Code ▼

HTO Analysis

This is a notebook for HTO Analysis, according to Stoeckius, et al (https://genomebiology.biomedcentral.com/articles/10.1186/s13059-018-1603-1)

After we run CellRanger for the gene expression part, we run the CellRanger for the feature barcodes.

Now it is time to load the samples in R, following Satija's lab hashing vignette (https://satijalab.org/seurat/v3.1/hashing_vignette.html)

Basic setup

Hide

```
# Load packages
library(Seurat)
```

Read in data

Hide

```
# Load the data (Change the paths according to the location of the files on your computer)
ge.data <- Read10X("filtered_ge_020_bc_matrix")
hto.data <- Read10X("filtered_hto_020_bc_matrix", gene.column = 1)</pre>
```

10% data contains more than one type and is being returned as a list containing matrices of each type.

Hide

```
# Select cell barcodes detected by both RNA and HTO In the example datasets we have already
# filtered the cells for you, but perform this step for clarity.
joint.bcs <- intersect(colnames(ge.data), colnames(hto.data$`Antibody Capture`))
# Subset RNA and HTO counts by joint cell barcodes
ex.umis <- ge.data[, joint.bcs]
ex.htos <- as.matrix(hto.data$`Antibody Capture`[, joint.bcs])
# Confirm that the HTO have the correct names
rownames(ex.htos)</pre>
```

```
[1] "0251" "0252" "0253" "0254" "0255" "0256" "0257" "0258"
```

Setup Seurat object and add in the HTO data

```
# Setup Seurat object
ex.hashtag <- CreateSeuratObject(counts = ex.umis)
# Normalize RNA data with log normalization
ex.hashtag <- NormalizeData(ex.hashtag)</pre>
```

```
Performing log-normalization
          30
                     60 70
       20
[----|----|----|----|----|
************
                                                                   Hide
# Find and scale variable features
ex.hashtag <- FindVariableFeatures(ex.hashtag, selection.method = "mean.var.plot")
Calculating gene means
  10 20
          30
                 50
                     60 70 80 90 100%
[----|----|----|----|
************
Calculating gene variance to mean ratios
   10 20
          30
             40
                 50
                     60 70 80
[----|----|----|----|
                                                                   Hide
ex.hashtag <- ScaleData(ex.hashtag, features = VariableFeatures(ex.hashtag))</pre>
Centering and scaling data matrix
  0 %
  _____
```

Adding HTO data as an independent assay

=====| 100%

You can read more about working with multi-modal data here (https://satijalab.org/seurat/multimodal_vignette.html)

```
# Add HTO data as a new assay independent from RNA
ex.hashtag[["HTO"]] <- CreateAssayObject(counts = ex.htos)
# Normalize HTO data, here we use centered log-ratio (CLR) transformation
ex.hashtag <- NormalizeData(ex.hashtag, assay = "HTO", normalization.method = "CLR")</pre>
```

Hide

|-----

Normalizing across features

```
0 % ~calculating
++++++
                       12% ~00s
                      25% ~00s
++++++++++++
++++++++++++++++++
                       38% ~00s
+++++++++++++++++++++++++
                       50% ~00s
62% ~00s
75% ~00s
                      | 88% ~00s
```

Demultiplex cells based on HTO enrichment

Here we use the Seurat function HTODemux() to assign single cells back to their sample origins.

```
Hide
```

```
# If you have a very large dataset we suggest using k function = 'clara'. This is a k-medoid
# clustering function for large applications. You can also play with additional parameters (see
# documentation for HTODemux()) to adjust the threshold for classification. Here we are using the
# default settings
ex.hashtag <- HTODemux(ex.hashtag, assay = "HTO", positive.quantile = 0.99) # , verbose=T
```

```
Cutoff for 0251 : 58 reads
Cutoff for 0252 : 166 reads
Cutoff for 0253: 115 reads
Cutoff for 0254 : 45 reads
Cutoff for 0255 : 39 reads
Cutoff for 0256: 46 reads
Cutoff for 0257 : 140 reads
Cutoff for 0258 : 54 reads
```

Visualize demultiplexing results

Output from running HTODemux() is saved in the object metadata. We can visualize how many cells are classified as singlets, doublets and negative/ambiguous cells.

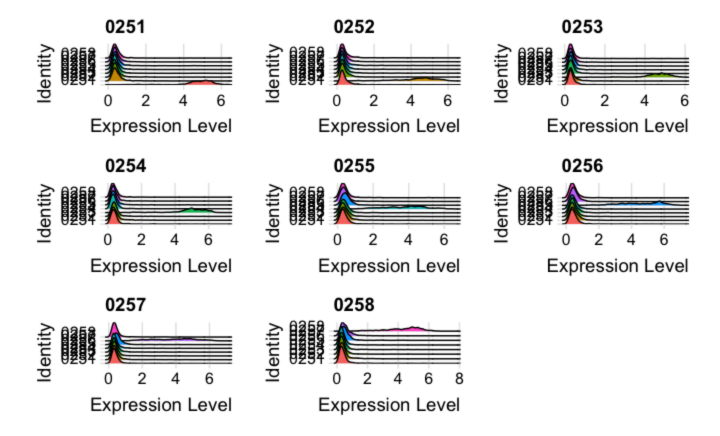
```
# Global classification results
table(ex.hashtag$HTO classification.global)
```

```
Doublet Negative Singlet
   1157
             225
                     6109
```

```
Hide
```

```
# Save sum
n cells <- sum(table(ex.hashtag$HTO classification.global))</pre>
```

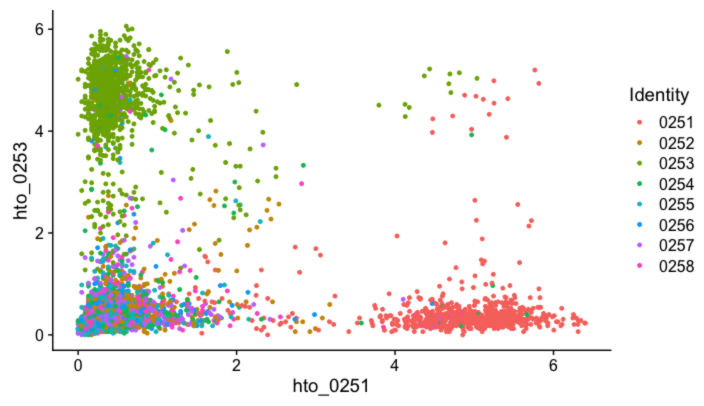
```
# Group cells based on the max HTO signal
Idents(ex.hashtag) <- "HTO_maxID"
RidgePlot(ex.hashtag, assay = "HTO", features = rownames(ex.hashtag[["HTO"]]), ncol = 3)</pre>
```



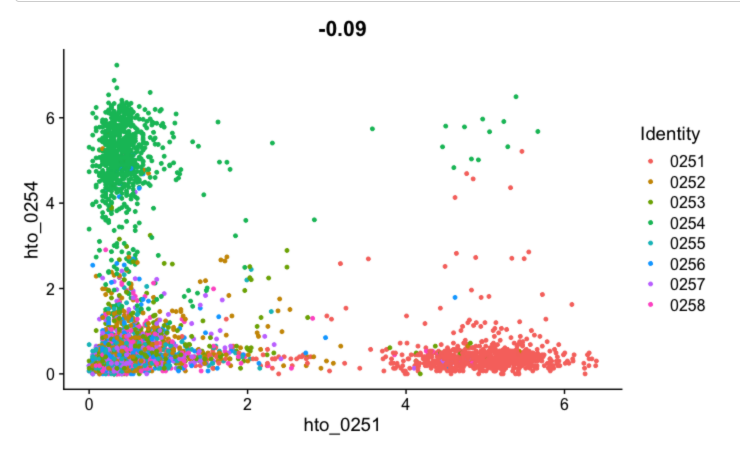
Visualize pairs of HTO signals to confirm mutual exclusivity in singlets

```
FeatureScatter(ex.hashtag, feature1 = "hto_0251", feature2 = "hto_0253")
```





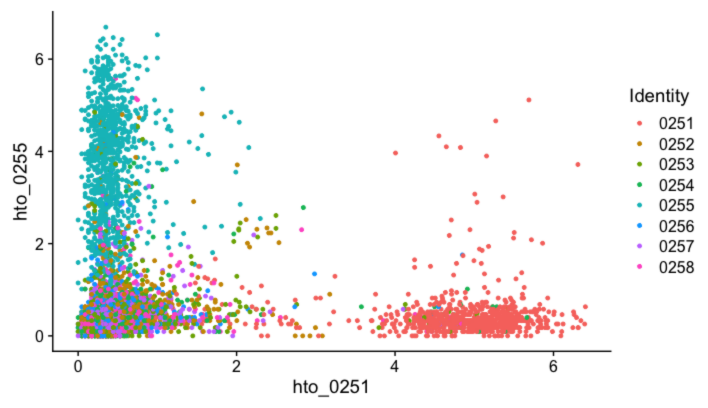
FeatureScatter(ex.hashtag, feature1 = "hto_0251", feature2 = "hto_0254")



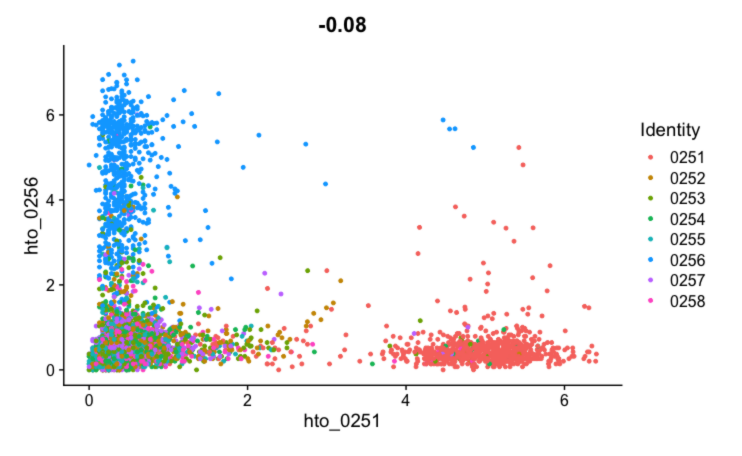
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0251", feature2 = "hto_0255")





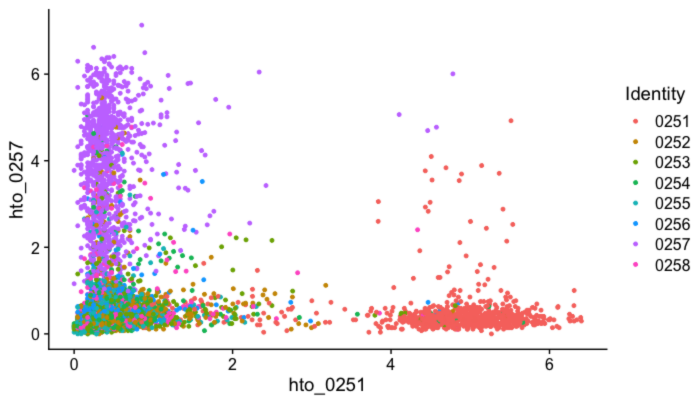
FeatureScatter(ex.hashtag, feature1 = "hto_0251", feature2 = "hto_0256")



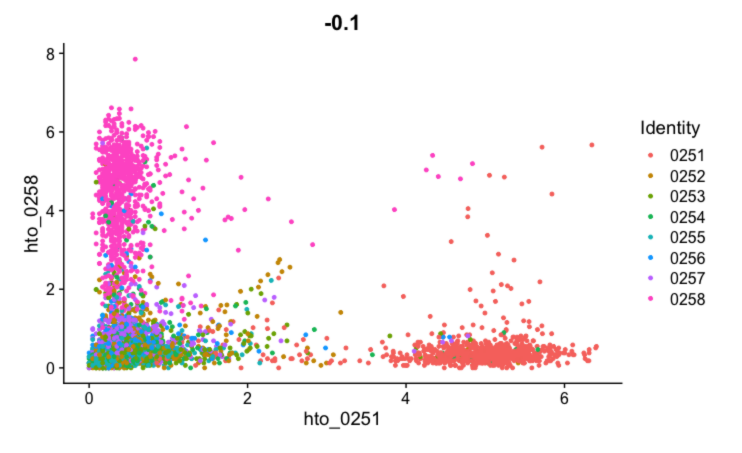
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0251", feature2 = "hto_0257")





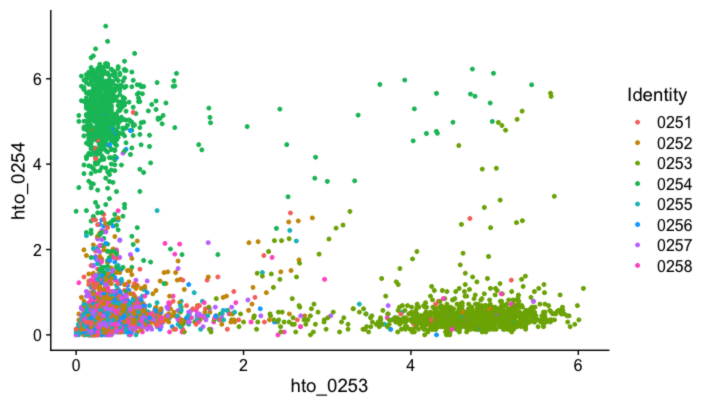
FeatureScatter(ex.hashtag, feature1 = "hto_0251", feature2 = "hto_0258")



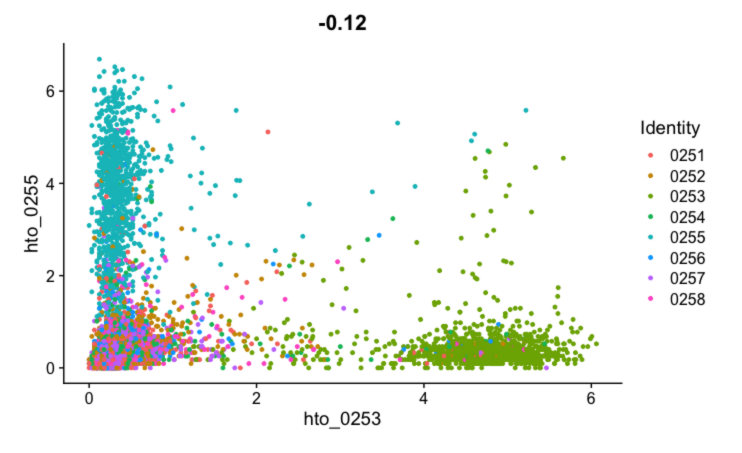
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0253", feature2 = "hto_0254")





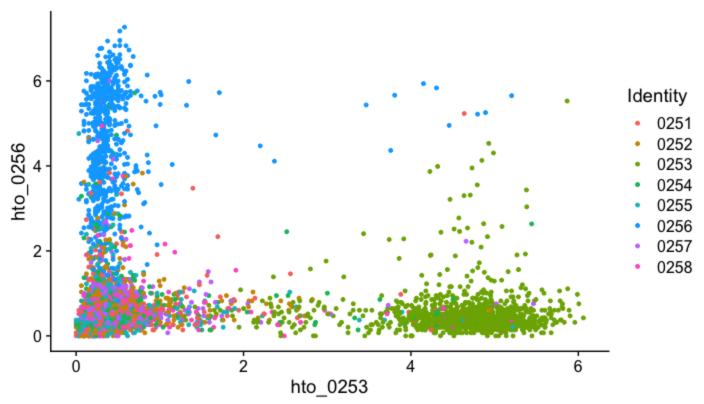
FeatureScatter(ex.hashtag, feature1 = "hto_0253", feature2 = "hto_0255")



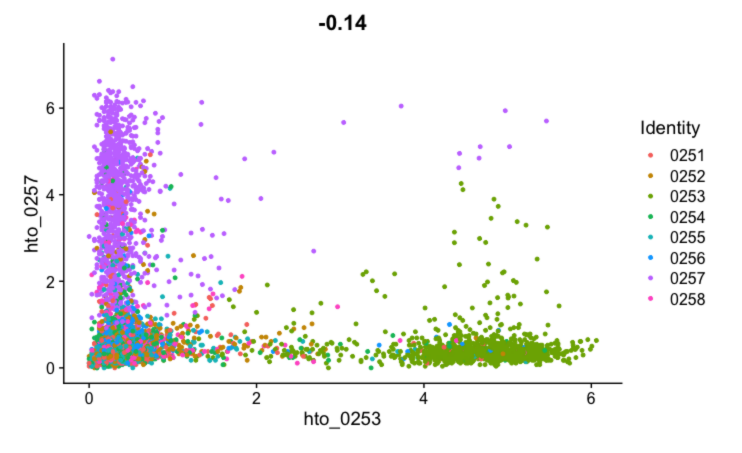
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0253", feature2 = "hto_0256")





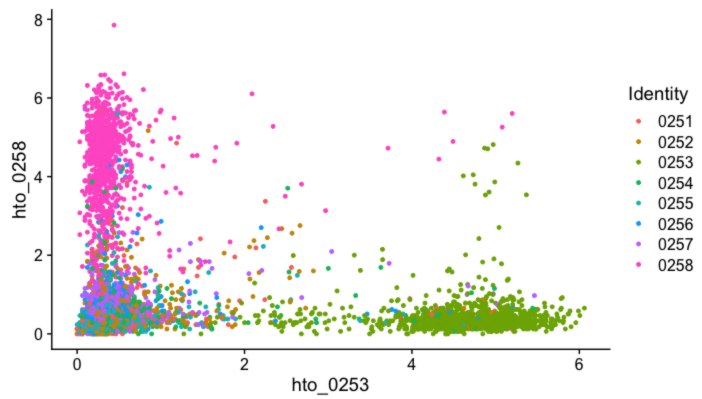
FeatureScatter(ex.hashtag, feature1 = "hto_0253", feature2 = "hto_0257")



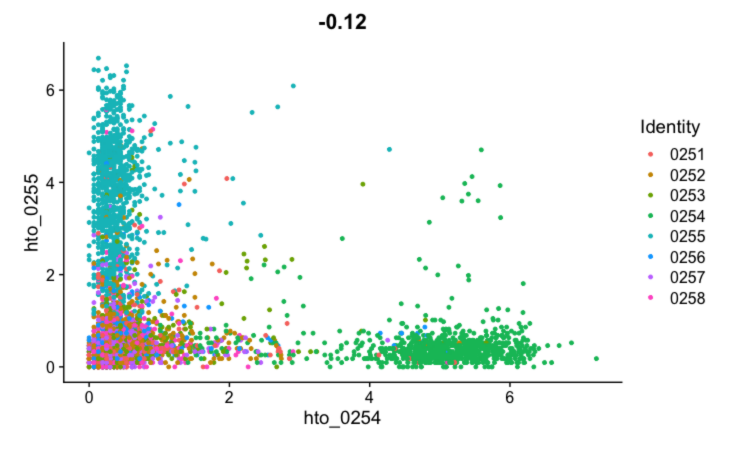
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0253", feature2 = "hto_0258")





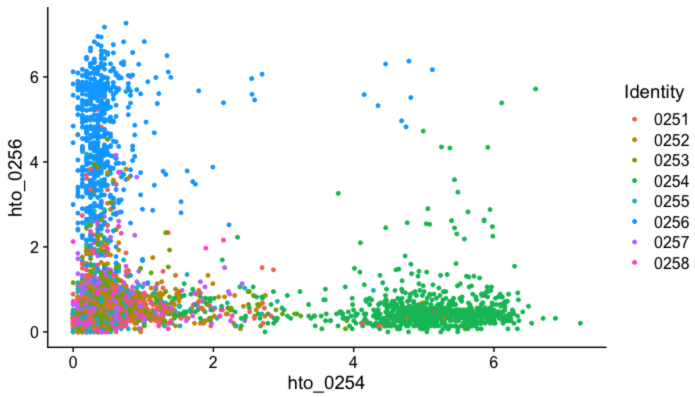
FeatureScatter(ex.hashtag, feature1 = "hto_0254", feature2 = "hto_0255")



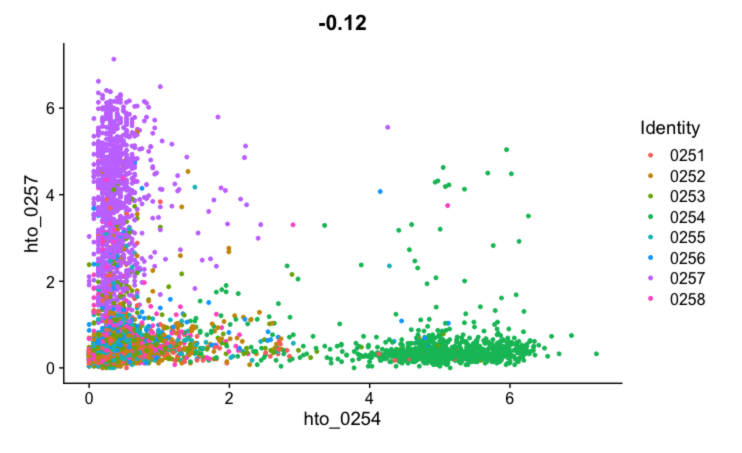
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0254", feature2 = "hto_0256")





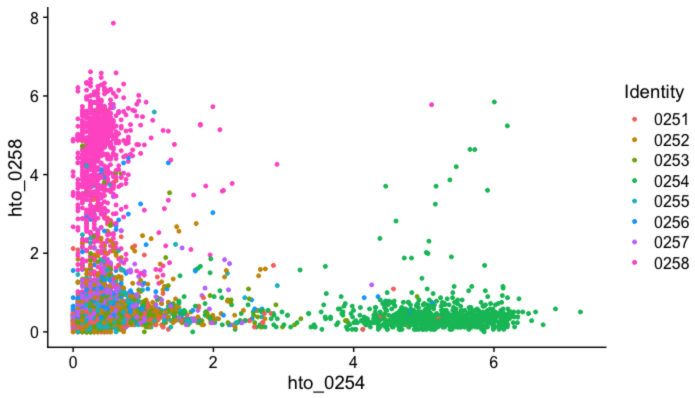
FeatureScatter(ex.hashtag, feature1 = "hto_0254", feature2 = "hto_0257")



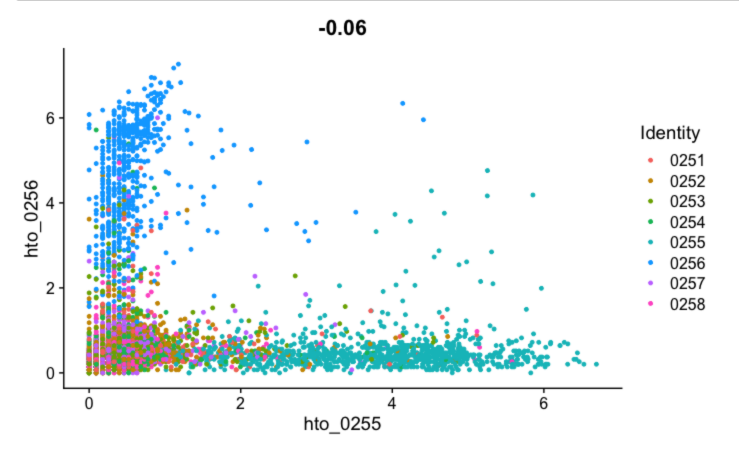
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0254", feature2 = "hto_0258")





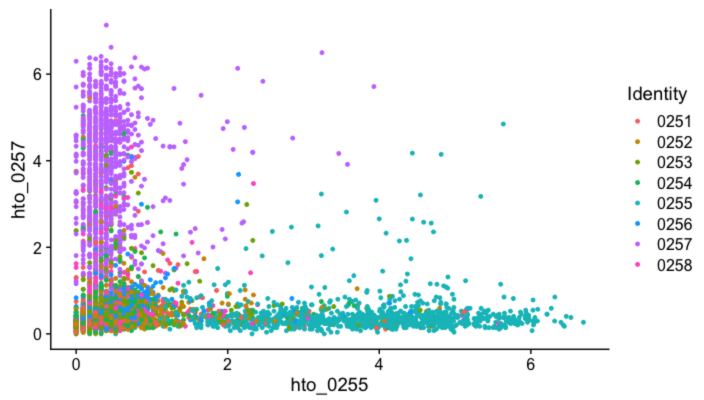
FeatureScatter(ex.hashtag, feature1 = "hto_0255", feature2 = "hto_0256")



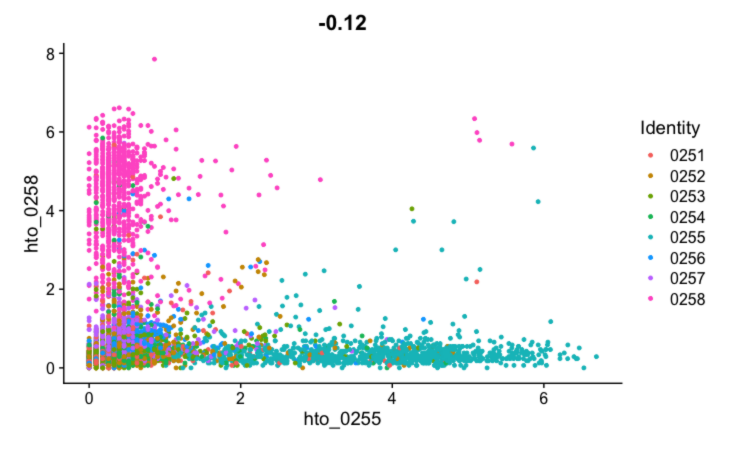
Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0255", feature2 = "hto_0257")



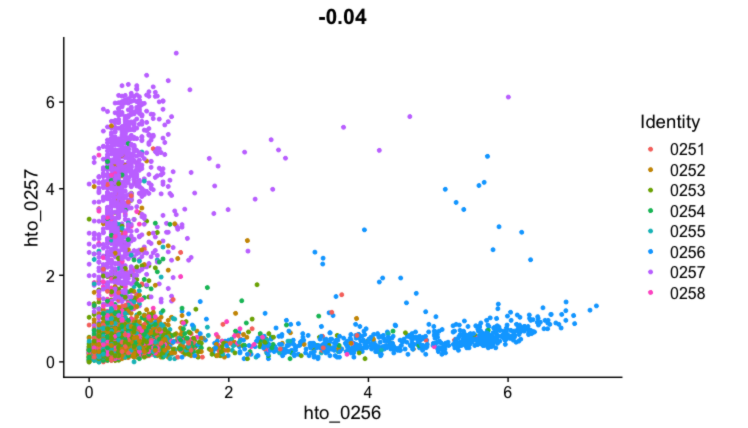


FeatureScatter(ex.hashtag, feature1 = "hto_0255", feature2 = "hto_0258")

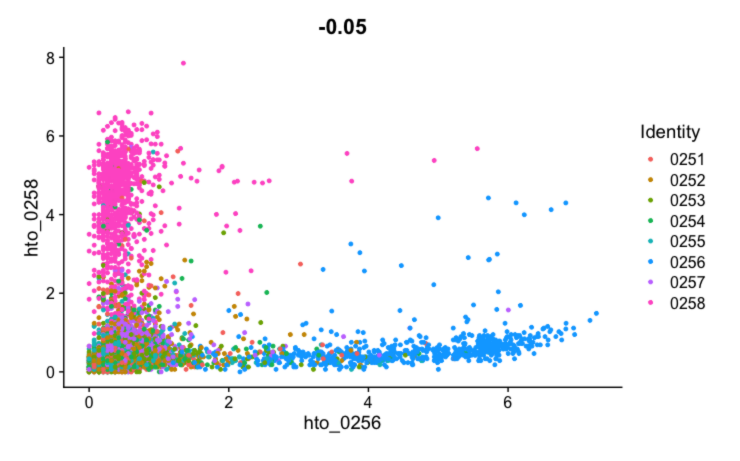


Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0256", feature2 = "hto_0257")

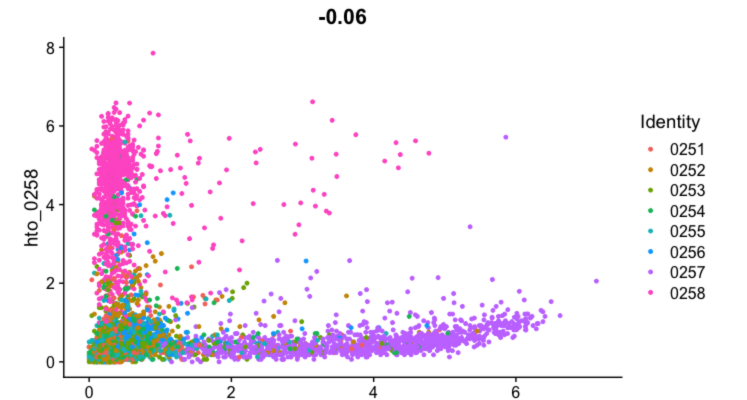


FeatureScatter(ex.hashtag, feature1 = "hto_0256", feature2 = "hto_0258")



Hide

FeatureScatter(ex.hashtag, feature1 = "hto_0257", feature2 = "hto_0258")



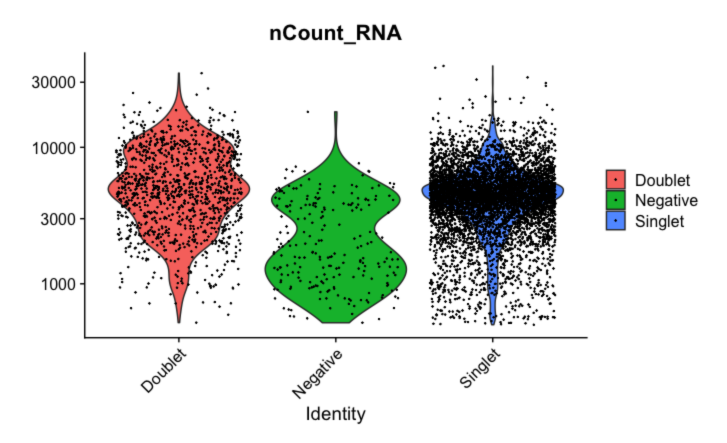
hto_0257

Compare number of UMIs for singlets, doublets and negative cells

```
Idents(ex.hashtag) <- "HTO_classification.global"</pre>
VlnPlot(ex.hashtag, features = "nCount_RNA", pt.size = 0.1, log = TRUE)
```

6

Hide



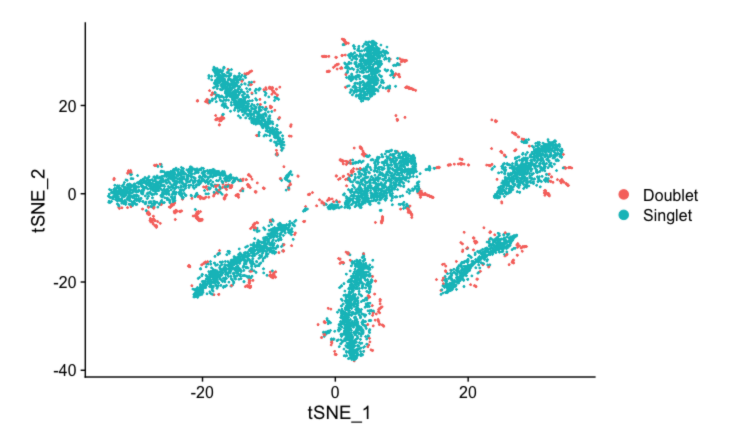
Generate a two dimensional tSNE embedding for HTOs. Here we are grouping cells by singlets and doublets for simplicity.

```
# First, we will remove negative cells from the object (if any)
ex.hashtag.subset <- subset(ex.hashtag, idents = "Negative", invert = TRUE)
# Calculate a distance matrix using HTO
# (use the subset if you just created in case you had negative cells removed)
hto.dist.mtx <- as.matrix(dist(t(GetAssayData(object = ex.hashtag.subset, assay = "HTO"))))
# Calculate tSNE embeddings with a distance matrix
# (use the subset if you just created in case you had negative cells removed)
ex.hashtag.subset <- RunTSNE(ex.hashtag.subset, distance.matrix = hto.dist.mtx, perplexity = 100)</pre>
```

Adding a command log without an assay associated with it

Hide

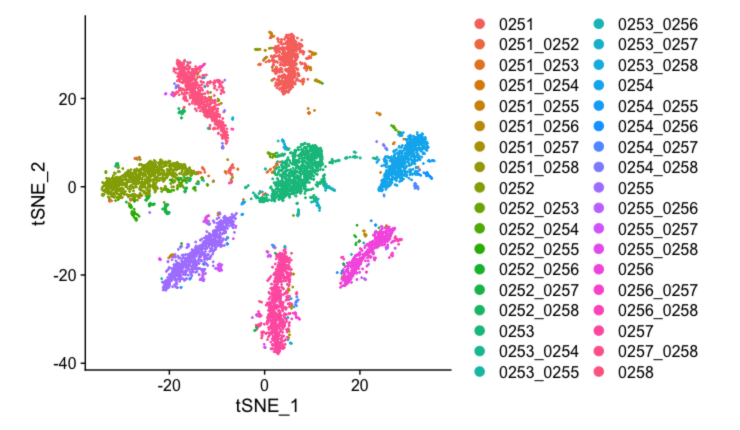
DimPlot(ex.hashtag.subset)



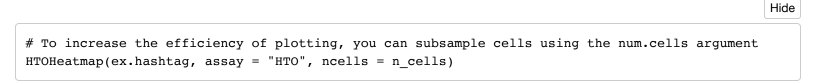
Visualize the more detailed classification result. Here, you should be able to see that each of the small clouds on the tSNE plot corresponds to one of the possible doublet combinations.

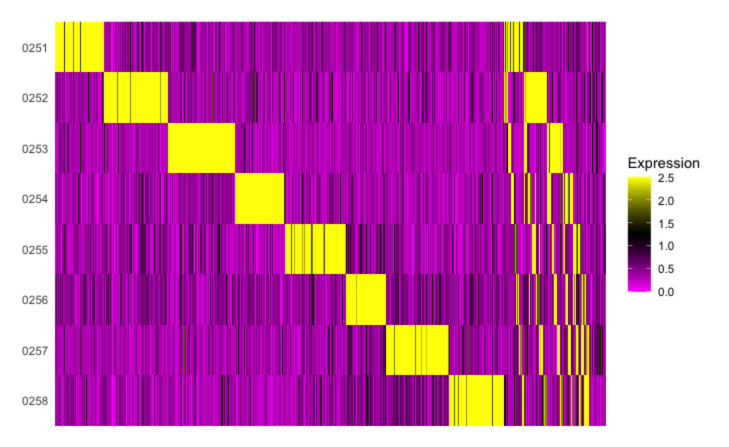
Hide

Idents(ex.hashtag.subset) <- 'HTO_classification'
DimPlot(ex.hashtag.subset)</pre>



Create an HTO heatmap, based on Figure 1C in the Cell Hashing paper.





Cluster and visualize cells using the usual scRNA-seq workflow, and examine for the potential presence of batch effects.

```
Hide
```

```
# Extract the singlets
ex.singlets <- subset(ex.hashtag, idents = "Singlet")
# Select the top 1000 most variable features
ex.singlets <- FindVariableFeatures(ex.singlets, selection.method = "mean.var.plot")</pre>
```

```
Calculating gene means
             50
                60 70
  10
     20
                     80
[----|----|----|----|
***********
Calculating gene variance to mean ratios
    20
        30
           40
             50
                60 70
                         90
[----|----|----|----|----|
***********
```

```
# Scaling RNA data, we only scale the variable features here for efficiency
ex.singlets <- ScaleData(ex.singlets, features = VariableFeatures(ex.singlets))</pre>
```

```
# Run PCA
ex.singlets <- RunPCA(ex.singlets, features = VariableFeatures(ex.singlets))</pre>
```

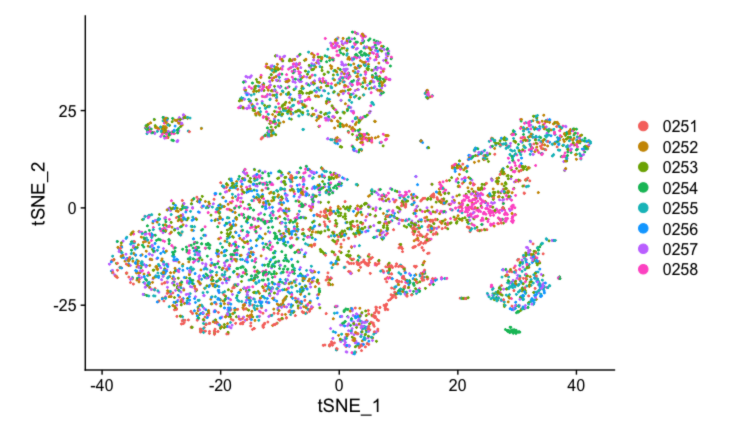
```
Positive: IL32, IL7R, RPS27, LINCO0861, CCR7, CTSW, SYNE2, TRBC1, CST7, GZMA
       CCL5, KLRK1, NKG7, PRF1, PYHIN1, RPS8, KLRD1, CD8A, KLRB1, GNLY
       GZMH, KLRG1, IL2RB, MATK, CD8B, PRDM1, MT-CYB, LINC01871, HOPX, FGFBP2
Negative: FCN1, IFI30, CST3, LYZ, MNDA, S100A9, SERPINA1, SPI1, S100A8, AIF1
       CD68, CYBB, CTSS, LST1, VCAN, CSTA, TNFAIP2, TYMP, CD14, MS4A6A
       CFD, FGL2, CEBPD, GRN, PSAP, CSF3R, FPR1, TYROBP, CLEC7A, HCK
PC 2
Positive: NKG7, PRF1, CST7, GNLY, GZMB, GZMA, KLRD1, FGFBP2, CTSW, SPON2
       CCL5, KLRF1, GZMH, ADGRG1, HOPX, CX3CR1, MATK, CCL4, FCGR3A, TBX21
       KLRK1, S1PR5, CLIC3, IL2RB, TTC38, FCRL6, SH2D1B, PRSS23, PTGDR, PLEK
Negative: CD79A, MS4A1, BANK1, FCRLA, RALGPS2, IGHM, LINC00926, NIBAN3, CD19, BLK
       AFF3, SPIB, CCR7, CD79B, BLNK, RPS8, FCER2, TNFRSF13C, FCRL1, POU2AF1
       CD22, P2RX5, HLA-DOB, HLA-DQA1, GNG7, CD24, IGKC, VPREB3, SWAP70, TCL1A
PC 3
Positive: MS4A1, CD79A, BANK1, FCRLA, IGHM, LINC00926, CD19, NIBAN3, HLA-DQA1, CD79B
       SPIB, RALGPS2, BLK, HLA-DPA1, BLNK, FCRL1, HLA-DPB1, AFF3, FCER2, HLA-DQB1
       SWAP70, CD22, CD74, POU2AF1, TNFRSF13C, PDLIM1, CD72, HLA-DOB, P2RX5, HLA-DRA
Negative: IL7R, IL32, CCR7, SERINC5, LINC00861, CISH, LINC01550, ADTRP, TSHZ2, DPP4
       PDE3B, LINC00402, S100A12, RETREG1, BEX3, S100A8, AIF1, S100A9, RNF157, LRRN3
       VCAN, EPHA1-AS1, CD14, S100A11, HNRNPLL, PI16, PLCL1, HAPLN3, RBP7, AC009041.2
PC 4
Positive: MT-CO1, MT-ND5, MT-CO2, MT-CYB, MT-ND3, MT-ND6, MTRNR2L12, MT-ATP8, XIST, NEAT1
       MTRNR2L8, NKTR, MACF1, NPIPB5, LINC00342, AHNAK, PCNX1, KIAA1109, DENND4A, SUGP2
       ARHGAP26, SYNE2, MDM4, ANKRD36C, RBM5, RICTOR, FTX, PARP14, NLRC5, GOLGA8A
Negative: RPS8, RPS27, FTH1, FTL, CDKN1C, FCGR3A, IFITM3, LYPD2, HMOX1, COTL1
       SMIM25, TCF7L2, SIGLEC10, FCER1G, HLA-DPA1, CDK2AP1, CXCL16, FAM110A, LRRC25, WARS
       DUSP5, MAFB, CHST2, MS4A7, PSAP, LST1, AIF1, MT2A, PILRA, CRIP1
PC 5
Positive: CDKN1C, TCF7L2, SIGLEC10, LYPD2, SMIM25, MS4A7, FCGR3A, CXCL16, HMOX1, WARS
       MYOF, LRRC25, MAFB, NR4A1, FAM110A, DUSP5, IFITM3, PECAM1, SLC2A6, EPB41L3
       MT-CO1, MT-CYB, PILRA, GCH1, MT-ATP8, IFIT2, MT-CO2, HCK, LST1, C5AR1
Negative: S100A12, VCAN, S100A8, PADI4, S100A9, MCEMP1, CD14, PLBD1, NCF1, RBP7
       CSF3R, CD36, JUN, CLEC4E, FOLR3, AC007952.4, NAIP, CYP1B1, AC020656.1, AC020916.1
      RPS8, CDA, CXCL8, CES1, AC245014.3, THBS1, NFE2, CYP27A1, MS4A6A, BLVRB
```

We select the top 10 PCs for clustering and tSNE based on PCElbowPlot
ex.singlets <- FindNeighbors(ex.singlets, reduction = "pca", dims = 1:10)</pre>

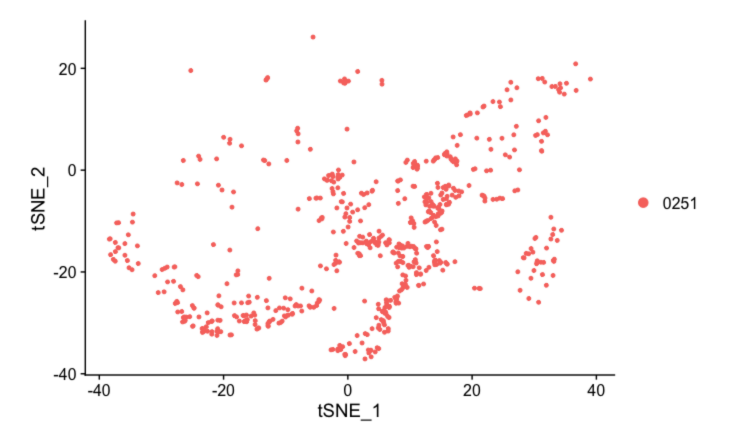
Computing nearest neighbor graph Computing SNN

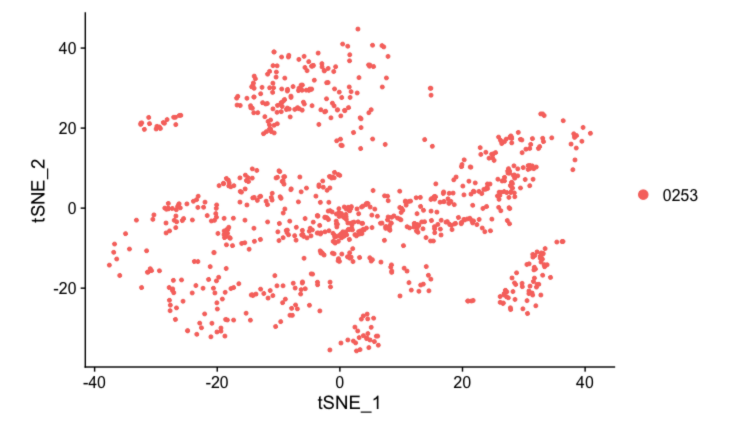
PC 1

```
ex.singlets <- FindClusters(ex.singlets, resolution = 0.6, verbose = FALSE)
ex.singlets <- RunTSNE(ex.singlets, reduction = "pca", dims = 1:10)
# Projecting singlet identities on TSNE visualization
DimPlot(ex.singlets, group.by = "HTO_classification")</pre>
```

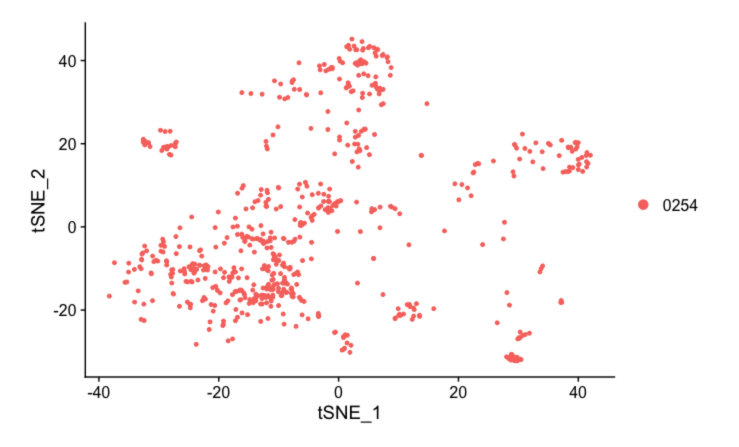


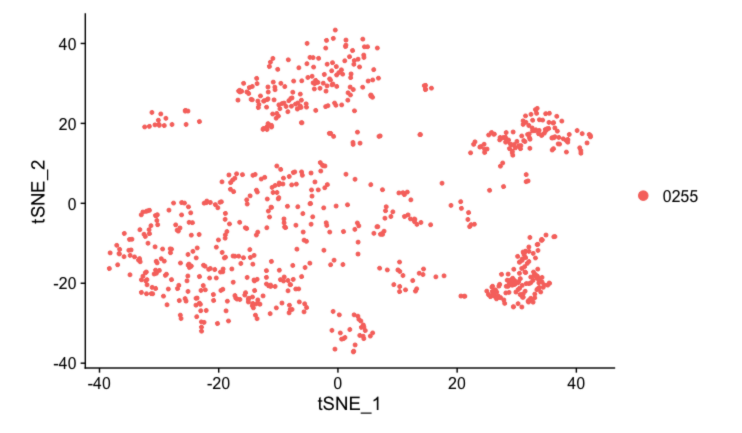
Projecting singlets for each hash ID separately
DimPlot(ex.singlets[, ex.singlets\$hash.ID == "0251"], group.by = "HTO_classification")



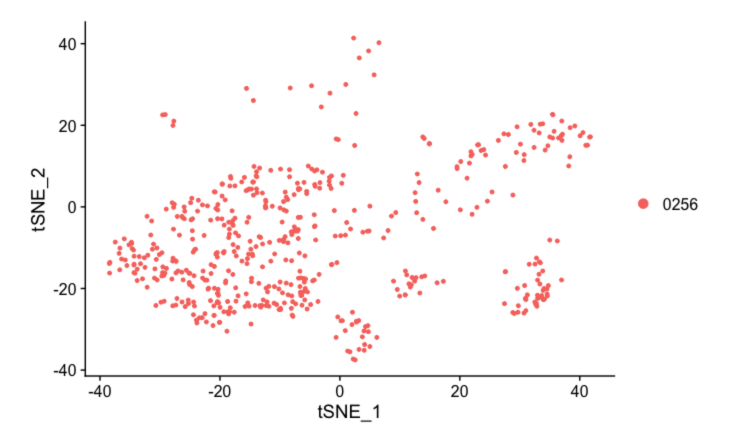


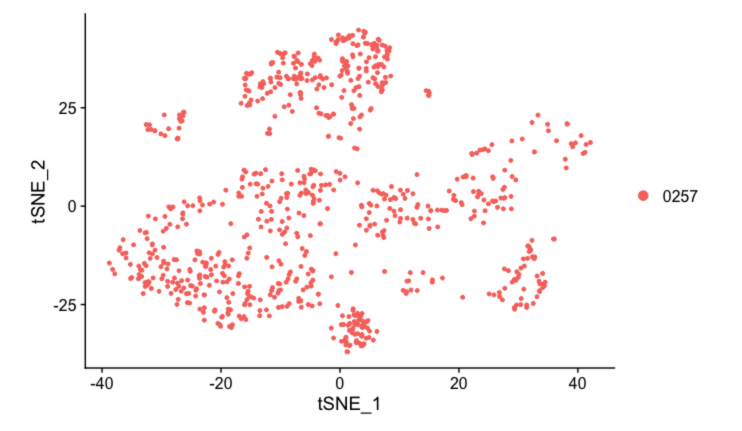
 $\label{eq:decomposition} \mbox{DimPlot(ex.singlets[, ex.singlets$hash.ID == "0254"], group.by = "HTO_classification")}$



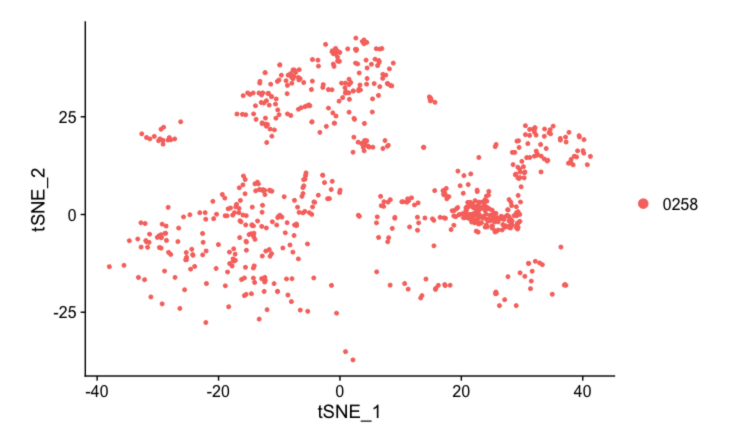


 $\label{eq:decomposition} \mbox{DimPlot(ex.singlets[, ex.singlets$hash.ID == "0256"], group.by = "HTO_classification")}$





```
\label{eq:decomposition} \mbox{DimPlot(ex.singlets[, ex.singlets$hash.ID == "0258"], group.by = "HTO_classification")}
```



```
# Visualize HTOs on RNA clusters
FeaturePlot(ex.singlets, features = rownames(ex.hashtag[["HTO"]]), ncol = 3)
```

Could not find 0251 in the default search locations, found in HTO assay insteadCould not find 0252 in the default search locations, found in HTO assay insteadCould not find 0253 in the default search locations, found in HTO assay insteadCould not find 0254 in the default search locations, found in HTO assay insteadCould not find 0255 in the default search locations, found in HTO assay insteadCould not find 0256 in the default search locations, found in HTO assay insteadCould not find 0257 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations, found in HTO assay insteadCould not find 0258 in the default search locations in HTO assay insteadCould not find 0258 in the default search locations in HTO assay insteadCould not find 0258 in the default search locations in HTO assay insteadCould not fin

