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Single cell analysis of iPSC-derived midbrain organoids

In 6 collections

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ABSTRACT

The following script was used for analysis of gene corrected (GC) versus GBA1 mutant (MUT) midbrain organoids. The purpose was to combine, filter, integrate, and identify clusters and differentially expressed genes sets.

ATTACHMENTS

404-875.docx

GUIDELINES

This script is based on the Seurat tutorials

- https://satijalab.org/seurat/articles/integration_introduction.html
- https://ucdavis-bioinformatics-training.github.io/2019-single-cell-RNA-sequencing-Workshop-UCD_UCSF/scrnaseq_analysis/scRNA_Workshop-PART1.html

1

```
# Install and load the libraries
library(Seurat)
library(patchwork)
library(ggplot2)
library(cowplot)
library(magrittr)
library(tidyverse)
# Load and combine 10x Runs
setwd("D:.../analysis results")
GC1.data <-Read10X h5("1GC filtered feature bc matrix.h5")
GC1 <-CreateSeuratObject(counts = GC1.data)</pre>
GC1
MUT1.data <-Read10X h5("1MUT filtered feature bc matrix.h5")
MUT1 <-CreateSeuratObject(counts = MUT1.data)</pre>
MUT1
GC2.data <-Read10X h5("2GC filtered feature bc matrix.h5")
GC2 <-CreateSeuratObject(counts = GC2.data)</pre>
GC2
MUT2.data <-Read10X_h5("2MUT_filtered_feature_bc_matrix.h5")</pre>
MUT2 <-CreateSeuratObject(counts = MUT2.data)</pre>
MUT2
GC3.data <-Read10X h5("3GC filtered feature bc matrix.h5")
GC3 <-CreateSeuratObject(counts = GC3.data)</pre>
GC3
MUT3.data <-Read10X_h5("3MUT_filtered_feature_bc_matrix.h5")</pre>
MUT3 <-CreateSeuratObject(counts = MUT3.data)</pre>
MUT3
# Merge more than two objects
MUT.combined <-Reduce(function(x,y) merge (x,y, all=T), list (GC1,
MUT1, GC2, MUT2, GC3, MUT3))
```

Part 2. Quality control

```
# Add number of genes per UMI for each cell to metadata
MUT.combined$log10GenesPerUMI <-log10(MUT.combined$nFeature RNA)</pre>
/log10(MUT.combined$nCount RNA)
# Compute percent mito ratio
MUT.combined$mitoRatio <-PercentageFeatureSet(object =
MUT.combined, pattern ="^MT-")
MUT.combined$mitoRatio <- MUT.combined@meta.data$mitoRatio /100
# Create .RData object
save(MUT.combined, file ="Seurat Object.RData")
# Filter out low quality reads using selected thresholds
filtered seurat <-subset(x =MUT.combined,subset= (nCount RNA >=500)
&(nFeature RNA >=250) &(log10GenesPerUMI >0.80) & (mitoRatio
<0.20)
# Extract counts
counts <-GetAssayData(object =filtered seurat, slot ="counts")</pre>
# Output a logical vector for every gene on whether the more than
zero counts per cell
nonzero <- counts >0
# Sums all TRUE values and returns TRUE if more than 10 TRUE values
per gene
keep genes <- Matrix::rowSums(nonzero) >=10
# Only keeping those genes expressed in more than 10 cells
filtered counts <-counts[keep genes, ]</pre>
# Reassign to filtered Seurat object
filtered seurat <-CreateSeuratObject(filtered counts, meta.data</pre>
=filtered seurat@meta.data)
# Save.RData object
save(filtered seurat, file="seurat_filtered.RData")
```

Part 3. Data preparation and normalization

```
#Cell cycle scoring. Normalize the counts
seurat phase <-NormalizeData(filtered seurat)</pre>
# Load cell cycle markers
load("cycle.rda")
# Score cells for cell cycle
seurat phase <-CellCycleScoring(seurat phase,g2m.features =</pre>
q2m genes,s.features =s genes)
# View cell cycle scores and phases assigned to cells
View(seurat phase@meta.data)
# Identify the most variable genes
seurat phase <-FindVariableFeatures(seurat phase, selection.method
="vst",nfeatures =2000, verbose =FALSE)
# Scale the counts
seurat phase <-ScaleData(seurat phase)</pre>
# Perform PCA
seurat_phase <-RunPCA(seurat_phase)</pre>
# Plot the PCA colored by cell cycle phase
DimPlot(seurat phase, reduction = "pca", group.by="Phase", split.by
="Phase")
# Split seurat object by condition to perform cell cycle scoring
and SCT on all samples
split seurat <-SplitObject(filtered seurat, split.by ="orig.ident")</pre>
for (i in1:length(split_seurat)) { split_seurat[[i]] <-</pre>
NormalizeData(split seurat[[i]], verbose =TRUE)
split seurat[[i]] <-CellCycleScoring(split seurat[[i]],</pre>
g2m.features=g2m genes, s.features=s genes)
split seurat[[i]] <-SCTransform(split seurat[[i]], vars.to.regress</pre>
=c("mitoRatio", "S.Score", "G2M.Score")) }
```

Part 4. Data integration and visualization (directly after Pa..

4

Select the most variable features to use for integration
integ_features <-SelectIntegrationFeatures(object.list</pre>

```
=split seurat,nfeatures =3000)
# Prepare the SCT list object for integration
split seurat <-PrepSCTIntegration(object.list</pre>
=split seurat,anchor.features =integ features)
# Find anchors
integ anchors <-FindIntegrationAnchors(object.list</pre>
=split seurat,normalization.method ="SCT", anchor.features
=integ features)
# Integrate across conditions
organoids.combined.sct <-IntegrateData(anchorset</pre>
=integ anchors,normalization.method ="SCT")
# Save integrated seurat object
saveRDS(organoids.combined.sct, "integrated seurat.rds")
## Run the standard workflow for visualization and clustering
DefaultAssay(organoids.combined.sct) <-"integrated"</pre>
organoids.combined.sct <-RunPCA(organoids.combined.sct, verbose
=FALSE)
PCAPlot(organoids.combined.sct,split.by ="orig.ident")
organoids.combined.sct <-RunUMAP(organoids.combined.sct, reduction
="pca", dims =1:40)
# Plot UMAP
DimPlot(organoids.combined.sct, split.by ="orig.ident")
DimPlot(organoids.combined.sct, group.by ="orig.ident")
# Elbow plot
ElbowPlot(object =organoids.combined.sct,ndims =40)
#Find neighbors for cluster analysis
organoids.combined.sct <-FindNeighbors(organoids.combined.sct,
reduction ="pca", dims =1:40)
organoids.combined.sct <-FindClusters(organoids.combined.sct,</pre>
resolution =c(0.6))
# Assign identity of clusters
Idents(object =organoids.combined.sct) <-"integrated snn res.0.6"</pre>
# Visualization
all.genes <-rownames(organoids.combined.sct)
organoids.combined.sct <-ScaleData(organoids.combined.sct, features</pre>
=all.genes)
```

```
DimPlot(organoids.combined.sct, reduction ="umap", group.by
="seurat_clusters", label =TRUE, repel =TRUE)

DimPlot(organoids.combined.sct, reduction ="umap", split.by
="orig.ident")
```

Part 5. Obtain information from the datasets

```
#Extract identity and sample information from seurat object to
determine the number of cells per cluster per sample
n cells <-FetchData(organoids.combined.sct,vars =c("ident",</pre>
"orig.ident")) %>% dplyr::count(ident, orig.ident) %>%
tidyr::spread(ident, n)
write.csv(n cells, "n cells.csv")
## Identify conserved cell type markers
# Switch back to the original data
DefaultAssay(organoids.combined.sct) <- "RNA"</pre>
annotations <-read.csv("annotation.txt")</pre>
# conserved markers to any cluster
get conserved <-function(cluster)</pre>
{FindConservedMarkers(organoids.combined.sct, ident.1 =cluster,
grouping.var ="orig.ident", only.pos =TRUE)
%>%rownames to column(var = "gene") %>%left join(y
=unique(annotations[, c("gene_name", "description")]),by
=c("gene"="gene name")) %>%cbind(cluster id =cluster, .)}
# Iterate function across desired clusters.
conserved markers <-map dfr(0:20, get conserved)</pre>
# Extract top 100 markers per cluster
top100 <-conserved markers %>%mutate(avg fc =(GC1 avg log2FC
+GC2 avg log2FC +GC3 avg log2FC + MUT1 avg log2FC +MUT2 avg log2FC
+MUT3 avg log2FC) /6) %>%group by(cluster id) %>%top n(n =100,wt =
avg fc)
write.csv(top100, "Clusters top100.csv")
## Identifying gene markers for each cluster
```

```
Genes <-c ("SOX2", "DCX", "TH", "NEUROD4") # Change target genes
depending on the cell type

# UMAP plot
FeaturePlot(organoids.combined.sct,reduction ="umap",features =
Genes, sort.cell =TRUE, min.cutoff ='q10', max.cutoff =5,label = T,
pt.size =0.5)

# Violin plot
plots <-VlnPlot(organoids.combined.sct, features =Genes, split.by
="orig.ident", pt.size =0, combine =FALSE)
wrap_plots(plots = plots, ncol =1)

# Dot plot
DotPlot(organoids.combined.sct, features =Genes) +RotatedAxis()</pre>
```

Part 6. Merge and analyse subclusters

```
## Combine the clusters according to the identity
new.cluster.ids <-c(1="Radial Glia",</pre>
2="Neurons",
3="NPC",
4="Oligodendrocytes",
5="Astrocytes", "...")
names(new.cluster.ids) <-levels(organoids.combined.sct)</pre>
organoids.combined.newnames <-RenameIdents(organoids.combined.sct,
new.cluster.ids)
DimPlot(organoids.combined.newnames, reduction = "umap", label
=TRUE, pt.size =0.5) +NoLegend()
#Using DE analysis in specific clusters (after merging) MAST
annotations <-read.csv("annotation.txt")</pre>
Markers <-FindMarkers(organoids.combined.newnames, ident.1 = "GC1",</pre>
ident.2 ="MUT1", group.by ="orig.ident", subset.ident ="Radial
Glia", min.pct =0.1, test.use ="MAST")%>%rownames to column(var
="gene") %>%left_join(y =unique(annotations[, c("gene_name",
"description")]),by =c("gene"="gene name"))
#Save
write.csv(Markers, "DEgenes cRadial Glia 1couple.csv")
```

Part 7. Create a subset of cells from a selected cluster to re.

```
Neurons_subset <-subset(organoids.combined.newnames, idents
="Neurons")
Neurons_subset

DefaultAssay(Neurons_subset) <-"integrated"

# Run the standard workflow for visualization and clustering
Neurons_subset <-RunPCA(Neurons_subset, verbose =FALSE)

# Plot PCA
PCAPlot(Neurons_subset,
split.by ="orig.ident")</pre>
```

```
#Run variable features
Neurons subset <-
FindVariableFeatures(Neurons subset, selection.method ="vst",
nfeatures =2000, verbose =FALSE)
# Scale the counts
Neurons subset <-ScaleData(Neurons subset)</pre>
#Find neighbors for cluster analysis
Neurons subset <-FindNeighbors(Neurons subset, reduction = "pca",
dims = 1:40)
Neurons subset <-FindClusters(Neurons subset, resolution =0.4)
# Assign identity of clusters
Idents(object =Neurons subset) <-"integrated snn res.0.4"</pre>
# Visualization
all.genes <-rownames(Neurons subset)</pre>
Neurons subset <-ScaleData(Neurons subset, features =all.genes)</pre>
DimPlot(Neurons subset, reduction ="umap", group.by ="orig.ident")
DimPlot(Neurons subset, reduction ="umap")
# to remove a non-specific cluster
Finalcluster <-subset(Neurons subset, idents =5, invert = T)</pre>
DimPlot(Finalcluster, reduction ="umap")
# To visualize the two conditions side-by-side
DimPlot(Neurons_subset, reduction ="umap", split.by ="orig.ident")
## Cluster identification. Find conserved markers to any cluster
DefaultAssay(organoids.combined.sct) <- "RNA"</pre>
get conserved <-function(cluster)</pre>
{FindConservedMarkers(Neurons subset,ident.1 = cluster,
grouping.var ="orig.ident",only.pos =TRUE)
%>%rownames to column(var = "gene") %>%left join(y
=unique(annotations[, c("gene_name", "description")]), by
=c("gene"="gene name")) %>%cbind(cluster id = cluster, .)}
# Iterate function across desired clusters.
conserved markers <-map dfr(0:8, get conserved)</pre>
# Extract top 100 markers per cluster
top100 <-conserved markers %>%
mutate(avg fc =(GC1 avg log2FC +GC2 avg log2FC +GC3 avg log2FC +
```

```
MUT1_avg_log2FC +MUT2_avg_log2FC +MUT3_avg_log2FC) /6) %>%
group_by(cluster_id) %>%
top_n(n =100,
wt = avg_fc)

#OR save
write.csv(top100, "Clusters_top100_Neuronsubset.csv")
```

Part 8. Differential expressed genes using FIndMarkers wit...

```
annotations <-read.csv("annotation.txt")</pre>
# couple 1
Markers <-FindMarkers(Neurons subset, ident.1 = "GC1", ident.2</pre>
="MUT1", group.by ="orig.ident", subset.ident ="11", min.pct =0.1,
test.use ="MAST")%>%
rownames to column(var = "gene") %>%left join(y
=unique(annotations[, c("gene name", "description")]), by
=c("gene"="gene name"))
write.csv(Markers, "DEgenes_C11Neuron_1couple.csv")
# couple 2
Markers <-FindMarkers(Neurons subset, ident.1 = "GC2", ident.2</pre>
="MUT2", group.by ="orig.ident", subset.ident ="11", min.pct =0.1,
test.use ="MAST")%>%rownames to column(var = "gene") %>%left join(y
=unique(annotations[, c("gene name", "description")]),by
=c("gene"="gene name"))
write.csv(Markers, "DEgenes C11Neuron 2couple.csv")
# couple 3
Markers <-FindMarkers(Neurons subset, ident.1 = "GC3", ident.2</pre>
="MUT3", group.by ="orig.ident", subset.ident ="11", min.pct =0.1,
test.use ="MAST")%>%
rownames to column(var = "gene") %>%left join(y
=unique(annotations[, c("gene name", "description")]), by
=c("gene"="gene name"))
write.csv(Markers, "DEgenes C11Neuron 3couple.csv")
```

```
# Merge datasets, example analysis for couple 1
organoids.combined.sct <-Reduce(function(x,y) merge(x,y, all=T),
list (split seurat$GC1, split seurat$MUT1))
integ features <-SelectIntegrationFeatures(object.list</pre>
=split seurat, nfeatures =3000)
VariableFeatures(organoids.combined.sct[["SCT"]]) <-integ_features</pre>
## Run the standard workflow for visualization and clustering
DefaultAssay(organoids.combined.sct) <- "integrated"</pre>
organoids.combined.sct <-RunPCA(organoids.combined.sct, verbose
=FALSE)
PCAPlot(organoids.combined.sct,split.by ="orig.ident")
organoids.combined.sct <-RunUMAP(organoids.combined.sct, reduction
="pca", dims =1:40)
# Plot UMAP
DimPlot(organoids.combined.sct, split.by ="orig.ident")
DimPlot(organoids.combined.sct, group.by ="orig.ident")
# Elbow plot
ElbowPlot(object =organoids.combined.sct,ndims =40)
#Find neighbors for cluster analysis
organoids.combined.sct <-FindNeighbors(organoids.combined.sct,
reduction ="pca", dims =1:40)
organoids.combined.sct <-FindClusters(organoids.combined.sct,</pre>
resolution =c(0.6))
# Assign identity of clusters
Idents(object =organoids.combined.sct) <-"integrated snn res.0.6"</pre>
# Visualization
all.genes <-rownames(organoids.combined.sct)</pre>
organoids.combined.sct <-ScaleData(organoids.combined.sct, features
=all.genes)
DimPlot(organoids.combined.sct, reduction ="umap", group.by
="seurat_clusters", label =TRUE, repel =TRUE)
```

```
DimPlot(organoids.combined.sct, reduction ="umap", split.by
="orig.ident")

## Continue withPart 5 and/or 6 if needed

## To analyze the clusters with resolution of 0.6
annotations <-read.csv("annotation.txt")

# Change the cluster as needed
Markers <-FindMarkers(organoids.combined.sct, ident.1 ="GC1", ident.2 ="MUT1", group.by ="orig.ident", subset.ident ="20", min.pct =0.1, test.use ="MAST")%>%rownames_to_column(var ="gene")%>%left_join(y =unique(annotations[, c("gene_name", "description")]), by =c("gene"="gene_name"))

#save
write.csv(Markers, "DEgenes_Cluster20_couple1.csv")
```