Seurat - Clustering Tutorial

Dennis Dimitri Krutkin

```
2023-06-16
```

```
#Load necessary libraries
library(dplyr)
## Attaching package: 'dplyr'
```

```
## The following objects are masked from 'package:stats':
##
##
       filter, lag
```

```
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
```

```
library(Seurat)
```

```
## Attaching SeuratObject
library(patchwork)
```

Import 10X dataset # Load the PBMC data set:

```
pbmc.data <- Read10X(data.dir = "./filtered_gene_bc_matrices/hg19/")</pre>
# Initialize the Seurat object with the raw (non-normalized data):
pbmc <- CreateSeuratObject(counts = pbmc.data, project = "pbmc3k", min.cells = 3, min.features = 200)</pre>
## Warning: Feature names cannot have underscores ('_'), replacing with dashes
## ('-')
```

```
pbmc
## An object of class Seurat
## 13714 features across 2700 samples within 1 assay
## Active assay: RNA (13714 features, 0 variable features)
```

Examine a 3 genes in the first 25 cells: pbmc.data[c("CD3D", "TCL1A", "MS4A1"), 1:25]

Examine the loaded dataset

```
## 3 x 25 sparse Matrix of class "dgCMatrix"
    [[ suppressing 25 column names 'AAACATACAACCAC-1', 'AAACATTGAGCTAC-1', 'AAACATTGATCAGC-1' ...
##
]]
##
## CD3D 4 . 10 . . 1 2 3 1 . . 2 7 1 . . 1 3 . 2 3 . . . .
```

#Check the full size of object: dense.size <- object.size(as.matrix(pbmc.data))</pre>

#Check the ratio of full-size:sparse-size:

dense.size/sparse.size

23.7 bytes

highly variable features.

The "." values in the matrix represent 0s (no molecules detected).

significant memory and speed savings for Drop-seq/inDrop/10x data.

dense.size

Most values in a scRNA-seq matrix are 0 - Seurat uses a sparse-matrix representation whenever possible. This results in

```
## 709591472 bytes
#Check the sparse-represented size:
sparse.size <- object.size(pbmc.data)</pre>
```

```
sparse.size
## 29905192 bytes
```

```
Standard pre-processing workflow
The steps below demonstrate the standard pre-processing workflow for scRNA-seq data using Seurat.
The steps include selection and filtration of cells based on QC metrics, data normalization and scaling, and the detection of
```

Some QC metrics used by the community include: The number of unique genes detected in each cell

Quality Control and selecting cells to further analyze

 The percentage of reads which map to the mitochondrial genome Poor quality and dying cells frequently show extensive mitochondrial contamination The PercentageFeatureSet() function calculates the percentage of counts originating from a set of features to

The total number of molecules detected within a cell, which correlate strongly with unique genes

In Seurat, QC metrics can be explored and cells can be filtered according to any user-defined criteria.

Genes starting with MT- are pooled as a set of mitochondrial genes

#Add QC stats for mitochondrial genes to the object metadata:

calculate mitochondrial QC metrics

They are stored in the object metadata

Low-quality cells or empty droplets often have very few genes

Cell doublets or multiplets may have unusually high gene counts

- pbmc[["percent.mt"]] = PercentageFeatureSet(pbmc, pattern = "^MT-") #pbmc[["percent.mt"]]
- Where are QC metrics stored in Seurat? The number of unique genes and total molecules are automatically calculated during CreateSeuratObject()

```
#Show QC metrics for the first 5 cells:
head(pbmc@meta.data, 5)
##
                    orig.ident nCount_RNA nFeature_RNA percent.mt
```

Cells are filtered that have unique feature counts over 2,500 or less than 200

VlnPlot(pbmc, features = c("nFeature_RNA", "nCount_RNA", "percent.mt"), ncol = 3)

nCount_RNA

percent.mt

20

```
## AAACATACAACCAC-1
                                                        779 3.0177759
                                         2419
                           pbmc3k
                                                       1352 3.7935958
                                         4903
 ## AAACATTGAGCTAC-1
                           pbmc3k
 ## AAACATTGATCAGC-1
                                         3147
                                                       1129 0.8897363
                           pbmc3k
 ## AAACCGTGCTTCCG-1
                           pbmc3k
                                         2639
                                                        960 1.7430845
 ## AAACCGTGTATGCG-1
                                          980
                                                        521 1.2244898
                           pbmc3k
Below, QC metrics are visualized and then used to filter cells
```

3000

Cells are filtered that have >5% mitochondrial counts

#Visualize QC metrics as violin plots:

nFeature_RNA

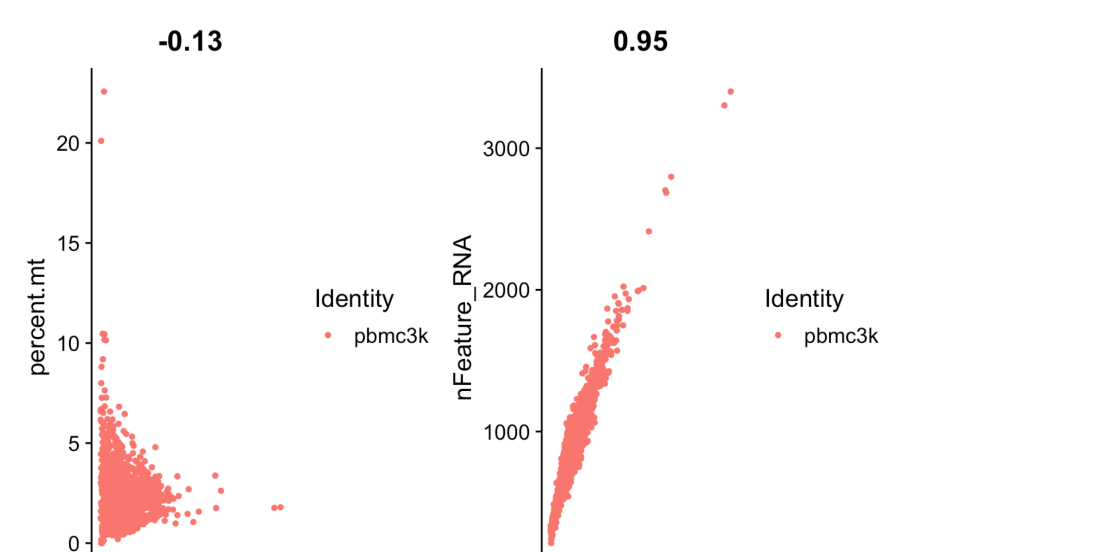
5000 1000015000

nCount_RNA

15 10000 2000

15000

```
10
                              5000
1000
                                                             5
                                                             0
                                  0
             Identity
                                           Identity
                                                                     Identity
#FeatureScatter is typically used to visualize feature-feature relationships
#It can be used for anything calculated by the object, such as columns in object metadata, PC scores,
etc.
plot1 = FeatureScatter(pbmc, feature1 = "nCount RNA", feature2 = "percent.mt")
plot2 = FeatureScatter(pbmc, feature1 = "nCount RNA", feature2 = "nFeature RNA")
plot1 + plot2
             -0.13
                                                          0.95
```



```
#Filter the PBMC data set:
pbmc_filtered <- subset(pbmc, subset = nFeature_RNA > 200 & nFeature_RNA < 2500 & percent.mt < 5)</pre>
```

5000 1000015000

nCount_RNA