Figure 5

${\bf EmptyNN}$ - ${\bf Figure}$ 5

The following code reproduces the Figure 5 in our EmptyNN manuscript.

Please download datasets and seurat objects before running this analysis (run download_datasets.sh in terminal)

Load libraries

```
library("Seurat")

## Warning: package 'Seurat' was built under R version 4.0.5

## Attaching SeuratObject

library("Matrix")

## Warning: package 'Matrix' was built under R version 4.0.5

library("ggplot2")

## Warning: package 'ggplot2' was built under R version 4.0.5

library("pheatmap")
load("./../data/BlueYellowColormaps_V1.RData")

## Version 3.6.3, Seurat_3.2.3, Matrix_1.3-2, ggplot_2_3.3.3, pheatmap_1.0.12
```

Load seurat objects containing barcodes retained by EmptyNN or CellRanger 2.0

```
pbmc8k_retained <- readRDS("./../data/pbmc_8k_retained.rds")
neuron900_retained <- readRDS("./../data/neuron900_retained.rds")</pre>
```

Define functions: runSeurat() and plot_heatmap()

```
runSeurat <- function(seu,RNA.thres,mt.thres,resolution,verbose){</pre>
        seu[["percent.mt"]] <- PercentageFeatureSet(seu, pattern = "^MT|^mt")</pre>
        seu <- subset(seu, subset = nFeature_RNA > RNA.thres &
                                percent.mt < mt.thres)</pre>
        seu <- NormalizeData(seu, verbose = verbose)</pre>
        seu <- FindVariableFeatures(seu, verbose = verbose)</pre>
        seu <- ScaleData(seu, features = VariableFeatures(seu), verbose = verbose)</pre>
        seu <- RunPCA(seu,features=VariableFeatures(seu),verbose = verbose)</pre>
        seu <- FindNeighbors(seu, dims = 1:10, verbose = verbose)</pre>
        seu <- FindClusters(seu, resolution = resolution, verbose = verbose)</pre>
        seu <- RunTSNE(seu, dims = 1:10, check_duplicates = verbose, verbose = verbose)</pre>
        return(seu)
plot_heatmap <- function(seu,n_top,title){</pre>
        seu <- seu[-grep("^RPL|^RPS|^MT|^Rps|^Rp1|^mt", rownames(seu)),]</pre>
        des <- FindAllMarkers(seu, only.pos = TRUE, min.pct = 0.25,</pre>
                                logfc.threshold = 0.25, verbose=FALSE)
        asplit_genes <- split(1:nrow(des), des$cluster)</pre>
        # take top n genes
        genes <- unlist(lapply(asplit_genes, function(x) des[x[1:n_top], "gene"]))</pre>
        # Average cells within each cluster
        asplit_cells <- split(rownames(seu@meta.data), seu@active.ident)</pre>
        means <- do.call(cbind, lapply(asplit_cells, function(x){</pre>
        s1 <- Matrix::rowMeans(seu@assays$RNA@data[genes, sample(unlist(x), 10)])
        s2 <- Matrix::rowMeans(seu@assays$RNA@data[genes, sample(unlist(x), 10)])
        s3 <- Matrix::rowMeans(seu@assays$RNA@data[genes, sample(unlist(x), 10)])
        cbind(s1, s2, s3)
}))
        cell_type <- unlist(lapply(names(asplit_cells), function(x) rep(x, 3)))</pre>
        # Create heatmap (sample 3 "replicates")
        anno_col <- data.frame(cell_type)</pre>
        rownames(anno_col) <- colnames(means) <- paste(colnames(means), cell_type)</pre>
        pheatmap(means,cluster_rows = F, cluster_cols = F, scale = "row",
                  breaks = seq(-2, 2, length = length(yellow2blue) + 1), col = yellow2blue,
                  annotation_col = anno_col,show_colnames = F,main=title)
```

Figure 5A

```
## Warning in simpleLoess(y, x, w, span, degree = degree, parametric =
## parametric, : reciprocal condition number 2.8276e-028

## Warning in simpleLoess(y, x, w, span, degree = degree, parametric =
## parametric, : There are other near singularities as well. 0.090619

new.cluster <- c("CD4", "CD14 Mono", "Platelet")
names(new.cluster) <- levels(recover)
recover <- RenameIdents(recover,new.cluster)
DimPlot(recover,label=T)+NoLegend()+labs(title="PBMC_8k_dataset_recovered")</pre>
```

PBMC_8k_dataset_recovered

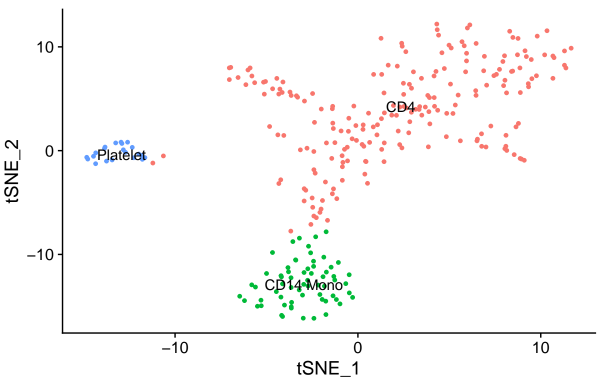


Figure 5B

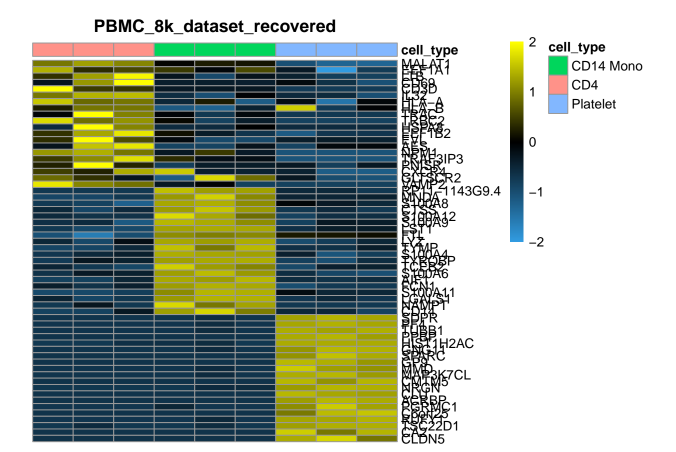


Figure 5C

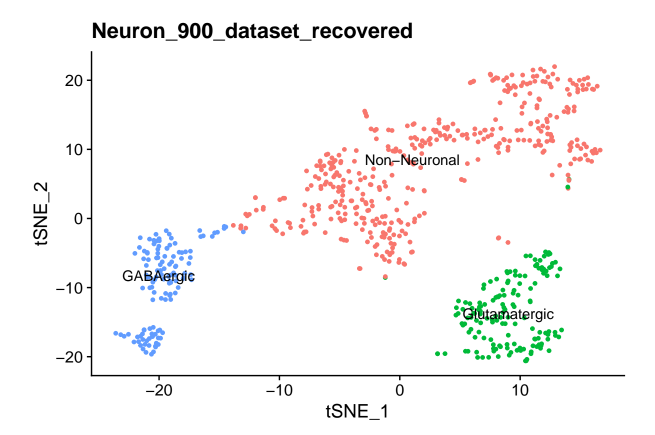


Figure 5D

plot_heatmap(recover,n_top=5,"Neuron_900_dataset_recovered")



