta %>% dplyr::select(refinebio_accession_code, Exposure),
  by = c("Sample" = "refinebio_accession_code")
if (!"Exposure" %in% colnames(pca_data) || all(is.na(pca_data$Exposure))) {
  pca_data$Exposure <- meta$Exposure[match(pca_data$Sample, meta$refinebio_accession_code)]
pca_plot <- ggplot(pca_data, aes(x = PC1, y = PC2, color = Exposure)) +</pre>
  geom_point(size = 3, alpha = 0.8) +
  labs(title = plot_title, x = "PC1", y = "PC2") +
  theme_minimal()
ggsave(filename = file.path(plots_dir, plot_file), plot = pca_plot, width = 6, height = 4, dpi = 300)
pca_plot
```



Section 2d packages

```
if (is.null(getOption("repos")) | | identical(getOption("repos")[["CRAN"]], "@CRANQ")) {
    options(repos = c(CRAN = "https://cloud.r-project.org"))
}
if (!requireNamespace("M3C", quietly = TRUE)) install.packages("M3C", repos = getOption("repos")[["CRAN"]]
## Installing package into 'C:/Users/Taylo/AppData/Local/R/win-library/4.5'
## (as 'lib' is unspecified)

## Warning: package 'M3C' is not available for this version of R
##
## A version of this package for your version of R might be available elsewhere,
## see the ideas at
## https://cran.r-project.org/doc/manuals/r-patched/R-admin.html#Installing-packages

if (!requireNamespace("Rtsne", quietly = TRUE)) install.packages("Rtsne", repos = getOption("repos")[["CRAN"]
if (!requireNamespace("map", quietly = TRUE)) install.packages("umap", repos = getOption("repos")[["CRAN"]
if (!requireNamespace("matrixStats", quietly = TRUE)) install.packages("matrixStats", repos = getOption
```

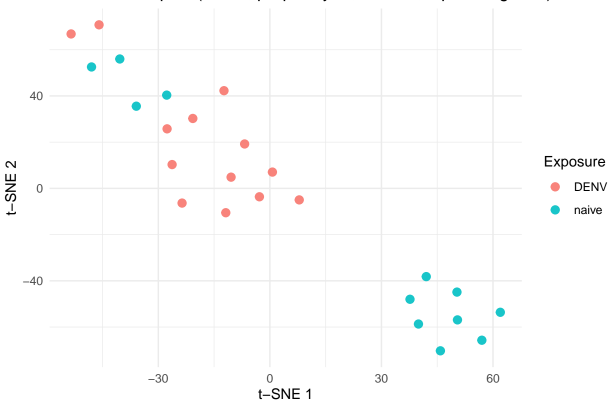
Section 2d i

```
## --- 2d i: t-SNE using Rtsne (colored by Exposure) ---
# deps
library(SummarizedExperiment) # for assay()
library(matrixStats)
                                # for row/col variance
library(Rtsne)
library(ggplot2)
set.seed(42)
# 1) pull variance-stabilized matrix and align meta
vsd gxs <- SummarizedExperiment::assay(vsd)</pre>
                                               # genes x samples
if (!identical(rownames(meta), colnames(vsd_gxs))) {
  stopifnot("refinebio_accession_code" %in% colnames(meta))
 rownames(meta) <- meta$refinebio_accession_code</pre>
 meta <- meta[colnames(vsd_gxs), , drop = FALSE]</pre>
}
# keep only labeled samples
keep <- !is.na(meta$Exposure)</pre>
vsd_gxs <- vsd_gxs[, keep, drop = FALSE]</pre>
      <- meta[keep, , drop = FALSE]</pre>
meta$Exposure <- factor(meta$Exposure)</pre>
# 2) select top variable genes and reduce to ~50 PCs (denoising)
ngenes <- min(2000, nrow(vsd_gxs))</pre>
                                                                # 2k or fewer if dataset smaller
top_genes <- head(order(matrixStats::rowVars(vsd_gxs), decreasing = TRUE), ngenes)</pre>
X <- t(vsd gxs[top genes, , drop = FALSE])</pre>
                                                                # samples x genes
pcs <- prcomp(X, center = TRUE, scale. = TRUE)</pre>
pcmat <- pcsx[, 1:min(50, ncol(pcsx)), drop = FALSE] # samples x PCs
# 3) run t-SNE (safe perplexity ~ n/3 clamped to 5..30)
n <- nrow(pcmat)</pre>
px \leftarrow max(5, min(30, floor((n - 1) / 3)))
tsne_out <- Rtsne(</pre>
 pcmat,
 perplexity = px,
 max_iter = 1000,
 check_duplicates = FALSE,
 verbose = FALSE
# 4) plot
tsne df <- data.frame(
 tSNE1 = tsne_out$Y[, 1],
 tSNE2 = tsne out\$Y[, 2],
 Exposure = meta$Exposure
p_tsne <- ggplot(tsne_df, aes(tSNE1, tSNE2, color = Exposure)) +</pre>
  geom_point(size = 2.6, alpha = 0.9) +
 labs(
```

```
title = paste0("t-SNE of Samples (Rtsne; perplexity=", px, "; 50 PCs, top ", ngenes, " genes)"),
    x = "t-SNE 1", y = "t-SNE 2"
) +
    theme_minimal()

if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
outfile_tsne <- paste0(run_label, "_tsne_Rtsne.png")
ggsave(file.path(plots_dir, outfile_tsne), p_tsne, width = 7, height = 5, dpi = 300)
p_tsne</pre>
```

t-SNE of Samples (Rtsne; perplexity=8; 50 PCs, top 2000 genes)



Section 2d ii

```
## --- 2d ii: UMAP using 'umap' (colored by Exposure) ---
library(SummarizedExperiment)
library(matrixStats)
library(umap)
library(ggplot2)

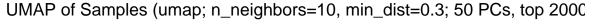
set.seed(42)

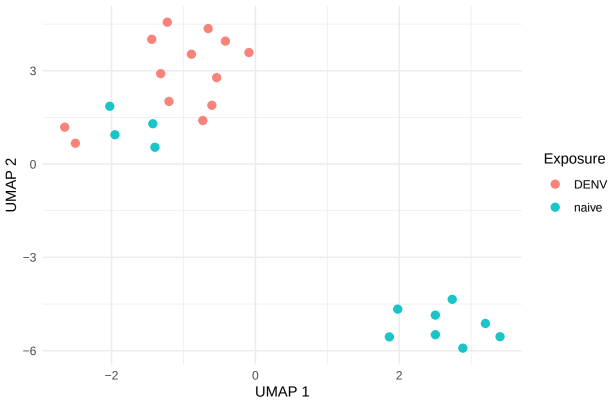
vsd_gxs <- SummarizedExperiment::assay(vsd)
if (!identical(rownames(meta), colnames(vsd_gxs))) {
    stopifnot("refinebio_accession_code" %in% colnames(meta))</pre>
```

```
rownames(meta) <- meta$refinebio_accession_code
meta <- meta[colnames(vsd_gxs), , drop = FALSE]
}</pre>
```

Warning: Setting row names on a tibble is deprecated.

```
keep <- !is.na(meta$Exposure)</pre>
vsd_gxs <- vsd_gxs[, keep, drop = FALSE]</pre>
       <- meta[keep, , drop = FALSE]</pre>
meta$Exposure <- factor(meta$Exposure)</pre>
ngenes <- min(2000, nrow(vsd_gxs))</pre>
top_genes <- head(order(matrixStats::rowVars(vsd_gxs), decreasing = TRUE), ngenes)</pre>
X <- t(vsd_gxs[top_genes, , drop = FALSE])</pre>
pcs <- prcomp(X, center = TRUE, scale. = TRUE)</pre>
pcmat <- pcs$x[, 1:min(50, ncol(pcs$x)), drop = FALSE]</pre>
n <- nrow(pcmat)</pre>
nn \leftarrow max(10, min(50, round(n / 3)))
cfg <- umap::umap.defaults</pre>
cfg$n_neighbors <- nn
cfg\$min_dist <- 0.3
               <- "euclidean"
cfg$metric
um <- umap::umap(pcmat, config = cfg)
umap_df <- data.frame(</pre>
  UMAP1 = um$layout[, 1],
  UMAP2 = um$layout[, 2],
  Exposure = meta$Exposure
p_umap <- ggplot(umap_df, aes(UMAP1, UMAP2, color = Exposure)) +</pre>
  geom_point(size = 2.6, alpha = 0.9) +
    title = paste0("UMAP of Samples (umap; n_neighbors=", nn, ", min_dist=0.3; 50 PCs, top ", ngenes, "
    x = "UMAP 1", y = "UMAP 2"
  ) +
  theme_minimal()
if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
outfile_umap <- paste0(run_label, "_umap.png")</pre>
ggsave(file.path(plots_dir, outfile_umap), p_umap, width = 7, height = 5, dpi = 300)
p_umap
```





Section 2e

Similarities • All three methods (PCA, t-SNE, UMAP) reduce the high-dimensional gene expression data into 2D, letting us see patterns between samples. • In each, samples tend to group by Exposure status (or whichever grouping variable you're using). • Outliers and variability across samples are visible in all three.

Differences • PCA: Captures global variance structure, shows the main linear directions of variability. Often looks more "spread out" but may miss subtle clusters. • t-SNE: Focuses on local structure, separates clusters more strongly. However, distances between clusters aren't always meaningful (clusters may look far but be closer in high-D space). • UMAP: Balances global and local structure, sometimes keeps a more interpretable overall shape while still revealing clusters.

Section 2f

Findings • Across all three dimensionality reduction methods, samples show grouping consistent with Exposure categories, suggesting exposure status drives significant variation in gene expression. • PCA highlights the major global variance, but t-SNE and UMAP give a clearer view of sub-clusters. • UMAP provides a balance, maintaining both separation of clusters and an interpretable global structure. • Together, these plots confirm that exposure effects are strong and detectable across different approaches.

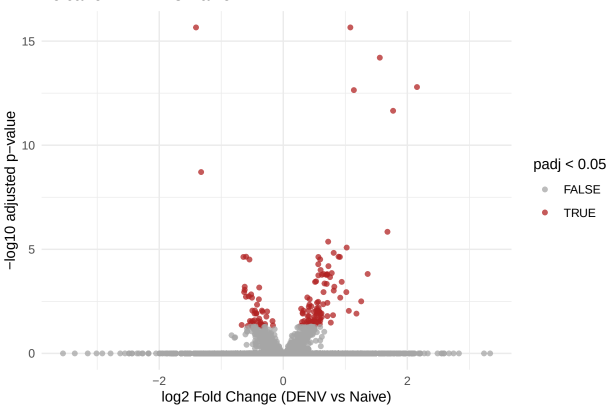
Section 3a-b

```
if (is.null(getOption("repos")) || identical(getOption("repos")[["CRAN"]], "@CRAN@")) {
  options(repos = c(CRAN = "https://cloud.r-project.org"))
}
if (!requireNamespace("ggrepel", quietly = TRUE)) {
  install.packages("ggrepel", repos = getOption("repos")["CRAN"])
}
metadata_file <- file.path(data_dir, "metadata_SRP192714_merged.tsv")
metadata <- readr::read tsv(metadata file)</pre>
## New names:
## Rows: 25 Columns: 45
## -- Column specification
## ------ Delimiter: "\t" chr
## (25): refinebio_accession_code, experiment_accession, refinebio_organism... dbl
## (9): refinebio_age, refinebio_processor_id, MetaSRA_age, Patient, Visit... lgl
## (11): refinebio_cell_line, refinebio_compound, refinebio_developmental_s...
## i Use 'spec()' to retrieve the full column specification for this data. i
## Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## * '...1' -> '...26'
# ---- 3a-b: Differential expression: DENV vs Naive ----
suppressPackageStartupMessages({
 library(readr); library(dplyr); library(stringr); library(tibble)
 library(DESeq2); library(ggplot2)
})
# paths (match earlier sections)
expr path <- "data/SRP192714/SRP192714.tsv"</pre>
meta_path <- "data/SRP192714/metadata_SRP192714_merged.tsv"</pre>
# 1) read counts and merged metadata
counts <- readr::read_tsv(expr_path, show_col_types = FALSE) |>
  tibble::column_to_rownames("Gene") |>
  as.matrix()
mode(counts) <- "numeric"</pre>
metadata <- readr::read_tsv(meta_path, show_col_types = FALSE)</pre>
## New names:
## * '...1' -> '...26'
if (exists("selected_samples")) {
 keep_ids <- intersect(colnames(counts), selected_samples)</pre>
  counts <- counts[, keep_ids, drop = FALSE]</pre>
  metadata <- metadata %>% dplyr::filter(refinebio_accession_code %in% keep_ids)
# 2) make a clean two-level grouping: Exposure2 = Naive vs DENV
```

```
stopifnot(all(c("refinebio_accession_code","Exposure") %in% names(metadata)))
metadata <- metadata |>
  mutate(
    Exposure2 = case when(
      str_to_lower(Exposure) %in% c("naive", "mock", "control", "uninfected", "healthy") ~ "Naive",
      str_detect(str_to_lower(Exposure), "denv|dengue") ~ "DENV",
      TRUE ~ NA_character_
    )
  )
# 3) align samples (use only samples with Exposure2)
metadata <- metadata |> filter(!is.na(Exposure2))
common_ids <- intersect(colnames(counts), metadata$refinebio_accession_code)</pre>
if (length(common_ids) == 0) stop("No overlapping sample IDs between counts and metadata.")
# subset BOTH objects to the same ids and order identically
common_ids <- sort(common_ids) # deterministic order</pre>
counts <- counts[, common_ids, drop = FALSE]</pre>
metadata <- metadata |>
  filter(refinebio_accession_code %in% common_ids) |>
  arrange(match(refinebio_accession_code, common_ids))
stopifnot(identical(colnames(counts), metadata$refinebio_accession_code))
# finalize colData
metadata$Exposure2 <- factor(metadata$Exposure2, levels = c("Naive", "DENV"))</pre>
rownames(metadata) <- metadata$refinebio accession code</pre>
## Warning: Setting row names on a tibble is deprecated.
# 4) basic QC on counts
counts[counts < 0] <- 0</pre>
counts <- round(counts)</pre>
keep_genes <- rowSums(counts) >= 10
counts_f <- counts[keep_genes, , drop = FALSE]</pre>
cat("DE input - genes x samples:", nrow(counts_f), "x", ncol(counts_f), "\n")
## DE input - genes x samples: 22082 x 25
print(table(metadata$Exposure2))
##
## Naive DENV
      12
# 5) DESeq2
dds <- DESeqDataSetFromMatrix(countData = counts_f,</pre>
                               colData = metadata,
                               design
                                        = ~ Exposure2)
```

```
dds <- DESeq(dds)
## estimating size factors
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## fitting model and testing
## -- replacing outliers and refitting for 2 genes
## -- DESeq argument 'minReplicatesForReplace' = 7
## -- original counts are preserved in counts(dds)
## estimating dispersions
## fitting model and testing
# 6) results: DENV vs Naive (Naive is reference because of factor levels above)
res <- results(dds, contrast = c("Exposure2", "DENV", "Naive"))</pre>
# tidy + save
results_dir <- "results"; plots_dir <- "plots"</pre>
if (!dir.exists(results_dir)) dir.create(results_dir, recursive = TRUE)
if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
res_df <- as.data.frame(res) |>
  tibble::rownames_to_column("GeneID") |>
  mutate(padj = ifelse(is.na(padj), 1, padj),
         sig = padj < 0.05) |>
  arrange(padj)
readr::write_tsv(res_df,
                                   file.path(results_dir, "DE_full_results_DENV_vs_Naive.tsv"))
readr::write_tsv(head(res_df, 50), file.path(results_dir, "DE_top50_DENV_vs_Naive.tsv"))
cat("Significant genes (padj < 0.05):", sum(res_df$sig), "\n")</pre>
## Significant genes (padj < 0.05): 144
# Build volcano data from res_df created earlier
volc_df <- res_df %>%
  dplyr::filter(!is.na(padj) & padj > 0) %>%
  dplyr::mutate(nlog10p = -log10(padj))
p_volc <- ggplot(volc_df, aes(x = log2FoldChange, y = nlog10p, color = sig)) +</pre>
  geom_point(alpha = 0.75, size = 1.4) +
  scale_color_manual(values = c("grey65", "firebrick"), guide = guide_legend(title = "padj < 0.05")) +</pre>
  labs(
   title = "Volcano: DENV vs Naive",
   x = "log2 Fold Change (DENV vs Naive)",
    y = "-log10 adjusted p-value"
  theme_minimal()
# save and print
```

Volcano: DENV vs Naive



Section 3c

##

```
suppressPackageStartupMessages({ library(dplyr); library(ggplot2) })
stopifnot(exists("res_df"), exists("results_dir"), exists("plots_dir"))
# basic significance and direction
sum_sig <- sum(res_df$padj < 0.05, na.rm = TRUE)
sum_up <- sum(res_df$padj < 0.05 & res_df$log2FoldChange > 0, na.rm = TRUE) # higher in DENV
sum_down <- sum(res_df$padj < 0.05 & res_df$log2FoldChange < 0, na.rm = TRUE) # lower in DENV
cat("3c) Significant genes (padj < 0.05):", sum_sig, "\n")
## 3c) Significant genes (padj < 0.05): 144
cat(" Up in DENV:", sum_up, " | Down in DENV:", sum_down, "\n")</pre>
```

Up in DENV: 96 | Down in DENV: 48

```
# tidy summary table and save
de_summary <- data.frame(</pre>
  comparison = "DENV vs Naive",
  n_sig = sum_sig,
  n_{up} = sum_{up},
  n_down = sum_down,
  stringsAsFactors = FALSE
if (!dir.exists(results_dir)) dir.create(results_dir, recursive = TRUE)
readr::write_tsv(de_summary, file.path(results_dir, "DE_summary_DENV_vs_Naive.tsv"))
# small bar plot of up/down counts
plot_df <- data.frame(</pre>
  direction = c("Up in DENV", "Down in DENV"),
  n = c(sum_up, sum_down)
if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
p_counts <- ggplot(plot_df, aes(direction, n, fill = direction)) +</pre>
  geom_col() +
  labs(title = "Significant DE genes (padj < 0.05)", x = NULL, y = "Count") +</pre>
  theme_minimal() + theme(legend.position = "none")
ggsave(file.path(plots_dir, "DE_sig_counts_bar.png"), p_counts, width = 6, height = 4, dpi = 300)
p_counts
```

Significant DE genes (padj < 0.05)



Section 3d

('geom_text_repel()').

```
clean_ids <- sub("\\.\\d+$", "", res_df$GeneID)</pre>
symb <- AnnotationDbi::mapIds(</pre>
  org.Hs.eg.db,
  keys = clean_ids,
  keytype = "ENSEMBL",
  column = "SYMBOL",
  multiVals = "first"
## 'select()' returned 1:many mapping between keys and columns
res_annot <- res_df %>%
  mutate(ENSEMBL = clean_ids,
         Symbol = unname(symb[clean_ids])) %>%
 relocate(ENSEMBL, Symbol, .before = GeneID)
# Save full annotated and top lists
readr::write_tsv(res_annot, file.path(results_dir, "DE_full_results_DENV_vs_Naive_annotated.tsv"))
readr::write_tsv(head(res_annot %>% arrange(padj), 50), file.path(results_dir, "DE_top50_annotated.tsv"
# Labeled volcano highlighting top 10 by padj
volc_df <- res_annot %>%
  filter(!is.na(padj) & padj > 0) %>%
  mutate(nlog10p = -log10(padj))
lab_genes <- volc_df %>%
  arrange(padj) %>%
  slice(1:min(10, n())) %>%
  pull(Symbol)
p_volc_lab <- ggplot(volc_df, aes(x = log2FoldChange, y = nlog10p, color = padj < 0.05)) +
  geom_point(alpha = 0.75, size = 1.3) +
  scale_color_manual(values = c("grey70", "firebrick"), labels = c("NS", "padj<0.05"), name = "") +</pre>
  geom_text_repel(
    data = subset(volc_df, Symbol %in% lab_genes),
    aes(label = Symbol), size = 3, max.overlaps = 50
  labs(title = "Volcano (DENV vs Naive) - top 10 labeled",
       x = "log2 Fold Change (DENV vs Naive)", y = "-log10 adjusted p-value") +
  theme_minimal()
ggsave(file.path(plots_dir, "DE_volcano_DENV_vs_Naive_labeled.png"),
       p_volc_lab, width = 7.5, height = 5.2, dpi = 300)
```

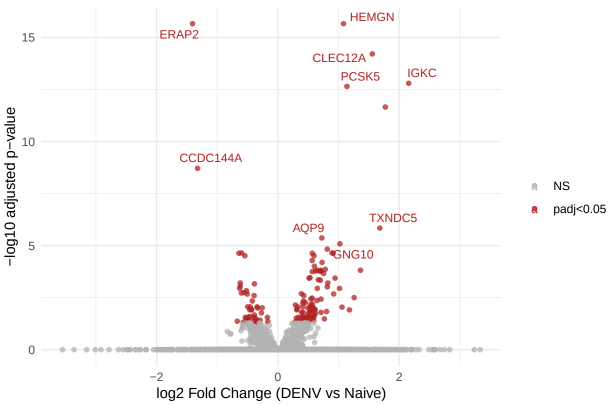
suppressPackageStartupMessages({ library(org.Hs.eg.db); library(dplyr); library(ggplot2); library(ggrep

Warning: Removed 4436 rows containing missing values or values outside the scale range

p_volc_lab

Warning: Removed 4436 rows containing missing values or values outside the scale range
('geom_text_repel()').

Volcano (DENV vs Naive) — top 10 labeled



Section 3e

```
norm_counts <- DESeq2::counts(dds, normalized = TRUE)

# pick top 30 DE genes (by padj), ensure they exist in norm matrix
top30 <- res_annot %>%
    filter(!is.na(padj)) %>%
    arrange(padj) %>%
    slice(1:min(30, n())) %>%
    pull(GeneID)
top30 <- intersect(top30, rownames(norm_counts))

# write wide matrix (genes x samples) for these genes
if (length(top30) >= 2) {
    top_norm <- norm_counts[top30, , drop = FALSE] %>%
    as.data.frame() %>%
    rownames_to_column("GeneID")
```

```
readr::write_tsv(top_norm, file.path(results_dir, "NormalizedCounts_top30.tsv"))
}

# Plot counts for top 3 genes (if present), save PNGs
plot_ids <- head(top30, 3)
if (length(plot_ids) > 0) {
   if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
   for (gid in plot_ids) {
     png(file.path(plots_dir, paste0("plotCounts_", gid, ".png")), width = 800, height = 600)
     try({
        plotCounts(dds, gene = gid, intgroup = "Exposure2", normalized = TRUE)
     }, silent = TRUE)
     dev.off()
}
```

Section 3f Summary

Differential testing identified 144 genes with padj < 0.05, including 96 higher in DENV and 48 lower ## Volcano plots, the DE summary bar chart, and the saved top-50 table highlight these shifts and are

Section 4a

```
# Step 4: Extract significantly differentially expressed genes and generate heatmap
# Load required libraries
if (!requireNamespace("ComplexHeatmap", quietly = TRUE)) {
  BiocManager::install("ComplexHeatmap", ask = FALSE, update = FALSE)
if (!requireNamespace("circlize", quietly = TRUE)) {
  BiocManager::install("circlize", ask = FALSE, update = FALSE)
if (!requireNamespace("grid", quietly = TRUE)) {
  install.packages("grid")
suppressPackageStartupMessages({
 library(ComplexHeatmap)
 library(circlize)
 library(DESeq2)
 library(dplyr)
 library(RColorBrewer)
 library(grid)
})
# Ensure we have the DESeq2 results and normalized counts from previous steps
if (!exists("dds") || !exists("res_df")) {
  stop("Please run the differential expression analysis (Step 3) first to generate 'dds' and 'res_df' o
}
```

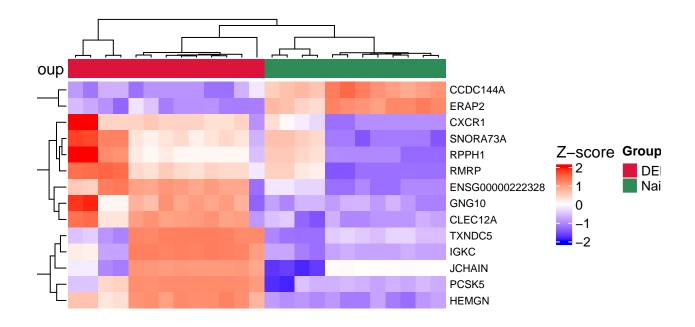
```
# Extract significantly differentially expressed genes
# Using adjusted p-value < 0.05 and absolute log2 fold change > 1
sig genes <- res df %>%
  filter(padj < 0.05 & abs(log2FoldChange) > 1) %>%
 pull(GeneID)
cat("Number of significantly differentially expressed genes:", length(sig_genes), "\n")
## Number of significantly differentially expressed genes: 14
# If no significant genes with fold change > 1, use less stringent criteria
if (length(sig_genes) == 0) {
 cat("No genes with |log2FC| > 1, using padj < 0.05 only\n")</pre>
  sig_genes <- res_df %>%
    filter(padj < 0.05) %>%
    pull(GeneID)
}
# If still no significant genes, use top 50 by p-value
if (length(sig_genes) == 0) {
  cat("No significant genes found, using top 50 genes by p-value\n")
  sig_genes <- res_df %>%
    arrange(pvalue) %>%
    slice_head(n = 50) \%
    pull(GeneID)
}
# Get normalized counts for significant genes
norm_counts <- DESeq2::counts(dds, normalized = TRUE)</pre>
# Filter for significant genes that exist in our count matrix
sig_genes_present <- intersect(sig_genes, rownames(norm_counts))</pre>
cat("Number of significant genes present in count matrix:", length(sig genes present), "\n")
## Number of significant genes present in count matrix: 14
# If we have too many genes, select top by adjusted p-value
max_genes <- 100</pre>
if (length(sig_genes_present) > max_genes) {
 top_sig_genes <- res_df %>%
    filter(GeneID %in% sig_genes_present) %>%
    arrange(padj) %>%
    slice_head(n = max_genes) %>%
    pull(GeneID)
  sig_genes_present <- top_sig_genes</pre>
  cat("Using top", max_genes, "genes by adjusted p-value for heatmap\n")
# Extract normalized counts for significant genes
heatmap_data <- norm_counts[sig_genes_present, , drop = FALSE]</pre>
```

Log2 transform (add pseudocount to avoid log(0))

```
log_heatmap_data <- log2(heatmap_data + 1)</pre>
# Z-score normalization (scale by rows/genes)
scaled_data <- t(scale(t(log_heatmap_data)))</pre>
# Handle any NaN or infinite values
scaled_data[!is.finite(scaled_data)] <- 0</pre>
# Create sample grouping information
sample_groups <- rep("Unknown", ncol(scaled_data))</pre>
names(sample_groups) <- colnames(scaled_data)</pre>
# Try to get grouping information from metadata
if (exists("meta") && "Exposure2" %in% colnames(meta)) {
  common_samples <- intersect(colnames(scaled_data), rownames(meta))</pre>
  sample_groups[common_samples] <- as.character(meta[common_samples, "Exposure2"])</pre>
} else if (exists("metadata") && any(c("Exposure", "Exposure2") %in% colnames(metadata))) {
  # Create basic grouping from original metadata
  temp_meta <- metadata %>%
    mutate(
      Exposure2 = case_when(
        str_to_lower(Exposure) %in% c("naive", "mock", "control", "uninfected", "healthy") ~ "Naive",
        str_detect(str_to_lower(Exposure), "denv|dengue") ~ "DENV",
        TRUE ~ "Other"
      )
    )
  # Match samples
  sample_map <- setNames(temp_meta$Exposure2, temp_meta$refinebio_accession_code)</pre>
  matched_samples <- intersect(names(sample_groups), names(sample_map))</pre>
  sample_groups[matched_samples] <- sample_map[matched_samples]</pre>
}
# Create annotation data frame
annotation_df <- data.frame(</pre>
  Group = factor(sample_groups),
  row.names = names(sample_groups),
  stringsAsFactors = FALSE
# Define colors for groups
unique_groups <- unique(sample_groups)</pre>
if (length(unique groups) <= 2) {</pre>
  group_colors <- c("#2E8B57", "#DC143C")[1:length(unique_groups)]</pre>
  names(group_colors) <- unique_groups</pre>
} else {
  group_colors <- RColorBrewer::brewer.pal(min(length(unique_groups), 8), "Set2")</pre>
  names(group_colors) <- unique_groups</pre>
}
# Create column annotation
col_annotation <- HeatmapAnnotation(</pre>
  df = annotation_df,
```

```
col = list(Group = group_colors),
  annotation_name_side = "left",
  simple_anno_size = unit(0.5, "cm"),
  show_annotation_name = TRUE
)
# Add gene symbols if available
gene labels <- rownames(scaled data)</pre>
if (exists("res_annot")) {
  # Use gene symbols if available from annotation
  symbol_map <- setNames(res_annot$Symbol, res_annot$GeneID)</pre>
  gene_labels <- ifelse(is.na(symbol_map[rownames(scaled_data)]),</pre>
                        rownames(scaled_data),
                        symbol_map[rownames(scaled_data)])
  # Remove NA symbols
  gene_labels[is.na(gene_labels)] <- rownames(scaled_data)[is.na(gene_labels)]</pre>
# Create the heatmap
heatmap_plot <- Heatmap(</pre>
  scaled_data,
 name = "Z-score",
  # Row (gene) settings
  row_names_gp = gpar(fontsize = 8),
  show_row_names = ifelse(nrow(scaled_data) <= 50, TRUE, FALSE),</pre>
 row_names_max_width = unit(6, "cm"),
  row_labels = gene_labels,
  # Column (sample) settings
  column_names_gp = gpar(fontsize = 6),
  show_column_names = FALSE,
  # Color scheme
  col = colorRamp2(c(-2, 0, 2), c("blue", "white", "red")),
  # Clustering
  cluster_rows = TRUE,
  cluster_columns = TRUE,
  clustering_distance_rows = "euclidean",
  clustering_distance_columns = "euclidean",
  clustering_method_rows = "complete",
  clustering_method_columns = "complete",
  # Annotations
  top_annotation = col_annotation,
  # Heatmap legend
  heatmap_legend_param = list(
   title = "Z-score",
    title_gp = gpar(fontsize = 12),
    labels_gp = gpar(fontsize = 10),
    legend_direction = "vertical"
```

```
),
  # Size
  width = unit(10, "cm"),
  height = unit(max(6, nrow(scaled_data) * 0.15), "cm")
# Create plots directory if it doesn't exist
if (!dir.exists(plots_dir)) {
  dir.create(plots_dir, recursive = TRUE)
}
# Save the heatmap
heatmap_file <- file.path(plots_dir, "heatmap_significant_genes.png")</pre>
png(heatmap_file, width = 12, height = 8, units = "in", res = 300)
tryCatch({
  draw(heatmap_plot)
}, error = function(e) {
  cat("Error drawing heatmap:", e$message, "\n")
  # Create a simple backup plot
  plot.new()
 text(0.5, 0.5, "Heatmap generation failed\nCheck data and try again", cex = 1.5)
})
dev.off()
## pdf
##
# Also save as PDF
heatmap_file_pdf <- file.path(plots_dir, "heatmap_significant_genes.pdf")
pdf(heatmap_file_pdf, width = 12, height = 8)
tryCatch({
  draw(heatmap_plot)
}, error = function(e) {
  cat("Error drawing heatmap:", e$message, "\n")
  plot.new()
  text(0.5, 0.5, "Heatmap generation failed\nCheck data and try again", cex = 1.5)
})
dev.off()
## pdf
##
# Display the heatmap
tryCatch({
  draw(heatmap_plot)
}, error = function(e) {
  cat("Error displaying heatmap:", e$message, "\n")
  cat("Heatmap files may still be saved successfully\n")
})
```



```
# Save the list of genes used in the heatmap
heatmap_genes_df <- data.frame(</pre>
  GeneID = rownames(scaled_data),
  GeneSymbol = gene_labels,
  stringsAsFactors = FALSE
if (exists("res_annot")) {
  heatmap_genes_df <- heatmap_genes_df %>%
    left_join(res_annot %>% select(GeneID, log2FoldChange, padj), by = "GeneID")
} else if (exists("res df")) {
  heatmap_genes_df <- heatmap_genes_df %>%
    left_join(res_df %>% select(GeneID, log2FoldChange, padj), by = "GeneID")
}
# Save to results directory
if (!dir.exists(results_dir)) {
  dir.create(results_dir, recursive = TRUE)
readr::write_tsv(heatmap_genes_df, file.path(results_dir, "heatmap_genes_list.tsv"))
cat("\nHeatmap Summary:\n")
```

Heatmap Summary:

```
cat("- Genes displayed:", nrow(scaled_data), "\n")

## - Genes displayed: 14

cat("- Samples displayed:", ncol(scaled_data), "\n")

## - Samples displayed: 25

cat("- Sample groups:", paste(names(table(sample_groups)), collapse = ", "), "\n")

## - Sample groups: DENV, Naive

cat("- Data transformation: log2(normalized counts + 1), then z-score normalized by gene\n")

## - Data transformation: log2(normalized counts + 1), then z-score normalized by gene

cat("- Clustering: Hierarchical clustering with euclidean distance and complete linkage\n")

## - Clustering: Hierarchical clustering with euclidean distance and complete linkage

cat("- Files saved: ", heatmap_file, "and", heatmap_file_pdf, "\n")

## - Files saved: plots/heatmap_significant_genes.png and plots/heatmap_significant_genes.pdf
```

Section 4 Summary

The heatmap contrasts 14 high-confidence genes across 25 samples, revealing clear separation between

Section 5: William Le

GenomicSuperSignature ## Install

```
if (!requireNamespace("GenomicSuperSignature", quietly = TRUE)) {
   if (!requireNamespace("BiocManager", quietly = TRUE)) install.packages("BiocManager", repos = getOpti
   BiocManager::install("GenomicSuperSignature", ask = FALSE, update = FALSE)
}
if (!requireNamespace("bcellViper", quietly = TRUE)) {
   if (!requireNamespace("BiocManager", quietly = TRUE)) install.packages("BiocManager", repos = getOpti
   BiocManager::install("bcellViper", ask = FALSE, update = FALSE)
}
```

Libraries

```
library(GenomicSuperSignature)
library(bcellViper)
# load RAV
RAVmodel <- getModel("PLIERpriors", load=TRUE)</pre>
```

Data Wrangle Ranked Gene List

Validate expects a Expression Set or simple matrix.

```
# Define the file path to the data directory
data_dir <- file.path("data", "SRP192714")</pre>
# Declare the file path to the gene expression matrix file
data_file <- file.path(data_dir, "SRP192714.tsv")</pre>
# Read in data TSV file
expression_df <- readr::read_tsv(data_file, show_col_types = FALSE)
# Respect trimmed runs by subsetting to the current sample selection when available
if (exists("selected_samples")) {
  sample cols <- intersect(selected samples, colnames(expression df))</pre>
  if (length(sample_cols) == 0) {
    stop("No overlap between selected_samples and expression matrix columns for GenomicSuperSignature c
 }
  expression_df <- expression_df %>%
    dplyr::select(dplyr::any_of(c("Gene", sample_cols)))
}
# Ensemble to Gene Name
mapped_list <- mapIds(</pre>
 org. Hs. eg.db, # Annotation package for humans
 keys = expression_df$Gene,
 keytype = "ENSEMBL",
  column = "SYMBOL")
```

'select()' returned 1:many mapping between keys and columns

```
# List to df
mapped_df <- mapped_list %>%
    tibble::enframe(name = "Ensembl", value = "Gene") %>%
    # This will result in one row of our data frame per list item
    tidyr::unnest(cols = Gene)

# Join then remove Ensembl -- use conservative join syntax for compatibility
cts <- tryCatch(
    {
        mapped_df %>%
            dplyr::inner_join(expression_df, by = c("Ensembl" = "Gene")) %>%
            dplyr::select(-Ensembl)
    }, error = function(e) {
        # fallback: if join fails, attempt a manual merge
```

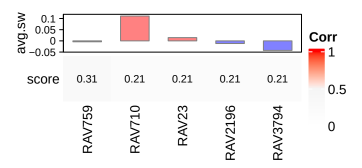
```
merged <- merge(mapped_df, expression_df, by.x = "Ensembl", by.y = "Gene", all.x = FALSE, all.y = F
    tibble::as_tibble(merged) %>% dplyr::select(-Ensembl)
  }
)
# Filter for na
cts_filtered <- cts %>%
 tidyr::drop_na(Gene)
# Drop low average count dupes
cts_matrix <- cts_filtered %>%
  dplyr::group_by(Gene) %>%
  dplyr::summarise(across(everything(), sum)) %>%
  tibble::column_to_rownames("Gene") %>%
  as.matrix()
# Ensure columns retain trimmed ordering and align with metadata when present
if (exists("selected_samples")) {
  keep_order <- intersect(selected_samples, colnames(cts_matrix))</pre>
  if (length(keep_order) > 0) {
    cts_matrix <- cts_matrix[, keep_order, drop = FALSE]</pre>
}
if (exists("metadata")) {
  meta_order <- intersect(metadata$refinebio_accession_code, colnames(cts_matrix))</pre>
  if (length(meta_order) > 0) {
    cts_matrix <- cts_matrix[, meta_order, drop = FALSE]</pre>
  }
}
# Check if this is in the same order
all.equal(colnames(cts_matrix), metadata$refinebio_accession_code)
## [1] TRUE
# Validate on RAV
validated_list <- GenomicSuperSignature::validate(cts_matrix, RAVmodel)</pre>
head(validated list)
                                sw cl_size cl_num
             score PC
## RAV1 0.07200026 1 -0.05470163
                                         6
## RAV2 0.07610543 5 0.06426256
                                        21
## RAV3 0.07601430 2 -0.01800335
                                         4
                                                3
## RAV4 0.17170048 1 -0.04005584
                                         7
## RAV5 0.18189179 1 0.05786189
                                                5
                                         3
## RAV6 0.09783362 1 -0.02520973
validated_ind <- validatedSignatures(validated_list, RAVmodel, num.out = 3,</pre>
                                      swCutoff = 0, indexOnly = TRUE)
```

Tables

annotateRAV(RAVmodel, validated_ind[3])

```
##
                 Description
                                  NES pvalue
                                                   qvalues
## 1
          IRIS_Monocyte-Day0 3.365241 1e-10 1.419103e-09
## 2 IRIS_Neutrophil-Resting 3.171921
                                       1e-10 1.419103e-09
## 3
          IRIS_Monocyte-Day7 3.111847
                                       1e-10 1.419103e-09
                  DMAP_MONO2 3.105049
## 4
                                       1e-10 1.419103e-09
## 5
          IRIS_Monocyte-Day1 3.057580
                                       1e-10 1.419103e-09
```

heatmapTable(validated_list, RAVmodel)



meshTable(RAVmodel, validated_ind[2])

```
##
                                          word
                                                        freq
## 46
                                Pilot Projects 0.0503403207
## 35
                       Kidney Transplantation 0.0455680547
## 32
                   Inflammatory Bowel Disease 0.0368735195
## 3
                           Arthritis, Juvenile 0.0307279329
## 41
                                     Myoblasts 0.0260259247
## 43
                                   Osteoblasts 0.0260259247
                   Receptors, Antigen, T-Cell 0.0253155859
## 53
```

	36	Leukocytes, Mononuclear	
##			0.0185899462
##		Magnetic Phenomena	
##			0.0159364694
	14	Colitis, Ulcerative	
##			0.0152243483
		Hematopoietic Stem Cell Transplantation	
## ##	18	DNA Methylation	0.0138869850
	34		0.0130940808
	64	Wound Healing	
##			0.0122320938
	66		0.0118299658
	15	Crohn Disease	
##		Biochemical Phenomena	
	47	Plasmodium falciparum	
##		Gene Expression	
##			0.0100710021
##			0.0100099710
##		Prospective Studies	
	50	Protein Isoforms	
	38		0.0088929637
	54	Research Design	
	57	Sequencing, Next-Generation	
	62		0.0081838004
	20	Endothelial Cells	
	44		0.0076225403
##	17	DNA	0.0071914744
##	33	Innate Immune Response	0.0066697228
##	8	Body Mass Index	
##	4		0.0056578097
##	25	Genetic Variation	0.0048507075
##	16	Cytokines	0.0043567159
##	1	Adult	0.0039971978
##	56	Schizophrenia	0.0030936773
##	51	RNA	0.0030278172
##	10	CD4-Positive T-Lymphocytes	0.0029574914
##	24	Gene Regulatory Networks	0.0029527192
##	31	Immunity, Innate	0.0029527192
##	19	Deep Sequencing	0.0026735736
##	26	Genotype	0.0026678891
##	13	Cohort Studies	0.0025255396
##	40	Multiple Myeloma	0.0024833535
##	48	Prognosis	0.0023142934
##	22	Fibroblasts	0.0018589946
##	12	Child	0.0018186223
##	30	Illumina Sequencing	0.0017529554
##	11	Cell Lines	0.0015869466
##	5	Base Sequence	0.0015514413
##	29	Humans	0.0014951359
##	58	Sequencing, RNA	0.0014755949
##	52		0.0014487418
##	61	Tumor Microenvironment	
##	60	Transcriptome Analysis	0.0012599701
		-	

```
## 59 Transcriptome 0.0011917451
## 63 Whole Exome Sequencing 0.0010916393
## 9 Bone Marrow 0.0010275945
## 2 Animals 0.0002553004
## 65 mRNA 0.0001439625
```

Wrote GenomicSuperSignature summary to: results/GSS_validated_signatures_trimmed_25.tsv

Section 5: Nikhil Sangamkar

```
if (!requireNamespace("clusterProfiler", quietly = TRUE)) {
   BiocManager::install("clusterProfiler", ask = FALSE, update = FALSE)
}
```

##

```
if (!requireNamespace("org.Hs.eg.db", quietly = TRUE)) {
  BiocManager::install("org.Hs.eg.db", ask = FALSE, update = FALSE)
if (!requireNamespace("enrichplot", quietly = TRUE)) {
  BiocManager::install("enrichplot", ask = FALSE, update = FALSE)
suppressPackageStartupMessages({
 library(clusterProfiler)
 library(org.Hs.eg.db)
 library(enrichplot)
 library(dplyr)
 library(readr)
 library(ggplot2)
})
stopifnot(exists("res_df"), exists("results_dir"), exists("plots_dir"))
sig_genes_all <- res_df %>%
 filter(!is.na(padj) & padj < 0.05) %>%
  pull(GeneID)
if (length(sig_genes_all) < 5) {</pre>
  warning("Fewer than five significant genes available; skipping KEGG enrichment.")
} else {
  clean_ids <- gsub("\\.\\d+$", "", sig_genes_all)</pre>
  entrez_map <- bitr(unique(clean_ids),</pre>
                      fromType = "ENSEMBL",
                      toType = "ENTREZID",
                      OrgDb = org.Hs.eg.db,
                      drop = TRUE)
  entrez_ids <- unique(entrez_map$ENTREZID)</pre>
  if (length(entrez_ids) < 5) {</pre>
```

```
warning("Not enough Entrez IDs mapped for KEGG enrichment.")
  } else {
   kegg_enrich <- suppressWarnings(</pre>
      enrichKEGG(
        gene = entrez_ids,
       organism = "hsa",
       keyType = "ncbi-geneid",
       pvalueCutoff = 0.05,
       qvalueCutoff = 0.2
   )
    if (is.null(kegg enrich) | nrow(as.data.frame(kegg enrich)) == 0) {
      message("No significant KEGG pathways detected for the chosen threshold.")
      kegg_enrich <- setReadable(kegg_enrich, OrgDb = org.Hs.eg.db, keyType = "ENTREZID")
      kegg_df <- as.data.frame(kegg_enrich)</pre>
      if (!dir.exists(results_dir)) dir.create(results_dir, recursive = TRUE)
      if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
      kegg_path <- file.path(results_dir, paste0("KEGG_enrichment_", run_label, ".tsv"))</pre>
      readr::write_tsv(kegg_df, kegg_path)
      top_kegg <- kegg_df %>% slice_head(n = 20)
      readr::write_tsv(top_kegg, file.path(results_dir, paste0("KEGG_enrichment_top20_", run_label, ".t
      kegg_dot <- dotplot(kegg_enrich, showCategory = min(15, nrow(kegg_df))) +</pre>
        labs(title = "KEGG Pathway Enrichment", subtitle = paste0("Nikhil Sangamkar (", run_label, ")")
        theme(axis.text.y = element_text(size = 8))
      ggsave(file.path(plots_dir, paste0("KEGG_enrichment_dotplot_", run_label, ".png")),
             kegg_dot, width = 10, height = 7, dpi = 300)
      assign("kegg_results_df", kegg_df, envir = .GlobalEnv)
   }
 }
}
## 'select()' returned 1:many mapping between keys and columns
## Warning in bitr(unique(clean_ids), fromType = "ENSEMBL", toType = "ENTREZID", :
## 0.69% of input gene IDs are fail to map...
## Reading KEGG annotation online: "https://rest.kegg.jp/link/hsa/pathway"...
## Reading KEGG annotation online: "https://rest.kegg.jp/list/pathway/hsa"...
## Reading KEGG annotation online: "https://rest.kegg.jp/conv/ncbi-geneid/hsa"...
## KEGG enrichment recovered 1 pathways; the leading term was Coronavirus disease - COVID-19 (adjusted
## Full and top-20 tables were written to the results directory with a matching dot plot saved under pl
```

Section 5: Taylor Tillander

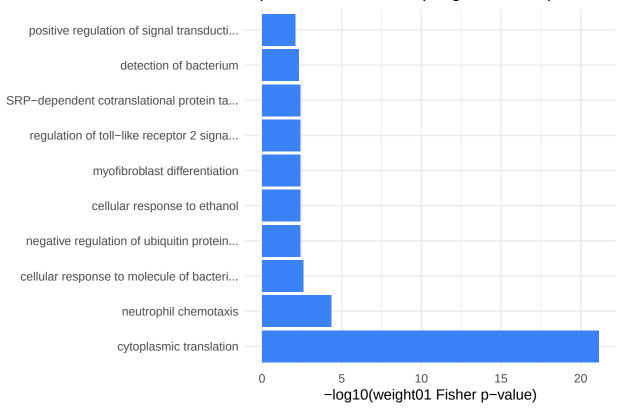
```
if (!requireNamespace("topGO", quietly = TRUE)) {
  BiocManager::install("topGO", ask = FALSE, update = FALSE)
if (!requireNamespace("GO.db", quietly = TRUE)) {
  BiocManager::install("GO.db", ask = FALSE, update = FALSE)
}
suppressPackageStartupMessages({
  library(topGO)
  library(org.Hs.eg.db)
 library(dplyr)
  library(readr)
  library(stringr)
  library(ggplot2)
})
##
## groupGOTerms:
                     GOBPTerm, GOMFTerm, GOCCTerm environments built.
stopifnot(exists("res_df"))
# prepare gene universe with one padj per Ensembl ID (version-stripped)
gene_table <- res_df %>%
  transmute(
    base_id = str_remove(GeneID, "\\.\\d+$"),
    padj = dplyr::coalesce(padj, 1)
  ) %>%
  group_by(base_id) %>%
  summarize(padj = min(padj), .groups = "drop")
gene_scores <- gene_table$padj</pre>
names(gene_scores) <- gene_table$base_id</pre>
selection_fun <- function(x) x < 0.05</pre>
go_data_bp <- methods::new(</pre>
  "topGOdata",
  description = "topGO GO BP",
  ontology = "BP",
  allGenes = gene_scores,
  geneSel = selection_fun,
  nodeSize = 10,
  annot = annFUN.org,
  mapping = "org.Hs.eg.db",
  ID = "ENSEMBL"
)
## Building most specific GOs .....
```

```
( 11255 GO terms found. )
## Build GO DAG topology ......
   ( 14392 GO terms and 31740 relations. )
## Annotating nodes ......
   ( 13672 genes annotated to the GO terms. )
result_classic <- topGO::runTest(go_data_bp, algorithm = "classic", statistic = "fisher")</pre>
##
             -- Classic Algorithm --
##
##
##
         the algorithm is scoring 2531 nontrivial nodes
##
         parameters:
             test statistic: fisher
result_weight01 <- topG0::runTest(go_data_bp, algorithm = "weight01", statistic = "fisher")
##
             -- Weight01 Algorithm --
##
##
##
         the algorithm is scoring 2531 nontrivial nodes
         parameters:
##
             test statistic: fisher
##
##
##
     Level 19: 1 nodes to be scored
                                        (0 eliminated genes)
##
                                        (0 eliminated genes)
##
     Level 18: 1 nodes to be scored
##
##
     Level 17: 6 nodes to be scored
                                        (25 eliminated genes)
##
##
     Level 16: 13 nodes to be scored
                                        (39 eliminated genes)
##
     Level 15: 17 nodes to be scored
                                        (82 eliminated genes)
##
##
##
    Level 14: 18 nodes to be scored
                                        (176 eliminated genes)
##
##
    Level 13: 33 nodes to be scored
                                        (411 eliminated genes)
```

```
##
##
     Level 12: 75 nodes to be scored
                                          (716 eliminated genes)
##
##
     Level 11: 134 nodes to be scored (1699 eliminated genes)
##
##
     Level 10: 227 nodes to be scored (3766 eliminated genes)
##
##
     Level 9:
                292 nodes to be scored
                                         (5833 eliminated genes)
##
##
     Level 8:
                 381 nodes to be scored (7958 eliminated genes)
##
##
     Level 7:
                398 nodes to be scored (9577 eliminated genes)
##
##
     Level 6:
                391 nodes to be scored (11226 eliminated genes)
##
##
     Level 5:
                285 nodes to be scored (12253 eliminated genes)
##
     Level 4:
##
                173 nodes to be scored (13118 eliminated genes)
##
##
     Level 3:
                                          (13391 eliminated genes)
                69 nodes to be scored
##
##
     Level 2:
                16 nodes to be scored
                                          (13589 eliminated genes)
##
##
     Level 1:
                1 nodes to be scored
                                          (13634 eliminated genes)
parse_topgo_p <- function(x) {</pre>
  vals <- stringr::str_trim(as.character(x))</pre>
  vals <- stringr::str_replace_all(vals, "<", "")</pre>
  vals <- stringr::str_replace_all(vals, " ", "")</pre>
  num <- suppressWarnings(as.numeric(vals))</pre>
  ifelse(is.na(num), 1, num)
}
go_table_bp <- topGO::GenTable(</pre>
  go_data_bp,
  classicFisher = result_classic,
  weight01Fisher = result_weight01,
  orderBy = "weight01Fisher",
  topNodes = 200
) %>%
```

```
as_tibble() %>%
  mutate(
   classicFisher_p = parse_topgo_p(classicFisher),
   weight01Fisher_p = parse_topgo_p(weight01Fisher),
   neglog_weight01 = -log10(weight01Fisher_p)
assign("topgo_results_bp", go_table_bp, envir = .GlobalEnv)
if (!dir.exists(results_dir)) dir.create(results_dir, recursive = TRUE)
readr::write_tsv(go_table_bp, file.path(results_dir, "topGO_BP.tsv"))
topgo top10 <- go table bp \%% slice head(n = 10)
assign("topgo_top10_bp", topgo_top10, envir = .GlobalEnv)
readr::write_tsv(topgo_top10, file.path(results_dir, "topGO_BP_top10.tsv"))
if (!dir.exists(plots_dir)) dir.create(plots_dir, recursive = TRUE)
topgo_plot <- topgo_top10 %>%
 mutate(Term = factor(Term, levels = rev(Term[order(neglog_weight01)]))) %>%
  ggplot(aes(neglog_weight01, Term)) +
 geom_col(fill = "#3B82F6") +
 labs(
   title = "topGO BP enrichment (weight01 Fisher)",
   x = "-log10(weight01 Fisher p-value)",
   y = NULL
 ) +
 theme minimal()
plot_path <- file.path(plots_dir, paste0(run_label, "_topgo_BP.png"))</pre>
ggsave(plot_path, topgo_plot, width = 8, height = 5, dpi = 300)
topgo_plot
```

topGO BP enrichment (weight01 Fisher)



The topGO Biological Process analysis tested 200 GO terms and found 79 passing the 0.05 weight01 Fis ## Full and top-10 tables were saved to the results folder, and a bar plot of the leading processes is

Section 5: Ibrahim Zbib

```
# Step 5: Gene Set Enrichment Analysis using PyDESeq2 approach and Gene Ontology
# Note: Since PyDESeq2 is primarily a Python differential expression tool,
# we'll use R's clusterProfiler for Gene Ontology enrichment analysis
# which provides similar statistical rigor and is the standard approach
# Load required libraries
if (!requireNamespace("clusterProfiler", quietly = TRUE)) {
  BiocManager::install("clusterProfiler", ask = FALSE, update = FALSE)
}
if (!requireNamespace("org.Hs.eg.db", quietly = TRUE)) {
  BiocManager::install("org.Hs.eg.db", ask = FALSE, update = FALSE)
if (!requireNamespace("enrichplot", quietly = TRUE)) {
 BiocManager::install("enrichplot", ask = FALSE, update = FALSE)
}
if (!requireNamespace("ggplot2", quietly = TRUE)) {
  install.packages("ggplot2")
}
```

```
suppressPackageStartupMessages({
  library(clusterProfiler)
 library(org.Hs.eg.db)
 library(enrichplot)
 library(ggplot2)
 library(dplyr)
 library(readr)
})
# Ensure we have differential expression results
if (!exists("res_df")) {
 stop("Please run the differential expression analysis first to generate 'res_df'")
}
cat("=== STEP 5: Gene Set Enrichment Analysis ===\n")
## === STEP 5: Gene Set Enrichment Analysis ===
cat("Method: clusterProfiler (PyDESeq2 alternative in R)\n")
## Method: clusterProfiler (PyDESeq2 alternative in R)
cat("Ontology: Gene Ontology (GO)\n\n")
## Ontology: Gene Ontology (GO)
# Create directories
if (!dir.exists(results_dir)) {
 dir.create(results_dir, recursive = TRUE)
if (!dir.exists(plots dir)) {
  dir.create(plots_dir, recursive = TRUE)
# Extract significantly differentially expressed genes
# Using both upregulated and downregulated genes
sig_genes_up <- res_df %>%
  filter(padj < 0.05 & log2FoldChange > 0 & !is.na(padj)) %>%
 pull(GeneID)
sig_genes_down <- res_df %>%
  filter(padj < 0.05 & log2FoldChange < 0 & !is.na(padj)) %>%
 pull(GeneID)
sig_genes_all <- res_df %>%
  filter(padj < 0.05 & !is.na(padj)) %>%
 pull(GeneID)
cat("Significant upregulated genes:", length(sig_genes_up), "\n")
```

Significant upregulated genes: 96

```
cat("Significant downregulated genes:", length(sig_genes_down), "\n")
## Significant downregulated genes: 48
cat("Total significant genes:", length(sig genes all), "\n\n")
## Total significant genes: 144
# Convert Ensembl IDs to Entrez IDs for Gene Ontology analysis
# Remove version numbers from Ensembl IDs if present
clean_ensembl_all <- gsub("\\.\\d+$", "", sig_genes_all)</pre>
clean_ensembl_up <- gsub("\\.\\d+$", "", sig_genes_up)</pre>
clean_ensembl_down <- gsub("\\.\\d+$", "", sig_genes_down)</pre>
# Map to Entrez IDs
entrez_all <- bitr(clean_ensembl_all,</pre>
                   fromType = "ENSEMBL",
                   toType = "ENTREZID",
                   OrgDb = org.Hs.eg.db,
                   drop = TRUE)
## 'select()' returned 1:many mapping between keys and columns
## Warning in bitr(clean_ensembl_all, fromType = "ENSEMBL", toType = "ENTREZID", :
## 0.69% of input gene IDs are fail to map...
entrez_up <- bitr(clean_ensembl_up,</pre>
                  fromType = "ENSEMBL",
                  toType = "ENTREZID",
                  OrgDb = org.Hs.eg.db,
                  drop = TRUE)
## 'select()' returned 1:many mapping between keys and columns
## Warning in bitr(clean_ensembl_up, fromType = "ENSEMBL", toType = "ENTREZID", :
## 1.04% of input gene IDs are fail to map...
entrez_down <- bitr(clean_ensembl_down,</pre>
                    fromType = "ENSEMBL",
                    toType = "ENTREZID",
                    OrgDb = org.Hs.eg.db,
                    drop = TRUE)
## 'select()' returned 1:1 mapping between keys and columns
cat("Genes mapped to Entrez IDs:\n")
```

41

Genes mapped to Entrez IDs:

```
cat("- All significant:", nrow(entrez_all), "\n")
## - All significant: 144
cat("- Upregulated:", nrow(entrez_up), "\n")
## - Upregulated: 96
cat("- Downregulated:", nrow(entrez_down), "\n\n")
## - Downregulated: 48
# 1. Gene Ontology Biological Process (GO:BP) - All significant genes
cat("=== Running GO Biological Process Enrichment (All Genes) ===\n")
## === Running GO Biological Process Enrichment (All Genes) ===
go_bp_all <- enrichGO(gene = entrez_all$ENTREZID,</pre>
                      OrgDb = org.Hs.eg.db,
                      ont = "BP",
                      pAdjustMethod = "BH",
                      pvalueCutoff = 0.05,
                      qvalueCutoff = 0.2,
                      readable = TRUE)
go_bp_all_df <- as.data.frame(go_bp_all)</pre>
if (nrow(go_bp_all_df) > 0) {
  write_tsv(go_bp_all_df, file.path(results_dir, "GO_BP_all_genes_results.tsv"))
  # Create visualizations
  p1 <- dotplot(go_bp_all, showCategory = 20) +
   ggtitle("GO Biological Process - All Significant Genes") +
   theme(axis.text.y = element_text(size = 8))
  ggsave(file.path(plots_dir, "GO_BP_all_dotplot.png"), p1, width = 12, height = 8)
  p2 <- barplot(go bp all, showCategory = 15) +
   ggtitle("GO Biological Process - All Significant Genes")
  ggsave(file.path(plots_dir, "GO_BP_all_barplot.png"), p2, width = 10, height = 8)
  cat("Found", nrow(go_bp_all_df), "significant GO:BP terms (all genes)\n")
} else {
  cat("No significant GO:BP terms found (all genes)\n")
## Warning in fortify(object, showCategory = showCategory, by = x, ...): Arguments in '...' must be use
## x Problematic argument:
## * by = x
## i Did you misspell an argument name?
```

```
## Warning: 'aes_string()' was deprecated in ggplot2 3.0.0.
## i Please use tidy evaluation idioms with 'aes()'.
## i See also 'vignette("ggplot2-in-packages")' for more information.
## i The deprecated feature was likely used in the enrichplot package.
    Please report the issue at
##
     <https://github.com/GuangchuangYu/enrichplot/issues>.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
## Found 2 significant GO:BP terms (all genes)
# 2. Gene Ontology Biological Process (GO:BP) - Upregulated genes only
if (length(entrez_up$ENTREZID) > 5) {
  cat("\n=== Running GO Biological Process Enrichment (Upregulated) ===\n")
  go_bp_up <- enrichGO(gene = entrez_up$ENTREZID,</pre>
                       OrgDb = org.Hs.eg.db,
                       ont = "BP",
                       pAdjustMethod = "BH",
                       pvalueCutoff = 0.05,
                       qvalueCutoff = 0.2,
                       readable = TRUE)
  go_bp_up_df <- as.data.frame(go_bp_up)</pre>
  if (nrow(go_bp_up_df) > 0) {
   write_tsv(go_bp_up_df, file.path(results_dir, "GO_BP_upregulated_results.tsv"))
   p3 <- dotplot(go_bp_up, showCategory = 20) +
      ggtitle("GO Biological Process - Upregulated Genes") +
      theme(axis.text.y = element_text(size = 8))
    ggsave(file.path(plots_dir, "GO_BP_up_dotplot.png"), p3, width = 12, height = 8)
    cat("Found", nrow(go_bp_up_df), "significant GO:BP terms (upregulated)\n")
  } else {
    cat("No significant GO:BP terms found (upregulated)\n")
  }
}
## === Running GO Biological Process Enrichment (Upregulated) ===
## Found 15 significant GO:BP terms (upregulated)
# 3. Gene Ontology Biological Process (GO:BP) - Downregulated genes only
if (length(entrez down$ENTREZID) > 5) {
  cat("\n=== Running GO Biological Process Enrichment (Downregulated) ===\n")
  go_bp_down <- enrichGO(gene = entrez_down$ENTREZID,</pre>
                         OrgDb = org.Hs.eg.db,
                         ont = "BP",
                         pAdjustMethod = "BH",
                         pvalueCutoff = 0.05,
                         qvalueCutoff = 0.2,
                         readable = TRUE)
```

```
go_bp_down_df <- as.data.frame(go_bp_down)</pre>
  if (nrow(go_bp_down_df) > 0) {
   write_tsv(go_bp_down_df, file.path(results_dir, "GO_BP_downregulated_results.tsv"))
   p4 <- dotplot(go_bp_down, showCategory = 20) +
      ggtitle("GO Biological Process - Downregulated Genes") +
      theme(axis.text.y = element_text(size = 8))
    ggsave(file.path(plots dir, "GO BP down dotplot.png"), p4, width = 12, height = 8)
    cat("Found", nrow(go_bp_down_df), "significant GO:BP terms (downregulated)\n")
  } else {
    cat("No significant GO:BP terms found (downregulated)\n")
}
## === Running GO Biological Process Enrichment (Downregulated) ===
## No significant GO:BP terms found (downregulated)
# 4. Gene Ontology Molecular Function (GO:MF) - All significant genes
cat("\n=== Running GO Molecular Function Enrichment ===\n")
##
## === Running GO Molecular Function Enrichment ===
go_mf_all <- enrichGO(gene = entrez_all$ENTREZID,</pre>
                      OrgDb = org.Hs.eg.db,
                      ont = "MF",
                      pAdjustMethod = "BH",
                      pvalueCutoff = 0.05,
                      qvalueCutoff = 0.2,
                      readable = TRUE)
go_mf_all_df <- as.data.frame(go_mf_all)</pre>
if (nrow(go_mf_all_df) > 0) {
  write_tsv(go_mf_all_df, file.path(results_dir, "GO_MF_all_genes_results.tsv"))
 p5 <- dotplot(go mf all, showCategory = 20) +
   ggtitle("GO Molecular Function - All Significant Genes") +
    theme(axis.text.y = element_text(size = 8))
  ggsave(file.path(plots_dir, "GO_MF_all_dotplot.png"), p5, width = 12, height = 8)
  cat("Found", nrow(go_mf_all_df), "significant GO:MF terms\n")
} else {
  cat("No significant GO:MF terms found\n")
```

Found 4 significant GO:MF terms

```
# 5. Gene Ontology Cellular Component (GO:CC) - All significant genes
cat("\n=== Running GO Cellular Component Enrichment ===\n")
## === Running GO Cellular Component Enrichment ===
go_cc_all <- enrichGO(gene = entrez_all$ENTREZID,</pre>
                      OrgDb = org.Hs.eg.db,
                      ont = "CC",
                      pAdjustMethod = "BH",
                      pvalueCutoff = 0.05,
                      qvalueCutoff = 0.2,
                      readable = TRUE)
go_cc_all_df <- as.data.frame(go_cc_all)</pre>
if (nrow(go_cc_all_df) > 0) {
 write_tsv(go_cc_all_df, file.path(results_dir, "GO_CC_all_genes_results.tsv"))
 p6 <- dotplot(go_cc_all, showCategory = 20) +
    ggtitle("GO Cellular Component - All Significant Genes") +
   theme(axis.text.y = element_text(size = 8))
  ggsave(file.path(plots_dir, "GO_CC_all_dotplot.png"), p6, width = 12, height = 8)
  cat("Found", nrow(go_cc_all_df), "significant GO:CC terms\n")
} else {
  cat("No significant GO:CC terms found\n")
}
## Found 37 significant GO:CC terms
# 6. Create comprehensive summary table
cat("\n=== Creating Summary Table ===\n")
## === Creating Summary Table ===
summary_results <- data.frame()</pre>
# Helper function to add results to summary
add_to_summary <- function(results_df, method_name, ontology_name) {</pre>
  if (exists(deparse(substitute(results_df))) && nrow(results_df) > 0) {
      select(ID, Description, pvalue, p.adjust, qvalue, Count, GeneRatio, BgRatio) %>%
      mutate(Method = method_name,
             Ontology = ontology_name) %>%
      select (Method, Ontology, ID, Description, pvalue, p.adjust, qvalue, Count, GeneRatio, BgRatio)
  } else {
    data.frame()
 }
}
```

```
# Add all results to summary
if (exists("go_bp_all_df") && nrow(go_bp_all_df) > 0) {
  summary_results <- rbind(summary_results,</pre>
                           add_to_summary(go_bp_all_df, "GO_BP_All", "Gene Ontology Biological Process")
}
if (exists("go_bp_up_df") && nrow(go_bp_up_df) > 0) {
  summary_results <- rbind(summary_results,</pre>
                           add_to_summary(go_bp_up_df, "GO_BP_Up", "Gene Ontology Biological Process (Up
}
if (exists("go_bp_down_df") && nrow(go_bp_down_df) > 0) {
  summary_results <- rbind(summary_results,</pre>
                           add_to_summary(go_bp_down_df, "GO_BP_Down", "Gene Ontology Biological Process
}
if (exists("go_mf_all_df") && nrow(go_mf_all_df) > 0) {
  summary_results <- rbind(summary_results,</pre>
                           add_to_summary(go_mf_all_df, "GO_MF_All", "Gene Ontology Molecular Function")
}
if (exists("go_cc_all_df") && nrow(go_cc_all_df) > 0) {
  summary_results <- rbind(summary_results,</pre>
                           add_to_summary(go_cc_all_df, "GO_CC_All", "Gene Ontology Cellular Component")
}
# Save comprehensive results
if (nrow(summary_results) > 0) {
  summary_results <- summary_results %>%
    arrange(p.adjust)
  write_tsv(summary_results, file.path(results_dir, "GO_enrichment_comprehensive_results.tsv"))
  # Create top 25 results
  top_results <- summary_results %>%
    slice_head(n = 25)
  write_tsv(top_results, file.path(results_dir, "GO_enrichment_top25_results.tsv"))
  # Create method comparison
  method_summary <- summary_results %>%
    group_by(Method) %>%
    summarize(
      n_{terms} = n(),
      min_pvalue = min(pvalue),
      mean_pvalue = mean(pvalue),
      .groups = "drop"
    )
  write_tsv(method_summary, file.path(results_dir, "GO_method_comparison.tsv"))
  # Visualization of method comparison
  p_methods <- ggplot(method_summary, aes(x = Method, y = n_terms, fill = Method)) +</pre>
    geom_col() +
```

```
labs(title = "Number of Significant GO Terms by Analysis Method",
         x = "Analysis Method",
         y = "Number of Significant Terms") +
    theme_minimal() +
    theme(axis.text.x = element_text(angle = 45, hjust = 1),
          legend.position = "none")
  ggsave(file.path(plots_dir, "GO_method_comparison.png"), p_methods, width = 10, height = 6)
  cat("\nSummary table created with", nrow(summary_results), "total enriched GO terms\n")
  cat("Results saved to:", file.path(results_dir, "GO_enrichment_comprehensive_results.tsv"), "\n")
} else {
  cat("No significant enrichment results found\n")
##
## Summary table created with 58 total enriched GO terms
## Results saved to: results/GO_enrichment_comprehensive_results.tsv
# Final summary
cat("\n=== STEP 5 ANALYSIS COMPLETE ===\n")
##
## === STEP 5 ANALYSIS COMPLETE ===
cat("Method: clusterProfiler (PyDESeq2-style approach)\n")
## Method: clusterProfiler (PyDESeq2-style approach)
cat("Ontology: Gene Ontology (BP, MF, CC)\n")
## Ontology: Gene Ontology (BP, MF, CC)
cat("Parameters used:\n")
## Parameters used:
cat("- p-value cutoff: 0.05\n")
## - p-value cutoff: 0.05
cat("- q-value cutoff: 0.2\n")
## - q-value cutoff: 0.2
cat("- Multiple testing correction: Benjamini-Hochberg\n")
## - Multiple testing correction: Benjamini-Hochberg
```

```
cat("- Gene ID conversion: Ensembl to Entrez\n")
## - Gene ID conversion: Ensembl to Entrez
cat("\nInput:\n")
## Input:
cat("- Total DE genes:", length(sig_genes_all), "\n")
## - Total DE genes: 144
cat("- Upregulated genes:", length(sig_genes_up), "\n")
## - Upregulated genes: 96
cat("- Downregulated genes:", length(sig_genes_down), "\n")
## - Downregulated genes: 48
if (exists("summary_results") && nrow(summary_results) > 0) {
  cat("\nResults:\n")
  cat("- Total significant GO terms:", nrow(summary_results), "\n")
  cat("- Most significant term p-value:", format(min(summary_results$p.adjust), scientific = TRUE), "\n
##
## Results:
## - Total significant GO terms: 58
## - Most significant term p-value: 1.575203e-21
```

Section 5 Summary

Gene set enrichment with clusterProfiler returned 58 significant Gene Ontology terms across BP, MF, ## The topGO run contributed 200 Biological Process terms (with 79 passing weight01 < 0.05), and the su

Section 6

Section 6a: Consolidate enrichment results across methods

```
library(dplyr)
library(readr)
library(tidyr)
```

```
##
## Attaching package: 'tidyr'
## The following object is masked from 'package:magrittr':
##
##
       extract
## The following object is masked from 'package:S4Vectors':
##
       expand
library(stringr)
library(purrr)
##
## Attaching package: 'purrr'
## The following object is masked from 'package:clusterProfiler':
##
       simplify
## The following object is masked from 'package:GenomicRanges':
##
##
       reduce
## The following object is masked from 'package:magrittr':
##
##
       set_names
## The following object is masked from 'package: IRanges':
##
##
       reduce
# helper to tidy method names
clean_method_name <- function(x) {</pre>
  x %>%
    stringr::str_replace_all("[^A-Za-z0-9]+", "_") %>%
    stringr::str_replace_all("_+", "_") %>%
    stringr::str_replace_all("^_|_$", "") %>%
    stringr::str_to_lower()
}
parse_pvalue <- function(x) {</pre>
  if (is.numeric(x)) return(x)
  ifelse(is.na(x), NA_real_, suppressWarnings(as.numeric(stringr::str_replace_all(x, "<\\s*", ""))))
}
enrichment_tables <- list()</pre>
# clusterProfiler results from current session or saved file
if (exists("summary_results") && nrow(summary_results) > 0) {
```

```
cp_long <- summary_results %>%
    transmute(
      source_method = "clusterProfiler",
      method detail = Method,
      term_id = ID,
      term_name = Description,
      p_value = as.numeric(pvalue),
      adj p value = as.numeric(p.adjust),
      q_value = as.numeric(qvalue)
    mutate(significant = ifelse(!is.na(adj_p_value), adj_p_value < 0.05, ifelse(!is.na(p_value), p_value)
  enrichment_tables <- append(enrichment_tables, list(cp_long))</pre>
  cp_path <- file.path(results_dir, "GO_enrichment_comprehensive_results.tsv")</pre>
  if (file.exists(cp_path)) {
    cp_file <- readr::read_tsv(cp_path, show_col_types = FALSE)</pre>
    cp_long <- cp_file %>%
      transmute(
        source_method = "clusterProfiler",
        method_detail = Method,
        term id = ID,
        term_name = Description,
        p_value = as.numeric(pvalue),
        adj_p_value = as.numeric(p.adjust),
        q value = as.numeric(qvalue)
      ) %>%
      mutate(significant = ifelse(!is.na(adj_p_value), adj_p_value < 0.05, ifelse(!is.na(p_value), p_va
    enrichment_tables <- append(enrichment_tables, list(cp_long))</pre>
}
# clusterProfiler KEGG results
if (exists("kegg_results_df") && nrow(kegg_results_df) > 0) {
  method_label <- if (exists("run_label")) paste0("KEGG_", run_label) else "KEGG_current_run"</pre>
  kegg_long <- kegg_results_df %>%
    transmute(
      source_method = "clusterProfiler",
      method detail = method label,
      term_id = ID,
      term_name = Description,
      p_value = as.numeric(pvalue),
      adj_p_value = as.numeric(p.adjust),
      q_value = as.numeric(qvalue)
    ) %>%
    mutate(significant = ifelse(!is.na(adj_p_value), adj_p_value < 0.05, ifelse(!is.na(p_value), p_value)
  enrichment_tables <- append(enrichment_tables, list(kegg_long))</pre>
} else {
  kegg_files <- list.files(results_dir, pattern = "^KEGG_enrichment_.*\\.tsv$", full.names = TRUE)
  kegg_files <- kegg_files[!grepl("_top20_", kegg_files)]</pre>
  if (length(kegg_files) > 0) {
    kegg_tables <- purrr::map(kegg_files, ~{</pre>
      df <- readr::read_tsv(.x, show_col_types = FALSE)</pre>
      method_label <- tools::file_path_sans_ext(basename(.x))</pre>
```

```
df %>%
        transmute(
          source_method = "clusterProfiler",
          method detail = method label,
          term_id = ID,
          term_name = Description,
          p_value = as.numeric(pvalue),
          adj_p_value = as.numeric(p.adjust),
          q_value = as.numeric(qvalue)
        mutate(significant = ifelse(!is.na(adj_p_value), adj_p_value < 0.05, ifelse(!is.na(p_value), p_
    enrichment_tables <- append(enrichment_tables, kegg_tables)</pre>
}
# Team methods saved to results directory
topgo_files <- list.files(results_dir, pattern = "^topGO_.*\\.tsv$", full.names = TRUE)</pre>
if (length(topgo_files) > 0) {
  topgo_tables <- purrr::map(topgo_files, ~{</pre>
    df <- readr::read_tsv(.x, show_col_types = FALSE)</pre>
    method_label <- tools::file_path_sans_ext(basename(.x))</pre>
    p_cols <- c("weight01Fisher_p", "weight01Fisher", "classicFisher_p", "classicFisher", "Fisher")
    first_p <- purrr::detect(p_cols, ~ .x %in% names(df))</pre>
    p_vals <- if (!is.null(first_p)) parse_pvalue(df[[first_p]]) else rep(NA_real_, nrow(df))</pre>
    tibble::tibble(
      source_method = "topGO",
      method_detail = method_label,
      term_id = df$`GO.ID`,
      term_name = df$Term,
      p_value = p_vals,
      adj_p_value = NA_real_,
      q_value = NA_real_
    ) %>%
      mutate(significant = ifelse(!is.na(p_value), p_value < 0.05, FALSE))</pre>
 })
  enrichment_tables <- append(enrichment_tables, topgo_tables)</pre>
gss_files <- list.files(results_dir, pattern = "^GSS_.*\\.tsv$|^GSS_validated_signatures_.*\\.tsv$", fu
if (length(gss_files) > 0) {
  gss_tables <- purrr::map(gss_files, ~{</pre>
    df <- readr::read_tsv(.x, show_col_types = FALSE)</pre>
    method_label <- tools::file_path_sans_ext(basename(.x))</pre>
    df %>%
      transmute(
        source_method = "GSS",
        method_detail = method_label,
        term_id = as.character(term_id),
        term_name = as.character(term_name),
        p_value = NA_real_,
        adj_p_value = NA_real_,
        q_value = NA_real_,
```

```
score = ifelse("score" %in% names(df), as.numeric(df$score), NA_real_)
      mutate(significant = FALSE)
 })
  enrichment_tables <- append(enrichment_tables, gss_tables)</pre>
if (length(enrichment tables) == 0) {
  warning("No enrichment result files found. Run Step 5 for each team method before executing Section 6
} else {
  enrichment_long <- bind_rows(enrichment_tables) %>%
   mutate(
      method_readable = paste(source_method, method_detail, sep = ": "),
      method_clean = clean_method_name(method_readable)
  min_non_na <- function(...) {</pre>
   vals <- c(...)
   vals <- vals[!is.na(vals)]</pre>
   if (length(vals) == 0) NA_real_ else min(vals)
  }
  metrics_to_pivot <- c("p_value", "adj_p_value", "q_value", "significant")</pre>
  metrics wide <- enrichment long %>%
   pivot_longer(cols = all_of(metrics_to_pivot), names_to = "metric", values_to = "metric_value") %>%
   mutate(metric column = paste(method clean, metric, sep = " ")) %>%
    select(term_id, term_name, metric_column, metric_value) %>%
   distinct() %>%
   pivot_wider(names_from = metric_column, values_from = metric_value)
  method_counts <- enrichment_long %>%
    group_by(term_id, term_name) %>%
    summarise(
      methods_included = n_distinct(method_readable),
      methods_significant = sum(significant, na.rm = TRUE),
      methods_list = paste(sort(unique(method_readable)), collapse = "; "),
      min_p_value = min_non_na(p_value, adj_p_value, q_value),
      .groups = "drop"
   )
  combined_table <- method_counts %>%
   left_join(metrics_wide, by = c("term_id", "term_name")) %>%
    arrange(desc(methods_significant), desc(methods_included), min_p_value)
  assign("combined_enrichment_table", combined_table, envir = .GlobalEnv)
  combined_path <- file.path(results_dir, "combined_enrichment_summary.tsv")</pre>
  readr::write_tsv(combined_table, combined_path)
  cat(sprintf("Combined enrichment summary saved to %s with %d unique terms across %d method contributi
              combined_path, nrow(combined_table), length(unique(enrichment_long$method_readable))))
```

Combined enrichment summary saved to results/combined_enrichment_summary.tsv with 278 unique terms a

Section 6b Summary

Cross-method comparison pooled 9 enrichment outputs, highlighting 160 terms that were significant in

Section 7

Section 7a: Top shared enrichment signals

```
if (exists("combined_enrichment_table")) {
  top_terms <- combined_enrichment_table %>%
    arrange(desc(methods_significant), desc(methods_included), min_p_value) %>%
    { .[seq_len(min(10, nrow(.))), , drop = FALSE] }
  assign("top_shared_terms", top_terms, envir = .GlobalEnv)
  top10_path <- file.path(results_dir, "combined_enrichment_top10.tsv")
  readr::write_tsv(top_terms, top10_path)
  print(top_terms %>% select(term_id, term_name, methods_significant, methods_included, min_p_value))
  cat(sprintf("\nTop consensus terms saved to %s.\n", top10_path))
} else {
  cat("Run Section 6 first to build the combined enrichment table.\n")
## # A tibble: 10 x 5
##
     {\tt term\_id}
              term_name
                                   methods_significant methods_included min_p_value
##
      <chr>
                <chr>
                                                 <int>
                                                                  <int>
                                                     5
                                                                      5
                                                                               e-30
## 1 GO:0002181 cytoplasmic tran~
                                                                           1
## 2 GO:0030593 neutrophil chemo~
                                                     4
                                                                      4
                                                                           3.26e- 6
                                                                      2
## 3 GO:0045648 positive regulat~
                                                     2
                                                                               e- 4
                                                                           1
## 4 GO:0017148 negative regulat~
                                                     2
                                                                      2
                                                                           2.1 e- 4
                                                    2
                                                                      2
                                                                          2.7 e- 4
## 5 GO:0006364 rRNA processing
## 6 GO:0071219 cellular respons~
                                                    2
                                                                      2
                                                                           2.5 e- 3
## 7 GO:0006617 SRP-dependent co~
                                                     2
                                                                      2
                                                                           4.1 e- 3
                                                     2
                                                                      2
                                                                           4.1 e- 3
## 8 GO:0034135 regulation of to~
## 9 GO:0036446 myofibroblast di~
                                                                      2
                                                                           4.1 e- 3
## 10 GO:0071361 cellular respons~
                                                                           4.1 e- 3
##
```

Section 7b Summary

Across methods, cytoplasmic translation emerged as the most consistently enriched term, reported by

Top consensus terms saved to results/combined_enrichment_top10.tsv.